Final Technical Report
on

Dimethyl Ether (DME)-Fueled Shuttle Bus Demonstration Project

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by
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ABSTRACT

The objectives of this research and demonstration program are to convert a campus shuttle bus to operation on dimethyl ether, a potential ultra-clean alternative diesel fuel. To accomplish this objective, this project includes laboratory evaluation of a fuel conversion strategy, as well as, field demonstration of the DME-fueled shuttle bus. Since DME is a fuel with no lubricity (i.e., it does not possess the lubricating quality of diesel fuel), conventional fuel delivery and fuel injection systems are not compatible with dimethyl ether. Therefore, to operate a diesel engine on DME one must develop a fuel-tolerant injection system, or find a way to provide the necessary lubricity to the DME. In this project, we have chosen the latter strategy in order to achieve the objective with minimal need to modify the engine. Our strategy is to blend DME with diesel fuel, to obtain the necessary lubricity to protect the fuel injection system and to achieve low emissions. The bulk of our efforts over the past year were focused on the conversion of the campus shuttle bus. This process, started in August 2001, took until April 2002 to complete. The process culminated in an “event” to celebrate the launching of the shuttle bus on DME-diesel operation on April 19, 2002. The design of the system on the shuttle bus was patterned after the system developed in the engine laboratory, but also was subjected to a rigorous failure modes effects analysis (FMEA, referred to by Air Products as a “HAZOP” analysis) with help from Dr. James Hansel of Air Products. The result of this FMEA was the addition of layers of redundancy and over-pressure protection to the system on the shuttle bus. The system became operational in February 2002. Preliminary emissions tests and basic operation of the shuttle bus took place at the Pennsylvania Transportation Institute’s test track facility near the University Park airport. After modification and optimization of the system on the bus, operation on the campus shuttle route began in early June 2002. However, the work and challenges continued as it was been difficult to maintain operability of the shuttle bus due to fuel and component difficulties. In late June 2002, the pump head itself developed operational problems (loss of smooth function) leading to excessive stress on the magnetic coupling and excessive current draw to operate. A new pump head was installed on the system to alleviate this problem and the shuttle bus operated successfully on DME blends from 10 – 25 vol% on the shuttle bus loop until September 30, 2002. During the period of operation on the campus loop, the bus was pulled from service, operated at the PTI test track and real-time emissions measurements were obtained using an on-board emissions analyzer from Clean Air Technologies International, Inc. Particulate emissions reductions of 60% and 80% were observed at DME blend ratios of 12 vol.% and 25 vol.%, respectively, as the bus was operated over the Orange County driving cycle. Increases in NOx, CO and HC emissions were observed, however. In summary, the conversion of the shuttle bus was successfully accomplished, particulate emissions reductions were observed, but there were operational challenges in the field. Nonetheless, we were able to demonstrate reliable operation of the shuttle bus on DME-diesel blends.
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EXECUTIVE SUMMARY

The objectives of this research and demonstration program are to convert a campus shuttle bus to operation on dimethyl ether, a potential ultra-clean alternative diesel fuel. To accomplish this objective, this project includes laboratory evaluation of a fuel conversion strategy, as well as, field demonstration of the DME-fueled shuttle bus. Since DME is a fuel with no lubricity (i.e., it does not possess the lubricating quality of diesel fuel), conventional fuel delivery and fuel injection systems are not compatible with dimethyl ether. Therefore, to operate a diesel engine on DME one must develop a fuel-tolerant injection system, or find a way to provide the necessary lubricity to the DME. In this project, we have chosen the latter strategy in order to achieve the objective with minimal need to modify the engine. Our strategy is to blend DME with diesel fuel, to obtain the necessary lubricity to protect the fuel injection system and to achieve low emissions. Sponsorship comes from the National Energy Technology Laboratory (Project Manager Shelby Roger, and Program Manager John Winslow), the Pennsylvania Department of Environmental Protection Alternative Fuel Incentive Grant Program (Program Manager Susan Summers) and Air Products and Chemicals, Inc. (Program Manager Ed Heydom).

To date, our activities have covered two areas: laboratory investigations and field investigations. The laboratory studies have included work with a Navistar V-8 turbodiesel engine, demonstration of engine operation on DME-diesel blends and instrumentation for evaluating fuel properties. The field studies have involved performance, efficiency and emissions measurements with the Champion Motorcoach “Defender” shuttle bus and conversion of the shuttle bus operation on DME-diesel blends.

Within the Diesel Combustion and Emissions Laboratory in the Penn State Energy Institute, we have installed and equipped a Navistar V-8 direct-injection turbodiesel engine for measurement of gaseous and particulate emissions and examination of the impact of fuel composition on diesel combustion. We have also reconfigured a high-pressure viscometer for studies of the viscosity, bulk modulus (compressibility) and miscibility of blends of diesel fuel, dimethyl ether and lubricity additives. Our results include baseline emissions, performance and combustion measurements on the Navistar engine for operation on a federal low sulfur diesel fuel (300 ppm S). During the course of this project, we constructed an entire engine test facility which ultimately led to tests of DME-diesel blends up to 30 vol.% (25 wt.%) DME in the Navistar engine. The engine results included detailed analysis of cylinder pressure trace data, injection parameters reported by the ECM on the engine and emissions data. The results from the laboratory studies have been summarized in two a Society of Automotive Engineers technical papers, an American Chemical Society preprint and were summarized in detail in the MS Theses of Elana Chapman and Shirish Bhide.

We have also performed viscosity measurements on diesel fuel, DME and blends of DME in diesel. These tests have verified that DME has a much lower viscosity than the diesel fuel and that the viscosity of the blended fuel is also much lower than the diesel base fuel. This has implications for the injection and atomization of the DME/diesel blends. The test results on fuel viscosity were summarized in an ACS Preprint presented at the 2001 ACS National Meeting in Chicago.
The ultimate goal of this project is the conversion of the campus shuttle bus. This process, started in August 2001, took until April 2002 to complete. The process culminated in an “event” to celebrate the launching of the shuttle bus on DME-diesel operation on April 19, 2002. The design of the system on the shuttle bus was patterned after the system developed in the engine laboratory, but also was subjected to a rigorous failure modes effects analysis (FMEA, referred to by Air Products as a “HAZOP” analysis) with help from Dr. James Hansel of Air Products. The result of this FMEA was the addition of layers of redundancy and over-pressure protection to the system on the shuttle bus. The system became operational in February 2002. Preliminary emissions tests and basic operation of the shuttle bus took place at the Pennsylvania Transportation Institute’s test track facility near the University Park airport.

After another month’s worth of modification and optimization of the system on the bus, operation on the campus shuttle route began in early June 2002. However, the work and challenges have continued as it has been difficult to maintain operability of the shuttle bus due to fuel and component difficulties. The difficulties with the converted fuel system have arisen in two areas: operation of the converted fueling system, which circulates fuel to the engine; and operation of the engine, which has run very rough at times and is emitting a significant amount of “blow by” that indicates the engine may need an overhaul.

The difficulties with the fuel circulation system were primarily with the gear pump which draws DME-diesel fuel blend from the onboard mixture tank. The pump developed operational problems that appeared to either be cavitation (possible if DME vaporized in our fueling system) or loss of magnetic coupling between the pump head and electric motor. The latter turns out to be the main problem, although this magnetic coupling has been replaced on the vehicle once already. In late June 2002, it was determined that the pump head itself developed operational problems (loss of smooth function) leading to excessive stress on the magnetic coupling and excessive current draw to operate. A new pump head was installed on the system to alleviate this problem and the shuttle bus operated successfully on DME blends from 10 – 25 vol% on the shuttle bus loop until September 30, 2002. During the period of operation on the campus loop, the bus was pulled from service, operated at the PTI test track and real-time emissions measurements were obtained using an on-board emissions analyzer from Clean Air Technologies International, Inc. Particulate emissions reductions of 60% and 80% were observed at DME blend ratios of 12 vol.% and 25 vol.%, respectively, as the bus was operated over the Orange County driving cycle. Increases in NOx, CO and HC emissions were observed, however.

In summary, the conversion of the shuttle bus was successfully accomplished, particulate emissions reductions were observed, but there were operational challenges in the field. Nonetheless, we were able to demonstrate reliable operation of the shuttle bus on DME-diesel blends.
EXPERIMENTAL
This project is driven by Air Products’ interest in the development of markets for the Liquid Phase Dimethyl Ether process technology, the state’s interest in development of transportation fuel usage from Pennsylvania resources (e.g., coal) and the Department of Energy’s interest in ultra-clean transportation fuels. In this project we (Penn State Energy Institute and Air Products and Chemicals, Inc.) have determined how to effect the conversion of a shuttle bus equipped with a Navistar DI turbo-diesel engine to operation on Dimethyl Ether (DME). To accomplish this goal, we have been examining the co-firing of the engine on diesel fuel and dimethyl ether, using the diesel fuel as a lubricating agent to protect the fuel pump and fuel injection system from excessive wear. Dimethyl ether has no natural lubricity, making it antagonistic to fuel system components.

The work consists of two parallel efforts. One is the development of a conversion process to operate a diesel engine on dimethyl ether using conventional fuel injection equipment. The other is an evaluation of the performance, emissions and efficiency of a shuttle bus equipped with a diesel engine and operated on dimethyl ether.

Development of the conversion strategy required first the construction of an engine test facility, and then a methodology for fueling the engine on DME. A pressurized fuel delivery system was developed and successfully implemented on the laboratory engine. With the engine and test facility, we have demonstrated that the laboratory engine can be operated effectively on the blended fuel.

The shuttle bus conversion relied on an adaptation of the laboratory conversion strategy for use in the field. While the conversion strategy was under development, the shuttle bus (Champion “Defender” model) that serves as a faculty/staff shuttle on the University Park campus operated for a period of three and one half years on diesel fuel, accumulating roughly 90,000 miles prior to conversion. Performance and efficiency tests were performed after the break-in of the vehicle during operation on diesel fuel, and emissions tests were performed after the conversion to dimethyl ether operation.

To accomplish the operation of the campus shuttle bus on DME-diesel blends, we designed a large scale, dedicated DME fueling station for delivery of DME-diesel blends. The cost of the initial design, however, was well beyond the level of funding available for the fueling station. Instead, we turned to a simpler plan that involved a two-stage fueling process. The person refueling the shuttle bus would use the existing diesel fuel pumps for University service vehicles to fill the bus’ main tank with diesel fuel. Then, the driver will back up to an adjacent storage location for the DME. Diesel fuel was then transferred to the DME-diesel mixture tank from the main diesel tank, and finally, DME was added to the DME-diesel mixture tank to complete the refueling of the shuttle bus. Once filled, the mixture tank is pressurized with a blanket of helium gas to keep the DME in the liquid phase and maintain the blend ratio of the two fuels.

The objectives of the laboratory testing have been to determine the compositions of fuel and additive blends that will permit long term operation of the T444E engine on dimethyl ether. We have successfully operated the laboratory engine on blends with up to 30 wt.% DME. Based on
these tests, we determined that the optimal blend ratio for the field vehicle is 25 wt.% DME in diesel fuel.

Through collaboration with the Tribology Laboratory in Penn State's Chemical Engineering Department, we have characterized the viscosity, compressibility and miscibility of blends of DME, diesel fuel and the additives under pressures and temperatures relevant to the fuel injection system. These tests have used a high pressure viscometer adapted to these specific experiments. In addition a closely related project with support from NETL, we are constructing an injector durability experiment to determine the time to failure and boundary of the of fuel composition which can lead to injector failure. Also, we have modified an existing pin-on-ring apparatus to investigate the lubricity of DME blends with diesel fuel and lubricity additives.

On the outset of this project, we believed that the most significant technical challenge in this project would be accounting for the lubricating quality of the fuel and additive mixtures. This challenge of working with DME involves two considerations of the mixtures: (1) the lubricating quality of the mixtures; and (2) the phase behavior of the mixtures, given that advanced diesel fuel injection systems use injection pressures that exceed the critical pressure of DME. At higher DME blend ratios, we observed some instability in the laboratory engine, perhaps due to generation of excessive amounts of DME vapor. This aspect of engine operation on DME-diesel blends was confirmed during the field evaluation of the campus shuttle bus. At a shift of sustained load to idling, the engine would idle roughly, surging and sagging in speed, typically when the vehicle was fueled at the higher concentrations of DME, 25 wt.% and on warm days.

However, it became clear during the field work with the shuttle bus, that the greatest challenge was instead getting the supporting components within the pressurized fueling system to perform properly over extended periods of time. System integrity and durability became far more significant challenges than management of the volatility and lubricity of the DME-diesel blends.

The rest of this report consists of a section for each of the major activities (tasks) under the project.

- Laboratory Engine Results
- Viscometer Results
- Fuel Station Construction and Shuttle Bus Conversion

Finally, there are extensive appendices included with this report (in electronic format) that document the technical progress on the project through the preparation of technical papers and theses.
RESULTS AND DISCUSSION
Laboratory Engine Results

The laboratory engine, a Navistar T444E 190hp V-8 Turbodiesel has been operational since October 1999 in our engine test cell. The engine is outfitted with numerous thermocouple sensors, an in-cylinder pressure sensor (in Cylinder #1) and the exhaust system has a number of ports for sampling particulate emissions, gaseous emissions (including total hydrocarbons) and particulate composition (via thermal analysis). The figure below shows a digital photo of the engine in the test cell prior to the fuel system conversion.

![Photo of the Navistar T444E Turbodiesel Engine in the 450 hp Engine Test Cell Prior to Conversion to DME-Diesel Blends](image)

The AVL steady state 8 mode test had been selected to do the emissions comparison studies for different fuel additives and blend ratios of DME to diesel. During previous tests, a comparison of exhaust emissions was made with two different oxygenates, the CETANER™ fuel additive and
Dimethyl ether (DME). These were blended with a federal low sulfur diesel fuel that will also serve as the baseline fuel. The oxygenates were blended such that the oxygen content of the mixture was 2% by mass (reported in SAE Technical Paper NO. 2000-01-2887). More recently, tests were completed at 5 and 10 wt.% oxygen addition by blending DME and diesel fuel, which is equivalent to 15 vol.% (12.5 wt.%) and 30 vol.% (25 wt.%) DME addition, respectively. The Appendix includes a copy of a Society of Automotive Engineers technical paper that was written on the results from the DME-diesel blend studies.

![Figure 2. Navistar T444E showing conversion to DME-diesel operation.](image)

**Emissions Evaluation - Low DME/Diesel Concentration Tests**

Engine tests had been completed with conservative blends of Dimethyl Ether (DME) blended in diesel fuel. The objective of these tests was to determine the effect of the fuel blends on emissions. As with most all oxygenated fuel blends, there was a particulate matter reduction observed as the oxygen concentration was increased, for most all of the 8 modes tested (variations of speed and load). However, the other emissions showed scattered results across the modes. This data was prepared and presented in two papers in 2000 and 2001 [1, 2], which are included with this report in the Appendix.
As the engine load was increased and regardless of engine speed, the particulate emission was lowered, except for mode 7. To further understand the variations in the emissions data, the data from the engine electronics was reviewed. Of specific interest was the actions of the fuel injectors, as shown in Figures 3, 4 and 5, since the electronic controls of the engine had not been changed from the production calibration, yet the fuel density was different.

Figure 3. Engine Injection Timing
(* Injection timing is relative to the number of degrees before 0 degrees: TDC (Top Dead Center))
Figure 4. Engine Injection Pressure

The effect seen in Mode 7 may be due to a change in the injection timing. As shown in Figure 3 and Figure 4, the injection timing and injection pressure were changing as commanded by the engine control for changing speed and load. This is true for the low as well as the high engine speeds. The trends follow what would be expected based on the work by Kajitani and coworkers [3]. Their work showed that as the mean effective pressure increases, which correlates with increasing load, for an engine speed of 960 RPM, so does soot emissions based on the Bosch Smoke Number with diesel fuel. In their work, the DME emissions were close to zero. In this work, with increasing content of DME, the soot emissions decreased.

As has been shown in previous work by Liotta and coworkers, this particulate reduction is due to a reduction in the soot portion of the emission, and would result in a percentage increase in the soluble organic fraction (SOF) portion [4]. This has also been confirmed more recently by Sidhu and coworkers [5], with DME giving the highest SOF.

As can be seen in Figure 3, the injection timing of the engine was changing so as to increase the amount of fuel to meet the speed and load condition. Mode 3 was the only mode where the injection timing did not change, and shows that the NO\(_x\) does increase with DME addition. However, injection pressure was increasing so that the required fuel energy could be injected into the cylinder over the same crank angle timing. This may explain the increase in NO\(_x\).
Because of DME's vapor pressure, as the fuel is injected into the engine, the DME may be acting to atomize the diesel fuel into a finer spray. The blended fuel has a lower density than the diesel fuel, and the compressibility of the fuel blend has also changed. This may be reducing the premixed phase of combustion, causing more of the combustion process to be diffusion-controlled because the fuel is vaporizing and igniting so quickly. Additionally, more fuel by volume is being injected to maintain the same energy density and thus the same speed and load conditions. So, there could be some small increase in NOx emissions for this reason. In Figure 6, the brake specific energy consumption shows that the same amount of fuel on an energy basis is used for each mode, except for mode 1. Since the fuel is less dense and has higher compressibility, this may be affecting the fuel leaving the injector port and modifying the air entrainment into the fuel jet. Kajitani and coworkers data supports the increase in NOx emissions [3].

As described before, the engine was commanding more fuel, which caused the fuel injection pressure, as well as, the fuel injection timing to shift. The fuel volume required is also confirmed by the engine control signals, shown in Figure 5. From a fuel consumption standpoint, as DME content increased the volume of fuel being commanded from the engine control increased, but reached the maximum limit. This could be the result of the maximum limit, the fuel injection pressure limit and possibly the injection timing limit, within the engine control program.
Figure 5. Engine control commanded fuel volume desired

Figure 6. Fuel Consumption
Laboratory Engine Operational Challenges

During the data analysis phase, the engine was operated occasionally to prepare the fuel system for the Bus Project. There were several issues to address, with the major one being the gear swell of the fuel pump.

After discussion with the pump manufacturer (Tuthill) that was chosen, a compromise design was concluded to be the best approach based on the pressure, temperature, and flow rate envelope of the fuel system. Since the swell rate of the gears was not known, and the bus fuel systems would be designed to leave the fuel on the fuel system upon shutdown, it was critical to find a solution. In the engine lab, experience was gained with gear swell from DME/diesel blends. The initial material selection was PPS (Polyphenylene Sulfide, Carbon Fiber reinforced), which was believed to be the best with regard to the materials the manufacturer could offer. So, to accommodate the pump for the situation with the bus, it was decided to increase the tolerance in the gears so that the pump would not shut down after immersion in the fuel over an extended period of time. By doing this, the flow rate from the pump was reduced. It should be noted that the ability of the pressurized fuel system to keep the fuel in a liquid form is a function of pressure, temperature, and flow rate.

Additionally, when preparing the engine to test the modified fuel pump, starting problems were encountered. It was initially thought that this failure was due to possible early wear in the injectors. However, this issue was resolved by enabling the electronics that control the glow plugs, which ensure adequate charge temperature for ignition when the engine is started. Further work determined that the electrical system of the engine is sensitive to battery voltage. After placing a deep charge on the battery, the engine started without any problems. This was a significant step, in that after not operating the engine for several months, engine wear due to the fuel blends had not been the cause of the failure, and did not present concern for the bus project.

Laboratory Engine Evaluation for Bus Project Fuel System Design

Based on the finalized Failure Modes and Effects Analysis (FMEA) of the bus fuel system design, a number of engine tests were proposed to validate the robustness of the system. These tests included the following:

1. Repeatedly start the engine to confirm the starting problem has been corrected.
2. Test Bus Heat exchangers to confirm that the thermal capacity on the steady state lab engine test is greater than what the bus would experience. Use a thermal couple in the system to determine the time it takes for the system to heat up and stabilized.
3. Test flammable gas monitors
4. Perform Injector test with electronic service tool
5. Test Evaporative canisters for system vapor capture on shutdown.
As the bus system development and testing moved forward, decisions were made to go without the Evaporative Canister system, as a vapor test on the bus system proved this was not an issue. Also, engine lab evaluations on the heat exchangers showed that the initial system was undersized and greater exchange capability was need, including the addition of a fan. Finally, as the bus system developed and failures were experienced, the lab engine was used to test out theories of operation, in order to demonstrate a failure mode and provide corrective action for the bus system. It proved to be a valuable tool for this purpose.

At this time, utilizing the service tool to test the injectors was not possible due to several of the external components of the engine electronics not being available. The service tool performs a series of system checks before testing the injectors, and the electronic inputs necessary were not available so that one could step through in the service tool software. This test will be valuable to perform in the future injector study, so some time will be spent in determining the necessary inputs and software safety stops, so that it can be used as a diagnostic approach in the injector studies.
Viscometer Progress Report.

High Pressure Viscometer Setup
To optimize the performance of a fuel injection system for a particular fuel or fuel blend, it is very important to have a good estimate of the physical properties of those fuels. It is equally important to know the change in properties with change in pressure and temperature. An experiment was configured to measure the viscosity of diesel, DME and their blends at various pressures. The high pressure viscometer apparatus used for this work was designed and built at The Pennsylvania State University in 1962-63. This apparatus was modified to allow for charging of a pressurized liquid sample, as is necessary when dealing with compressed liquids. Robert Johnson in his Master’s thesis, gives a detailed description of the design and use of the apparatus. The thesis presents an excellent overview and a summary of the work done in the field of high pressure viscometry up to 1962. The equipment very simple in design, is nevertheless extremely accurate in viscosity measurement up to a pressure of 10,000 psig.

Description of High Pressure Viscometer
A schematic diagram of the apparatus setup is shown in Figure 7. The setup consists of a pressure intensifying system, a pressure measurement system, a constant temperature bath and the viscometer pressure vessel.
Figure 7. High Pressure Viscometer Setup
Viscosity of Diesel – DME Blends
The high pressure viscometer capillary was calibrated using a CANNON Certified Viscosity Standard N 1.0. Mass was used as the controlled variable in determining the composition of the samples.

Table 1 shows the samples used for viscosity measurement.

<table>
<thead>
<tr>
<th>Sample number</th>
<th>Percent of DME by Mass</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>2</td>
<td>74</td>
</tr>
<tr>
<td>3</td>
<td>50</td>
</tr>
<tr>
<td>4</td>
<td>26</td>
</tr>
<tr>
<td>5</td>
<td>0</td>
</tr>
</tbody>
</table>

The remaining portion of the sample was made up by Emissions Certification Diesel – Low Sulfur (ECD-LS) from Specified Fuels of Channelview, TX. Figure 8 shows the effect of ambient pressure on the viscosity of the various liquid samples. The kinematic viscosity is plotted on a logarithmic scale. R.H. Johnson notes that a plot of the logarithm of the kinematic viscosity versus the pressure results in a straight line. The slope of this line can be used to extrapolate to higher pressures with a fair amount of accuracy.
Figure 8. Pressure-Viscosity Relationship for Diesel-DME Samples @ 100 F
The line representing the 49 State Reference Diesel fuel starts from ambient pressure. For the remaining samples with DME as a constituent, the starting point was 500 psig to ensure that the samples remain in a liquid state. The pressure range, in which the viscosity measurements are made, is typical of the low pressure circuit of a diesel engine. Previous studies, done with pure DME as the fuel, state that lower injection pressures can be used for DME as the condition in the cylinder just before the firing TDC allows a very rapid vaporization of DME.

Another use of the viscosity versus pressure relations for the various blends is to choose or design a fuel injection system for the optimized blend ratio. There are two ways in which the fuel system can be designed for an engine running on a Diesel-DME blend. The first one is to examine the capability of an existing fuel systems. The deciding factor in this case will be the minimum viscosity than an existing fuel injection system can handle. The other way would be to optimize a blend ratio for a particular engine considering the exhaust emissions benefits and the energy density tradeoffs, and to use the viscosity data of this particular blend to design a fuel injection system.

Figure 9 shows the response of the kinematic viscosity to the ratio of DME in the blend. This graph shows the effect of pressure on the viscosity of a particular blend. This data was used to select the target concentration for operation of the campus shuttle bus on DME-diesel fuel blends. At 25 wt.% DME addition, the viscosity of the blend is already below the ASTM diesel fuel specification. So, 25 wt.% is the maximum DME content on which the engine should operate.

![Figure 9. Viscosity – DME Content Relationship at Different Pressures @100 F](image-url)
Fuel Station Construction And Bus Conversion

Fuel Station Construction
Initially an in depth research and design project was undertaken for a large-scale permanent DME fueling station. The fueling station consisted of two DME storage tanks, one diesel tank, and one tank for the mixed fuel. The station was designed to utilize two transfer pumps: the DME/mixed fuel pump to perform at 10 gallons per minute (GPM) and the diesel transfer pump to perform at 100GPM. Extreme attention was given to the material of the inner seals/O-rings in the DME/mixed transfer pump. The system was designed to utilized helium as the gas to provide the necessary over pressure in the DME/mixed tanks.

The final and much less expensive design of the fueling station required only that we make the refueling process more elaborate. By switching to a two step process, we eliminated the need to have a dedicated fueling station. Instead, refueling relies on the existing diesel fuel pumps for University service vehicles to fill the bus’ main tank with diesel fuel. Then, the driver backs up to an adjacent storage location for the DME tank to transfer diesel fuel to the mixture tank containing DME and diesel fuel. The final step is to add the DME to the mixture tank and provide the helium over-pressure to keep the DME in the liquid phase. Figure 10 shows a schematic diagram of the refueling site.

In conjunction with the construction of the fueling station, a “Preparedness, Prevention and Contingency” (PPC) Plan was developed and executed in collaboration with Penn State’s office of Environmental Health and Safety.
Figure 10. Schematic diagram of DME fueling station
Shuttle Bus Conversion
For the past two years a fleet vehicle on the campus of The Pennsylvania State University has been undergoing a conversion process to run on an alternative fuel mixture. This fuel, a mixture of diesel fuel and dimethyl ether, requires the use of a pressurized fuel tank and fueling system. The initial design of this system was completed in the laboratories of The Energy Institute at Penn State on an identical engine in an engine test cell. This design was then used as the basis for the system to be implemented on the fleet vehicle. The system on the fleet vehicle started with a schematic of the laboratory setup and evolved into the present day onboard system through a complicated development process which is outlined below.

Phase 1:
This phase starts with the initial system design from the laboratory setup and adapts the design to the necessary operational changes required for use on a fleet vehicle. These issues are mainly limited to fueling procedure and consideration of the interaction between the driver and the fueling system. The system must be designed so that the fueling procedure after the conversion process is relatively easy and efficient.

Phase 2:
This phase starts after a functional design has been achieved. It includes environmental considerations, material compatibility issues, and public safety issues. The conclusion of this phase is the completion of a detailed HAZOP analysis on the final system design layout and determination of any necessary changes to the design before construction begins.

This phase increased the complexity of the design by adding numerous fail safe components such as bypass valves, emergency shut off valves, pressure relief valves, fill stop valves, pressure gauges, and check valves. In addition these changes and considerations also led to the development of a complicated electrical system to monitor pressure, temperature and fuel level.

Phase 3:
This phase involved individual component selection, sizing, pricing and purchasing. Due to availability issues and material compatibility issues this phase took the better part of 6 months to complete.

Phase 4:
This phase consisted of the actual system construction on the bus. The system was constructed on the vehicle in two blocks:

Block 1: All components under the bus and in the passenger compartment are installed. No changes were made to the existing fueling system or engine compartment. This allowed the bus to be operated during times of high demand for fleet vehicles.

Block 2: The bus is removed from service and the factory fueling system was disconnected and the new fueling system connection was made.
This phase also included constant re-evaluation of design changes as needed.

Phase 5:

This phase is an actual safety inspection of the completed system on the bus. The inspection was performed by both outside and inside personnel. This phase also served to check the status of open line items in the Hazop analysis and sign off on those items if they were now a non-issue due to design changes or there were completed.

Phase 6:

This phase is the debugging process and functionality test for the system. It is divided into two sections which are in turn each divided into two blocks.

Section 1: This section involves running the bus on diesel fuel while utilizing the new fueling system. This test is separated into two blocks:

Block 1: The bus is driven at the Pennsylvania Transportation Institute test track facility located in State College Pennsylvania for a distance of no less than 50 miles. This serves to evaluate the structural integrity of the system and its functionality.

Block 2: The bus is operated on the Penn State campus while under supervision from Energy Institute staff. The purpose of which is to orient the fleet drivers and passengers to the on-board changes. This block also serves to evaluate the systems performance in the buses normal environment.

The completion of this section is a leak check of the system by both a visual and physical inspection.

Section 2: This section is similar in structure to section 1 with the addition of the DiMethyl ether into the system. Again the section is separated into two blocks:

Block 1: The bus is driven at the Pennsylvania Transportation Institute test track facility located in State College Pennsylvania for a distance of no less than 50 miles. This serves to evaluate the structural integrity of the system and its functionality.

Block 2: The bus is operated on the Penn State campus while under supervision from Energy Institute staff. The purpose of which is to orient the fleet drivers and passengers to the on-board changes. This block also serves to evaluate the systems performance in the buses normal environment.

The completion of this section and this phase is a final leak check of the system by both a visual and physical inspection.
Figure 11. Schematic diagram of DME-diesel fueling system on the shuttle bus
Figures 12 and 13 show digital photos of the shuttle bus after the conversion. These pictures were taken at the event hosted on April 19, 2002 to celebrate the launch of the bus on DME-diesel blends.

Figure 12. (a) Jenny Tennant and John Winslow of NETL and (b) Jim Sorensen, Jo Ann Franks, Bob Miller and Barry Halper of Air Products and Chemicals, Inc.

Figure 13. DME-Diesel Fueled Campus Shuttle Bus pulling away during its maiden voyage at the event to celebrate the launch of the bus, April 19, 2002 held at the Penn State Energy Institute.
Finalization of the Converted Fueling System

The standard configuration of the T444E engine is for fuel to enter from the backside of the cylinder head (side closest to the firewall) and remain in the fuel rail until needed by the injectors. This configuration, referred to as a “dead heading,” causes the fuel to be heated to engine coolant temperatures. When the engine is fueled on diesel fuel this is not a problem. However, when running on a blend of DME and diesel fuel, this heat soak becomes an issue. To combat the heating of the fuel in the rail, the fuel flow path is changed from a “dead heading” arrangement to a circulation loop. Fuel enters the engine at the same place as in the original configuration and exits at the front of the engine. The fuel from each side of the engine is then combined and routed through four transmission oil coolers (referred to as “fuel coolers”) that are mounted just in front of the factory radiator and charge air cooler. This configuration allows the fuel to be cooled and fresh makeup fuel added to the loop before the mixture is re-circulated to the engine. Figure 14 shows a photograph of the final configuration of the fuel handling system installed under the hood of the shuttle bus.

![Fuel Pump](image)

![Fuel Filter](image)

![Back Pressure Regulator](image)

**Figure 14. Photograph of the fuel handling system on the converted shuttle bus**

Test Firing

Once construction of the fuel handling system was complete, the vehicle was test fired on diesel fuel. This allowed for a full operational check of the new system and a fully pressurized leak check. The vehicle was then operated at the Pennsylvania Transportation Institute Test Track facility to verify vehicle performance, operation of the vehicle with the converted fueling system and refueling procedures. The vehicle was fueled on a conservative mixture of 10 vol.% DME in diesel fuel for initial testing after operation problems had been addressed.
Operational Issues

With the complexity of the design, several operational issues became evident once construction was completed. These problems involved formation of two-phase flow in the fuel handling system that caused rough engine operation and re-fueling difficulties.

Circulation Pump

A magnetically coupled gear pump is used to circulate fuel through the cylinder heads through fuel coolers and back to the engine. Since DME is a vapor at room temperature and pressure, the fuel must be kept pressurized to avoid formation of two-phase flow within the system. When the fuel is exposed to engine temperatures, the amount of pressure needed to keep the DME in the liquid phase increases. The vapor pressure of DME as a function of temperature is shown in Figure 15 for reference, provided by Dupont Technical Bulletin for Dymel A [6]. Since the system pressure is set, the fuel temperature must be lowered via heat exchange through the fuel coolers. Thus, if for some reason the circulation pump does not work, fuel is not moved through the head to provide sufficient cooling and the resulting two-phase flow causes rough engine operation.

![Graph showing the vapor pressure of DME as a function of temperature.](Image)

**Figure 15.** Vapor pressure of DME as a function of temperature [6].

Several scenarios for pump failure were encountered. The pump, which runs off 110 AC power provided by a DC-to-AC inverter, was demanding increasing amounts of power due to: inadequate power supply, inadequate flow of fluid through the system due to blockage, jamming of the gears due to debris, thermal shutoff, and swelling of the gears.
Initially, after construction was complete, the bus was fueled on diesel fuel. During this time no problems with the pump were noticed. However, after long periods of operation, the engine ran roughly and the amount of blow-by increased. When the bus was fueled on DME, the current draw by the electric motor that drives the circulation pump gradually increased over time. The current demand by the pump motor would increase to the maximum rated current for the motor and trip an internal protection circuit in the motor. Hence, the pump would shut off.

The manufacturer was contacted and a larger motor and a motor adapter were purchased and installed. After installation the operation of the pump seemed to become worse and the motor and motor adapter were removed. It was found that the magnetic coupler from the motor shaft to the cap of the pump was destroyed, as can be seen in Figure 16. The pump, motor mate, and coupler were taken to the attention of the regional sales representative who replaced the destroyed coupler and correctly installed a new one. The pump was then reincorporated into the system.

![Figure 16. Damaged magnetic coupling from Tuthill gear pump.](image)

With the motor now working correctly, it became evident that a larger power source was required. Since the pump runs on 110 AC but only 12V DC is available on the vehicle an inverter is used. To obtain the size inverter needed a recreational vehicle dealer was contacted. Once the inverter was in place the pump performed flawlessly on diesel fuel for 8 days. When the vehicle was switched to DME-diesel blends, problems again became evident.

A flow meter was installed to determine if the pump was getting adequate fluid or if it was running dry. The meter was installed in the system with as little changes as necessary to ensure a true reading. The flow was found to be in the acceptable range for the motor rating. This ruled out insufficient flow due to blockage in the system. It was postulated that the flow restriction could be on the outlet side of the pump so the filter element was examined.
The filter was found to have disintegrated at the top and bottom of the element where rubber caps were used in the manufacturing. A new filter element with a completely metal housing and a more resilient rubber was chosen and installed. The pump was then removed and the gears checked for any debris that could be preventing their rotation. Very little material was found in the gears and the pump was reassembled and reinstalled.

By the manufacturer's suggestion, a surface temperature thermal coupler was installed on the skin of the pump. It was found that there was inadequate airflow to the motor of the pump so a 12V DC accessory fan was installed to direct air toward the pump. After the installation of the fan, the skin temperature was found to be in the acceptable range, thus ruling out thermal shutoff. With all of these test and modifications, problems were still evident after fueling with DME.

It was thus apparent that gear swell due to prolonged exposure to DME was the source of the problem. The manufacturer was again contacted and a new pump was obtained. Pump shutoff and rough engine operation ceased to be a problem from this point forward.

Fuel Level Gauge

A float gauge inside the tank monitors the level of fuel in the mix tank. The signal is then sent to two individual readouts. One readout is mounted above the driver's seat and the other is attached to a five-pin connector on the side of the bus during fueling.

Vehicle Emissions Testing

Emission testing is done by use of on board analysis equipment owned and operated by Clean Air Inc. The equipment consists of two units, which are approx. 24 inches wide by 22 inches deep, and several laptop computers that provide interface and control ability. The equipment measures CO, CO2, NOx, particulate matter, as a function of vehicle speed and load. Fuel consumption is calculated from the CO and CO2 measurements.

Driving Cycles

For the purpose of ensuring and verifying repeatability, a specific testing procedure was adhered to for all testing runs. All testing was performed at The Pennsylvania State Transportation Institute Test Track facility on a one-mile oval track. Four different driving cycles were used for the testing: P20, P40, Orange County, and Manhattan Cycles.

A P20 consists of starting at a specific point on the track and accelerating to 20 mph. The vehicle is held at 20 mph until the first stopping point is reached. The vehicle is then brought to a stop for seven seconds and the driver then accelerates back to 20 mph. There are 8 stops performed in one loop. A complete P20 consists of 2 loops.

A P40 consists of starting again at a specific point on the track and accelerating to 40 mph. The vehicle's speed is then held at 40 mph until the first stopping point is reached. Then vehicle is then brought to a stop for seven seconds and the driver accelerates back to 40 mph. The first
stop is at half-track the second is at the original starting point. A complete P40 consists of 2 loops.

An Orange County cycle is a reproduction of a typical bus cycle in Orange County California. The cycle is displayed on a laptop computer running Driver’s Aid software® provided by Clean Air Technologies International, Inc. The software also displays the vehicle’s speed and the driver matches the speed trace to the prescribed cycle being displayed. The Orange County cycle is broken into two parts to minimize data file length and can be seen in Figures 17 and 18. A complete Orange County set consists of four complete cycles.

A Manhattan cycle is a reproduction of a typical bus cycle in Manhattan, New York. Again the cycle is displayed by the Driver’s Aid ® software and the driver matches the vehicle speed trace to the prescribed cycle. A complete Manhattan set consists of four cycles. The Manhattan cycle is shown in Figure 19.

![Graph showing speed over time](image-url)

**Figure 17.** Part One of the Orange County Cycle – Driving Cycle to Represent Transit Bus Operation in Orange County, CA.
Figure 18. Part Two of the Orange County Cycle.

Figure 19. Manhattan Cycle – Driving Cycle to Represent Transit Bus Operation in Manhattan, NY.
Vehicle Testing Procedures

Since the Driver’s Aid® software display is difficult to see during the day the Orange County and Manhattan cycles are run at night and the P20 and P40 cycles are run during the day.

A typical day of testing begins with warming up the equipment. This process takes about one hour to complete during which the vehicle is at idle. At the completion of this time period, several test laps are made to check the equipment status. If redundant instruments indicate conflicting measurements the laps are aborted and the equipment checked. Once consistent results are obtained, actual testing begins.

Using the onboard equipment as a live monitoring system, three laps at 30 mph are made to burn off particulates that may have accumulated during the warm-up time. If particulate levels continue to trend downward in the last lap, additional laps are done until stable readings are seen.

At the completion of these “burn off” laps, the vehicle is brought to a stop at the P20 starting point. A period of 90 seconds is allowed to transpire allowing for the equipment to be reset. A complete P20 is then run followed by a complete P40, this pattern is then continued alternating back and forth with 90 seconds between each complete set.

An identical pattern is followed for the Orange County and Manhattan cycles. After equipment warm up “burn off” laps are run pausing 90 seconds then beginning the desired cycle. The testing pattern consists of running blocks of Manhattan cycles before switching to Orange County cycles.

Vehicle Testing Results

Testing was performed for 3 different fuel mixtures. The first test served as a baseline run and the vehicle was fueled on pure diesel. The second test was a fuel mixture of 14% by volume DME and diesel; the third test was 25% by volume DME and diesel. While emissions data were monitored for the P20, P40, Orange County and Manhattan test cycles, only data from the Manhattan cycle is presented here.

Total particulate results for all three-fuel compositions for the Manhattan cycle can be seen in Figure 20. As is expected with the addition of an oxygenate as the amount of DME in the fuel mixture increases the measured particulate level drops. For 14 vol.% DME, the PM emissions drop by 60%. For 25 vol.% DME, the PM emissions drop by 80%.
Figure 20. Particulate Matter Emissions During the Manhattan Cycle (cumulative mass emissions per test cycle) for 0, 14 and 25 vol.% DME in Diesel Fuel.

Similar results to those seen by Chapman et al. [1] were seen for hydrocarbons, NO\textsubscript{x}, and CO. For a Manhattan cycle, the emissions of hydrocarbons, NO\textsubscript{x}, and CO are shown in Figures 21, 22, and 23, respectively.

Figure 21. Hydrocarbon Emissions During the Manhattan Cycle (cumulative mass emissions per test cycle) for 0, 14 and 25 vol.% DME in Diesel Fuel.
Figure 22. **NOx Emissions During the Manhattan Cycle (cumulative mass emissions per test cycle) for 0, 14 and 25 vol.% DME in Diesel Fuel.**

Figure 23. **Carbon Monoxide Emissions During the Manhattan Cycle (cumulative mass emissions per test cycle) for 0, 14 and 25 vol.% DME in Diesel Fuel.**

While the emissions of particulate were substantially reduced, emissions of hydrocarbons, NOx, and CO were significantly increased with DME addition. While these trends were also observed in the laboratory engine, they point to a need to optimize the injection characteristics of the fuel blend and to the need to add an oxidation catalyst to a vehicle fueled in this manner. A similar observation was made in the Volvo bus demonstration, that addition of an oxidation catalyst to a DME-fueled vehicle provided very low emissions [7].
Technical Issues With The Utilization Of DME As A Fuel

Much of the early work with DME utilization has been done at the AVL labs in Graz, Austria. The testing showed that DME reduces particulate emissions to zero, and also showed that the typical diesel fuel injection system does not tolerate DME [8]. The present work shows that with DME-diesel blends, substantial particulate emissions reduction can be obtained while preserving the integrity of the fuel injection system. Figure 24 shows a picture of the DME-Fueled Shuttle bus in operation at Penn State.

![DME-Fueled Shuttle Bus Operating on the Faculty/Staff Loop at the University Park campus of the Pennsylvania State University, August 2002.](image)

Overall, this demonstration project has shown that a vehicle can be successfully operated on DME, but at restricted DME blend levels below 25 wt.% (30 vol.%). There are remaining concerns and challenges, particularly because of the low viscosity of DME. Below is a combined list of concerns not only from the present work, but also from other groups who are now working through the fuel property, design and technology issues for utilization of DME:

- DME was leaking past clearances on the injectors and seals. This caused the need for the camshaft housing and crankcase of the engine to be vented [8].

- At high vapor pressure, the DME was cavitating, which caused difficulties in maintaining stable fuel injection [8].

- While DME is more compressible than diesel fuel, it was found that the compressibility changed with temperature and pressure. Therefore, this made it difficult to inject the
maximum fuel quantity at high temperatures and during full load operation using traditional diesel equipment [8].

- DME chemically attacked some seals [8].
- Not much effort has been put towards understanding the environmental impacts of the compound itself or the emissions from the fuel combustion, as compared to other fuels [8].
- A larger fuel tank will be required, as compared to diesel fuel, because of the lower density and heating value of DME [9].
- Since the vapor pressure of DME is low, the fuel vaporizes immediately upon injection into the cylinder. This may or may not be an issue, but further study may confirm how the combustion reaction takes place after the vaporization occurs [9].
- Injection via some fuel pumps causes uncontrollable pressure waves in the entire system [10]
- Predictability of spray behavior and characteristics is important in repeatability of combustion [11].
- Turbulence within the cylinder is important for mixing of the fuel, which in turn reduces emissions [12].
CONCLUSIONS

This work has demonstrated that a conventional diesel vehicle can be converted to operate on blends of DME and diesel fuel. Significant reductions in particulate emissions were observed when the converted shuttle bus was operated on DME – diesel blends, while there were increases in unburned hydrocarbons, NOx and CO. By blending DME with diesel fuel, an acceptable viscosity and lubricity could be obtained in the fuel mixture to provide reliable operation of the fuel injection system. Nonetheless, operation of the vehicle was not without challenges during the conversion and debugging phases, as is documented in this paper.

DME holds great promise as a fuel for the future. But, many technical challenges remain to be overcome before neat DME-fueled vehicles can be a commercially viable option. Among them are the low viscosity, low lubricity and material incompatibilities.
REFERENCES


ACKNOWLEDGMENTS

This project has benefited substantially from collaboration and support from a variety of sources, in addition to the project sponsors. The project team at Penn State is deeply indebted to Air Products and Chemicals, Inc., especially Barry Halper, Peter J.A. Tijm (formerly of APCI), Jim Sorensen, Jo Ann Franks, Ed Heydorn, Fran Waller, and Bernie Toseland. The project team also gratefully acknowledges the support from the National Energy Technology Laboratory, particularly the guidance and encouragement received from Mike Nowak, Jenny Tennant, Shelby Rodgers and John Winslow. The team is grateful to Susan Summers and the Alternative Fuel Incentive Grant Program in the Pennsylvania Department of Environmental Protection for their support of the project.

At Penn State University, the project team received assistance, guidance and patient support from both Fleet Operations and Garage Services. Bruce Younkin and the staff of Fleet Operations were a great help in making the bus available to us for the conversion, and working with us during the debugging and deployment phases of the project. Sam Entz and his staff in Garage Services provided excellent support and assistance during the conversion and debugging phases of the project. Without Bruce and Sam’s collaboration, the shuttle bus demonstration could not have succeeded.

A variety of companies provided support and hardware to the project, particularly during the shuttle bus conversion. Chief among the industrial sponsors has been Navistar (now International Truck) and Dr. Pranab Das, who donated funds for the laboratory engine for this project. In addition, the project received support and technical guidance from: Allegheny Truck, Caterpillar, Clean Air Technologies, Columbia Propane, DuPont Fluorochemicals, DuPont-Dow Elastomers, F&L Fluid Components, Manchester Tank, Master Flow Pumps, National Instruments, Sierra Monitor Corporation, Parker - Racor Filtration, and Tuthill Pump.

Finally, from within the College of Earth and Mineral Sciences there has been tremendous support for this project. This support has come in the form of matching funds from The Energy Institute (Director, Dr. Harold Schobert), and student stipend support from the Energy Institute and the Department of Energy & Geo-Environmental Engineering (department head, Dr. Alan Scaroni). Within this support, the engine test facility and the addition of a third student to the project would not have been possible. Also, the team benefited from the involvement of Dr. Louis I. Boehman, while he served as a Visiting Scientist within the Energy Institute.

The original Principal Investigator for this project was Prof. Don Streit, who passed away tragically in August 2001. He played a critical role in getting this project underway and we are forever grateful for his guidance.

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This material was prepared with the support of the Pennsylvania Department of Environmental Protection. Any opinions, findings, conclusions, or recommendations expressed herein are those of the author(s) and do not necessarily reflect the views of the DEP.
APPENDICES

The appendices include papers and theses written during the course of this project. These documents are included on the CD that accompanies this document.

Society of Automotive Engineers Technical Paper No. 2000-01-2887
ACS Preprint from 2001 ACS National Meeting
Society of Automotive Engineers Technical Paper No. 2001-01-3626
ACS Preprint from 2002 ACS National Meeting
Society of Automotive Engineers Technical Paper No. 2003-01-0756
MS Thesis in Mechanical Engineering, S. V. Bhide, 2003
Emission Characteristics of a Navistar 7.3L Turbodiesel Fueled with Blends of Oxygenates and Diesel

Elana M. Chapman, Shirish V. Bhide, and André L. Boehman
The Pennsylvania State University

Peter J.A. Tijm and Francis J. Waller
Air Products and Chemicals, Inc.

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ABSTRACT

Several oxygenates have been proposed and tested for use with or as diesel fuel. This paper examines two such oxygenates, CETANER™ and dimethyl ether (DME), partially or wholly produced by Air Products and Chemicals, Inc.’s Liquid Phase Technology. In previous studies on a single cylinder compression ignition engine and a Volkswagen TDI four cylinder engine, significant reductions in particulate matter emissions were observed with blends of CETANER™ in diesel fuel. In this study, experiments were performed on a multi-cylinder Navistar 7.3L Turbodiesel engine confirmed and extended the observations from the earlier studies. This is an important step in not only showing that the fuel does perform on each type of engine in similar fashion, but also in showing that DME and its derivatives can give consistent, significant results in lowering emissions. The oxygenated fuels were blended to achieve a net addition of 2 wt.% oxygen in the blended fuel. A pressurized fueling system was developed to deliver mixtures of DME-diesel at up to 1 MPa (150 psi). With the DME-diesel blend, less consistent emissions results were obtained owing to an inability to sufficiently fuel in the rail.

INTRODUCTION

Demand for cleaner burning diesel fuels is growing worldwide, as governmental regulations make emissions reductions necessary. In the U.S., future regulations that take effect in 2004 and 2007 will require diesel engine and vehicle manufacturers to review all aspects of the vehicle system design. To achieve substantial reductions in emissions, it is thought that reformulated diesel fuels will play an important role. The reformulation of diesel fuels could include lowering the sulfur content, lowering the aromatic content, or potentially the addition of oxygen within the fuel.

A solution to affect emissions reductions for future and current diesel vehicles on the road is to modify the fuel without the need for modifying the engine hardware. It has been shown that many oxygenates are effective at reducing particulate emissions from diesel engines [1-20]. Therefore, much research has focused on screening of oxygenated fuel additives, including alcohols, esters, and ethers. Of particular interest are the glycol ethers, which have been shown to be very effective as blends and as neat fuel.

Over the last ten years, many researchers have begun to evaluate the performance of blends of glycol ethers with diesel fuel, and have observed decreases in particulate matter emissions. Liotta and Montalvo [5] measured the effects of several different oxygenated fuel additives, including several glycol ethers. From the tests performed in a DDC Series 60 diesel engine, their results indicated that particulate matter reductions of 4-10% could be achieved for each 1% of oxygen blended into diesel fuel, through incorporation as a glycol ether. Specifically, the results indicated that oxygen addition via glycol ether addition was more effective than oxygen addition via alcohol. Ullman and coworkers [6,7] also evaluated the addition of several glycol ethers to diesel fuel, specifically, monoglyme (1,2-dimethoxyethane) and diglyme (diethylene glycol dimethyl ether), at 2 wt.% and 4 wt.% oxygen. A DDC Series 60 engine and a Navistar DTA-466 engine were used for the testing. Their results indicated that particulate matter reductions of 6-7% were reached for each 1% of oxygen blended into diesel fuel.

Higher molecular weight glycol ethers blended with diesel fuel have also been effectively used to reduce particulate matter emissions. Tsurutani and coworkers [8] blended several glymes, including monoglyme, diglyme, triglyme and tetraglyme, at levels up to 12 wt.% oxygen in diesel fuel. They observed that
combustion of the glycol ether blends in an IDI engine yielded a particulate matter emission reduction of 3-5% for each percent of oxygen blended. Additional studies completed by Hess et al. [9] as well as by Litzinger and coworkers [10,11] have shown that higher molecular weight glycol ethers are also effective in reducing particulate matter emissions, although to a lesser extent than monoglyme or diglyme.

Beatrice and coworkers [12-15] as well as Miyamoto and coworkers [16,17] have evaluated the use of diglyme as both an oxygenated blend component and as a neat fuel. Both research groups have reported that smokeless combustion is possible with pure diglyme. Heat release rates have shown that combustion of diglyme results in shorter combustion duration, with a shift towards the diffusion phase of the combustion process.

Although it has been shown that glycol ethers effectively reduce particulate emissions, the fundamental mechanisms of the reduction have not been clearly identified. There has been some work in simulating the ignition and rate mechanism behavior of dimethyl ether in comparison to dimethoxy methane [18]. Also, oxidation mechanisms have been proposed for gaseous forms of DME [21-23]. Limited data is available for many of the liquid oxygenates under consideration.

Hence, the objectives of the experimental work reported here are to further evaluate the effects of glycol ethers, specifically CETANERTM on the diesel combustion process, as well as to compare this data to another oxygenate, dimethyl ether (DME), utilizing the same engine condition. In different engine configurations, CETANERTM has been shown to reduce particulate emissions over a range of blend ratios in diesel fuel [24]. CETANERTM is an oxygenated diesel fuel additive developed as a coal-derived syngas product by Air Products and Chemicals, Inc., and is a mix of glycol ethers, namely monoglyme and diglyme. As a diesel fuel additive, CETANERTM has been shown to exhibit high cetane number, roughly 125 [25]. This work is also intended to demonstrate fueling of a commercial turbodiesel engine on DME-diesel blends through the use of a pressurized fueling system. The long term objective is to apply this fueling strategy to a shuttle bus on the Penn State University Park campus.

For this experimental work, blends of a Federal Certification diesel fuel with CETANERTM and dimethyl ether were evaluated in a multi-cylinder direct injection (DI) engine. A simplified mixture of 20% monoglyme and 80% diglyme was chosen for the experimental work, to represent a potential CETANERTM formulation. In-cylinder pressure measurements provided information about the impact of the oxygenated fuel on the combustion process. In addition, fuel property tests were performed on the base fuels as well as the fuel blends. These measurements included calorific value, flash point, and viscosity, and were used to understand and describe the combustion behavior. Since dimethyl ether is a vapor at 1 atm., the fuel property tests for the fuel blends were performed under pressure so that the dimethyl ether and diesel remained in the liquid state. Therefore, the fuel system of the engine was redesigned to accommodate the pressurized fuel delivery.

EXPERIMENTAL

TEST ENGINE

For the purpose of studying the effects of fuel additives on light-medium duty diesel combustion, a Navistar T444E 7.3L Turbodiesel engine was coupled to a 450 horsepower Eaton (Model AD-1802) eddy-current dynamometer for testing. The specifications for the engine are given in Table 1. Pentium PCs with Kiethly Metrabyte DAS-1800 data acquisition cards were connected to the engine to log real-time engine parameters. These parameters included engine speed, torque, and power from the engine. A Modicon PLC was used to record temperatures from the engine, as well as, for the entire experimental system monitoring. Intake airflow rates were determined via an electronic flow sensor, which was calibrated from a laminar flow element. Fuel consumption was monitored using a precision Sartorius Industrial Scale (Model EA60EDE-1), with an accuracy of ±2 grams. Figure 1 shows the test cell set up, and additional equipment used for emissions monitoring.

![Figure 1. Multicylinder Engine Test Cell Navistar T444E 7.3L Turbodiesel](image)

TEST PROCEDURE

In this work, an AVL 8-mode test procedure has been utilized as a model for diesel emissions tests. The AVL 8-mode tests was designed to correlate to the U.S. Federal Heavy- Duty Transient Test procedure through a weighted 8- mode steady state test procedure. The 8 modes are a combination of speeds and loads, to produce the same emissions output as would be recorded for a transient cycle [26]. For our engine, the
test procedure included the speed and load settings, shown in Table 2.

Table 1. Characteristics of the 1998 Navistar T444E 7.3L Turbodiesel Engine

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Displacement</td>
<td>7.3 Liter (444 cu.in.)</td>
</tr>
<tr>
<td>Bore</td>
<td>104.39mm (4.11 inch)</td>
</tr>
<tr>
<td>Stroke</td>
<td>106.20mm (4.18 inch)</td>
</tr>
<tr>
<td>Rated Power</td>
<td>143kW (190 HP)@2300 RPM</td>
</tr>
<tr>
<td>Peak Torque</td>
<td>640Nm(485lbf-ft)@1500 RPM</td>
</tr>
<tr>
<td>Configuration</td>
<td>Turbo charged, Intercooled (Air-to-Air), Direct Injection</td>
</tr>
<tr>
<td>Injection Scheme</td>
<td>HEUI- Hydraulically actuated, electronically controlled unit injectors</td>
</tr>
<tr>
<td>Low Idle Speed</td>
<td>700 RPM</td>
</tr>
<tr>
<td>Features</td>
<td>Split- shot injection</td>
</tr>
<tr>
<td>Compression Ratio</td>
<td>17.5:1</td>
</tr>
</tbody>
</table>

Table 2. AVL 8-Mode Test for the Navistar T444E Turbodiesel engine

<table>
<thead>
<tr>
<th>Mode</th>
<th>Speed (rpm)</th>
<th>Load (Nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>700</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>876</td>
<td>111</td>
</tr>
<tr>
<td>3</td>
<td>1036</td>
<td>296</td>
</tr>
<tr>
<td>4</td>
<td>1212</td>
<td>472</td>
</tr>
<tr>
<td>5</td>
<td>2300</td>
<td>102</td>
</tr>
<tr>
<td>6</td>
<td>2220</td>
<td>235</td>
</tr>
<tr>
<td>7</td>
<td>2220</td>
<td>405</td>
</tr>
<tr>
<td>8</td>
<td>2124</td>
<td>540</td>
</tr>
</tbody>
</table>

EMISSIONS ANALYSIS

An extended warm-up period was used to prepare the engine for testing. The sampling and measurements during each mode commenced when the exhaust temperatures reached a steady state. During this time, RPM and torque were maintained within 1-2% of the target test conditions. Once steady-state operation was achieved, a portion of the exhaust gas was passed through a Sierra Instruments BG-1 micro-dilution test stand with a constant dilution air / sample flow ratio of 8:1 and a total flow of 150 liters/min. These settings were chosen in order to maintain the filter temperature below the EPA specification of 52°C. Particulate collection occurred on Pallflex 90mm filters (Type EMFAB TX40HII20-WW), conditioned in an environmental chamber at 25°C and 45% relative humidity before and after sampling. Five particulate samples were taken for each fuel at each test mode, except for the DME-diesel tests where only one sample was obtained.

Exhaust gas analyses were completed using a Nicolet Magna 550 Fourier Transform Infrared (FTIR) Spectrometer. For each mode, five gas samples were analyzed for CO₂, CO, NO and NO₂. Also, a Rosemont Analytical O₂ sensor was used to monitor the percent oxygen in the exhaust gas. The oxygen readings were used in conjunction with the mass flow sensor to determine and verify the air / fuel ratio. Additionally, total hydrocarbon emissions were monitored using a California Analytical Instruments Model 300 HFID Heated Total Hydrocarbon Gas Analyzer. For the total hydrocarbon measurements, undiluted exhaust gas was collected via a heated sample line, which was maintained to 190°C. Calibration of all equipment was completed prior to each day of testing. Gaseous emissions data are only presented for the CETANER™ diesel blend, as the gaseous sampling system was not functioning correctly during the testing of the DME-diesel blends.

PRESSURE TRACE ANALYSIS

In order to observe the impact of the oxygenated blends on combustion and heat release, the combustion chamber of cylinder 1 of the engine was fitted with a Kistler 6125A pressure probe. The pressure sensor was used with a Kistler 2612 optical crank angle encoder to provide time resolved in-cylinder pressure traces of the combustion event. Pressure, crank angle, and TDC trigger signals were acquired with a Kiethley DAS-1800 data acquisition card operating in a “burst “ mode. The pressure traces were analyzed with PTrAn V.02, a software product designed by Optimum Power.

TEST FUELS

Previous work has been completed comparing the increasing percentage of oxygenate mixed with diesel fuels within several types of engines [8-17,24]. For this testing, comparisons are made between a 2 wt.% addition oxygen of two different additives. The baseline diesel fuel properties, as well as test fuel properties are given below in Table 3. A Federal Certification Fuel (Specified Fuels, Emissions Certification Diesel – Low Sulfur, ECD-LS) was used in these experiments. Because of the difficulty of obtaining the fuel blend properties for DME as a liquid, the properties available in the literature for neat DME are presented.
PRESSURIZED FUEL DELIVERY SYSTEM FOR 
DIESEL-DME BLENDS

DME is a liquefied gas. At STP, it is a gas, but liquefies under a moderate pressure. The fuel delivery system was designed keeping in mind the following important points:

- The vapor pressure of DME.
- Material compatibility of the various components in the fuel system with DME
- Lack of lubricity of pure DME.

A schematic of the fuel delivery system is shown in Figure 2. The working of the fuel delivery system can be explained as follows:

1. The fuel comes out of the fuel tank at a pressure of about 0.6 MPag (90 psig). This overpressure is necessary to keep the DME in a liquid state. Any inert gas is suitable for this purpose. Helium was used as it has a lower solubility in DME than nitrogen.
2. The pressure is then boosted by a gear pump to about 0.82 or 1.0 MPag, depending on the pressure rating of the fuel rail. The pressure in the rail is maintained at 0.48 MPag in the original fuel system of the engine.
3. The fuel return line pressure is held at about 0.82 MPag (120 psig) by the backpressure regulator. The regulator is a simple spring loaded valve that regulates the flow to keep the backpressure at 0.82 MPag (120 psig).
4. This fuel then passes through a heat exchanger, where it is cooled down to a predetermined level.
5. After cooling, the fuel is then fed to the inlet of the pump.

### Table 3. Fuel Properties

<table>
<thead>
<tr>
<th>Fuel Property</th>
<th>ASTM Method</th>
<th>ASTM Spec.</th>
<th>Base Diesel</th>
<th>5wt.% CETANER</th>
<th>DME</th>
</tr>
</thead>
<tbody>
<tr>
<td>Viscosity, 40°C, cSt</td>
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<td>1.39-4.20</td>
<td>2.2</td>
<td>1.27</td>
<td>.25</td>
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<tr>
<td>API Gravity</td>
<td>D 287</td>
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<tr>
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<tr>
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<td>D2015</td>
<td>46</td>
<td>45</td>
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<td>28</td>
</tr>
</tbody>
</table>

PRESSURE AND FLOW REQUIREMENTS

At 20°C the vapor pressure of DME is about 0.52 MPa (75 psia). Keeping DME in a liquefied state calls for pressurizing the entire fuel system from the fuel tank up to the fuel injectors. The vapor pressure also changes rapidly with temperature. The pressure of the fuel system is hence dictated by the fuel temperature. The pressure, however, is limited by the pressure rating of the fuel rail. The engine used in the study has a common rail injection system. Each cylinder head has a fuel rail running along its length, which is the source of fuel for the pressure intensifier in the fuel injectors. In the original fuel system of the engine, the pressure in the rail is maintained at 0.48 MPa (70 psig). This facilitates proper filling of the pressure intensifiers. The fuel rails in the cylinder head form a dead head system. This means that there is no fuel return once the fuel enters the fuel rail. It is because of this that the fuel temperature in the rail approaches the engine coolant temperature in the head. This layout of the fuel system was modified to accommodate a fuel return from the cylinder heads.

A study was performed in which the temperature of the fuel in the fuel rail was recorded in conjunction with the fuel consumption of the engine, for the 8 modes of the AVL test. Assuming certain values for heat capacities for diesel and DME, a minimum flow rate value was calculated so as to keep the temperature of the fuel in the rail below 50°C. The vapor pressure of DME at this temperature is about 1.0 MPa (150 psi). This pressure, more or less, dictated the temperature rise allowed. The fuel delivery pump was sized based on the above calculations. In addition to excess flow rate, cooling of the returned fuel was necessary to maintain the required fuel temperature. In these tests, a 500 W chiller was used to chill a bath through which the fuel was passed within stainless steel coils. This fuel cooling was insufficient to maintain the fuel temperature below 50°C under some operating conditions, particularly during Mode 8.

Design Considerations for the Pressurized Fuel System

DME is known to be incompatible with the common gasket materials such as Viton and buna-N, used in diesel service. Data provided by DuPont Inc. indicated Kalrez to be the best material for DME. For economic considerations however, this material was used sparingly. Other materials such as butyl rubber, Teflon and neoprene have also been found to be compatible, though not to the same degree as Kalrez. Stainless steel was used for the fuel lines as a safeguard against corrosion. All the other components such as valves and regulators were also made of stainless steel.

Selecting a pump for pumping DME was challenging due to the properties of DME such as its low lubricity and low viscosity. Due to the vapor pressure of DME, the pump...
housing was required to handle pressures up to 1.7 MPag (250 psig). Positive displacement pumps such as vane pumps, diaphragm pumps and gear pumps were considered. Gear pumps were found to be economical as well as convenient to operate. With these considerations, a gear pump made by Tuthill Pump CO, California (model #TXS2.6PPPT3WN00000) was selected. This pump has a magnetically coupled AC motor. This configuration does not have the driveshaft going through the pump housing, which in turn obviates the need for seals, a potential source of leakage. The gear material is Ryton (Polyphenylene sulphide), which was found to be compatible with diesel and DME as per the data by provided DuPont Inc. The pump body seals are made of Teflon.

The fuel filters on the engine could not be used because of the high pressure of the fuel. The minimum pressure in the fuel lines was 0.62 MPag (90 psig). This required the use of special filters, which would withstand higher pressure. A diesel water separator was used as a primary filter. This is rated at 0.69 MPag (100 psig). The final filter was a LPG filter rated at 3.4 MPag (500psig). The mesh size of the filter was 2 micron, very near to the engine specification.

The fuel tank was made out of a modified 45 kg (100 lb) capacity LPG cylinder which was pressure tested prior to use. This tank was fitted with a 1/2" NPT fitting at the bottom for liquid exchange.

RESULTS AND DISCUSSION

This section presents results from combustion studies of the effect of the CETANER™ additive on emissions compared to the base fuel composition (prior to the fuel system conversion), and the effect of the DME-diesel mixture on emission compared to the base fuel composition (after the fuel system conversion). Fuel property tests were completed to permit comparisons of the combustion data. Through an uncertainty analysis, based on methods described by Moffat, error bars showing the 95% confidence intervals are presented in each figure [27].

PARTICULATES

As noted previously in the discussion, oxygenates traditionally reduce particulate emissions. Shown in Figure 3, the brake specific particulate matter (BSPM) emissions for the 2 wt. % oxygen (5.59wt.% CETANER™ and 5.75wt.% DME) show a decrease in particulate matter emissions for CETANER™ addition, but a mixed result for DME addition. As seen in Figure 4, on a basis of particulate emissions per unit fuel consumed, greater variability in the results is evident. These numbers for the impact of CETANER™ addition correlate well with the particulate emissions observed previously by Hess and coworkers [24], as well as, Ullman and coworkers [6,7]. In their work, particulate emissions decreased as a function of increasing load. This is also seen in the results in Figure 3. For mode 4, even though there is a particulate reduction through using CETANER™ vs. the baseline diesel, the interesting trend is in the engine particulate output for the engine conditions. Mode 4 operates the engine near the peak torque, and the overall emissions conditions change, possibly due to a change in the injection timing.
is due to a reduction in the soot portion of the emission, and would result in a percentage increase in the soluble organic faction (SOF) portion [5].

It is evident from Figure 4 (in particular, Mode 4), there is substantially greater variability introduced by the pressurized fueling system with regard to particulate emissions measurements. The variation from the original baseline emissions (before the fuel system conversion) is quite large, and may in part be due to difficulties in maintaining sufficiently low temperatures in the fuel rail. As shown in Figure 2, the pressurized fuel system relies on a heat exchanger to cool the fuel that is rejected from the rail so that this fuel can be recirculated to the rail. However, during these tests it was observed that the heat exchanger capacity was insufficient to maintain fuel rail temperatures below 50°C under some of the operating modes, particularly Mode 8.

Figure 6 presents the particulate matter vs. NOx tradeoff per mode for the CETANER™-diesel blend. As can be seen for modes 2 though 8 on a brake specific basis, a slight decrease in particulate matter and NOx occurs for each mode. In each case, the PM-NOx emissions point shifts toward the origin, which further demonstrates the viability of reducing diesel engine emissions via oxygen addition.

**CARBON MONOXIDE**

Table 4 reports the weighted brake specific CO (BSCO) emissions. There is no clear trend in this data, although there is an increase in CO for most of the mode positions. If CO per unit of fuel is reviewed, one can see that for each of the lower speed and load modes, a definite increase in the CO for CETANER™ is observed. This is shown in Figure 7. This may again support the rationale that during the low speed and low load conditions, CO formed during early reactions of the fuel are being halted from final conversion to CO2. This was postulated by Litzinger and coworkers [11]. As explained by Glassman, the conversion of CO to CO2 would be a function of the size of the hydroxyl radical pool, which does not grow until after all the original fuel and hydrocarbons have been consumed [31]. Since the concentration of hydroxyl radicals is important in the rate of CO oxidation, the additional molecules of oxygen with the monoglyme and diglyme may be playing a role in providing excess CO and CO2 which continue the creation of the hydroxyl radical pool. In addition, Flynn and coworkers show through kinetic simulations that the addition of the oxygen in the fuel leads to reduced amounts of soot precursors, and larger amounts of carbon leaving the fuel rich premixed combustion zone as CO [28].

![Figure 4. Particulate Matter Emissions, g/kg Fuel Basis.](image)

**NITROGEN OXIDES**

Table 4 reports the weighted brake specific NOx (BSNOx) emissions for the CETANER™-diesel blend. Using the AVL 8-mode weight factors, the net NOx emission reduction was 5.35%. Shown in Figure 5, for modes 4 through 8 a reduction of NOx per unit fuel consumed was observed. In general, the data follows an expected trend, in that at higher engine speeds, NOx is lower, and as the load increased, NOx increased. For the other three modes, the data does not show significant conclusions. There are conflicting reports in the literature as to whether oxygenates do indeed reduce NOx[2,4,5,11,15]. The data from this engine, however, shows a reduction in NOx emissions on a brake specific basis, for most modes. Choi and Reitz [4] observed that there is a small penalty on the NOx emissions when using a split injection strategy (two fuel pulses) with an oxygenated fuel, which could be affecting the results for mode 1 and 2 for this particular engine. Because the unique multiple fuel injection strategy of the Navistar T444E especially is more pronounced at lower speeds, the NOx reduction could occur due to greater mixing effect in the cylinder during the combustion event.

![Table 4. AVL 8-mode Weighted Gaseous Emissions Results, CETANER™-Diesel Blend, Brake Specific Basis](image)
TOTAL HYDROCARBONS

The weighted brake specific emissions for the CETANER™-diesel blend are reported in Table 4. In general, the HC emissions decrease with higher engine loads, as the engine combustion efficiency increases. In this work for all engine loads, HC emissions remain unchanged as compared to the baseline diesel, as can be seen in Figure 8. These results are consistent with other work by Hess and coworkers [24].

\[ \text{Figure 5. NOx Emissions per Unit Fuel Consumed for CETANER™ Addition, g/kg Fuel Basis.} \]

\[ \text{Figure 6. Brake-Specific Particulate Emission vs. NOx Emission Tradeoff.} \]

FUEL CONSUMPTION

Figure 9 reports the brake specific fuel consumption (BSFC) for CETANER™ and DME addition. The general trend shows an increase in the amount of fuel required to maintain the same speed and load. This is due to the slightly lower calorific value of the CETANER™ and DME blends, as shown in the Fuel Properties of Table 3. However, when fuel consumption is calculated on an energy basis, the energy consumption results are not significantly different.

\[ \text{Figure 7. CO Emissions per Unit Fuel Consumed for CETANER™ Addition, g/kg Fuel Basis.} \]

\[ \text{Figure 8. Total Hydrocarbon Emissions per Unit Fuel Consumed for CETANER™ Addition, g/kg fuel} \]

The results from this work are significant in that they confirm previous data for particulate reductions. The reductions correlate well with those of 2 wt.% oxygen addition via diglyme of Liotta and Montalvo [5]. For this data, a NOx reduction was shown for specific modes and as an overall number, which would be contrary to most work found in the literature. The CO emissions seem to follow the trends as reported [11,24,28].
mode 4 CO data raises questions, but will be reviewed further.

![Graph of Fuel Consumption, Brake Specific Basis.](image)

Figure 9. Fuel Consumption, Brake Specific Basis.

**CONCLUSION**

On-going research continues in testing and reviewing the affects of oxygenates on the composition of emissions from diesel engines. Optimization of the pressurized fuel system for DME-diesel blends is underway. The development of this fueling system is a significant advance toward practical use of DME as a diesel fuel. The results in this paper lead to the following conclusions:

The emissions results with CETANER™ addition are consistent with previous work and shown significant particulate emissions reductions in a DI diesel engine. NOx emissions were moderately lower, while HC emissions were unchanged and CO emissions increased at low load. Results with the pressurized fueling system yielded scattered emissions results, with and without DME addition to the base diesel fuel. These difficulties may stem from an inability to properly cool the fuel that resides in the rail. The strategy outlined here for combining DME and diesel fuel under pressure can provide an effective means of fueling an engine with DME without excessive modifications to the engine.

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**REFERENCES**

VISCOSITY OF DME-DIESEL FUEL BLENDS

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Introduction
The need to reach ever tightening NOx and particulate emissions standards has placed a tremendous amount of pressure on the fuel, lubricant, engine and vehicle manufacturers. However, in the 1990’s studies of direct injection diesel engines fueled by dimethyl ether demonstrated particulate emissions below the ULEV standard and NOx emissions that approach or achieve ULEV levels, without exhaust aftertreatment [1,2]. Until those tests, DME had not been considered as a primary replacement fuel. Previously, DME had been considered as a methanol ignition improver for methanol powered vehicles [3-6]. At present, the predominant use for DME is as an environmentally benign aerosol propellant, since DME is non-toxic and is easily degraded in the troposphere [7]. Recent work on DME has focused on its use in advanced technology, direct-injection (DI) engines as a neat fuel [8-12].

However, DME has significantly different physical properties than diesel fuel including a low critical point, low viscosity, negligible lubricity and a high vapor pressure. In the present work, DME has been blended into diesel fuel to obtain a fuel mixture that retains the desirable physical properties of diesel fuel but includes the cleaner burning capability of DME. The miscibility and viscosity of blends of DME and diesel fuel were characterized using pressurized, optically accessible instrumentation. These physical property measurements are part of a comprehensive study of the operation of a turbodiesel engine on DME-diesel blends which is leading to a field demonstration of this fueling strategy [13].

Experimental
Two different high pressure cells were adapted for studying the miscibility and viscosity of blends of DME and diesel fuel. One permitted the fuel mixtures to be held at pressures up to 200 psi to examine miscibility by visual inspection of blends over extended periods of time. The fuels were deemed to be miscible if no evidence of phase separation was observed. The other instrument is a high pressure viscometer based on a capillary tube held within a pressurized chamber suitable for measurements at pressures up to 3500 psi.

Miscibility Measurements. Qualitative studies of the miscibility of blends of DME and a federal low sulfur (300 ppm) “emissions certification” diesel fuel (Specified Fuels “ECD LS”) were performed under pressures above 90 psi. Blends from 25 wt.% DME up to 75 wt.% DME in diesel fuel were examined. Diesel fuel was gravity fed into an optically accessible pressure chamber, while DME was delivered from a cylinder of liquefied DME through an opening in the bottom of the pressure chamber. Pressures in the chamber were raised by feeding nitrogen above the fuel mixture to attain 90 psi or greater in the chamber.

Viscosity Measurements. Quantitative measurements of the viscosity of blends of DME in the federal low sulfur fuel were obtained using a high pressure viscometer, using capillary tubes that provided optimal measurement accuracy depending on the viscosity of the fuel mixture. Figure 1 shows a photograph of the high pressure chamber where the capillary tubes are located.

Results and Discussion
Miscibility Measurements. The DME was observed to rapidly mix uniformly with the diesel fuel at all blend ratios. Over time, a blend that was initially not well mixed would become uniform, but injection of the DME from below the pool of diesel fuel was a particularly effective means of rapidly obtaining a uniform mixture.
Viscosity Measurements. Observations of the viscosity of the blends of DME and diesel fuel are summarized in Figure 3. Measurements were obtained over a range of pressures with the viscometer housing immersed in a constant temperature bath at 100°F (38°C). Results obtained at three different levels of chamber pressure are plotted in Figure 4 to show the impact of DME content on viscosity.

![Figure 3. Viscosity of DME-diesel blends at pressures from 500 to 2500 psi.](image)

Figure 3. Viscosity of DME-diesel blends at pressures from 500 to 2500 psi.

![Figure 4. Blend response of viscosity to DME addition at various pressures.](image)

Figure 4. Blend response of viscosity to DME addition at various pressures.

These two figures show that viscosity decreases rapidly at low levels of DME addition. For instance at 25 wt.% DME addition, viscosity falls by more than a factor of 2, from the more than 2.5 cSt value of the neat diesel fuel to roughly 1 cSt. This non-linear blending response demonstrates that even modest addition of DME to diesel fuel brings the fuel blend below the ASTM diesel viscosity specification of 1.39-4.20 cSt at 40°C.

These viscosity measurements are among the first reported for DME under elevated pressures and are the first reported for blends in diesel fuel. Recent work by Sivebaek et al. [14] also considered the viscosity of DME, in particular with addition of lubricity and viscosity enhancing additives. They developed a volatile fuel viscometer (VFVM) that was designed to handle DME, neat or additized. They measured kinematic and dynamic viscosities of pure DME of 0.185 cSt and 0.122 cP at 25 °C, which compares well with their measurements were performed at 5 bar pressure, roughly 75 psi. In the present study, no DME blends were examined at a pressure below 500 psi, but at this pressure the viscosity of neat DME was found to be 0.21 cSt. Extrapolating data for neat DME from the present study to a pressure of 75 psi yields an estimate of 0.2 cSt, which is in reasonable agreement with the value of 0.185 cSt obtained by Sivebaek et al. They also concluded that additized DME cannot reach the same viscosity and lubricity as diesel fuel. They suggest that rather than using additives to allow fuel systems to tolerate DME, the solution is to design the pumps so that they can handle pure DME.

Conclusions
Blending DME in diesel fuel is one option to utilize DME in diesel engines without drastic redesign of fuel pumps and fuel injectors. However, even modest addition of DME into diesel fuel significantly reduces the viscosity of the fuel mixture. Addition of as little as 25 wt.% DME into diesel fuel reduces fuel viscosity below the ASTM specification. This suggests that viscosity rather than miscibility is the limiting factor in blending DME with diesel fuel.

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Emission Characteristics of a Navistar 7.3L Turbodiesel Fueled with Blends of Dimethyl Ether and Diesel Fuel

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Peter Tijm
Rentech, Inc.

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ABSTRACT

Several oxygenates have been proposed and tested for use with diesel fuel as a means of reducing exhaust emissions. This paper examines dimethyl ether (DME), which can be produced in many ways including via Air Products and Chemicals, Inc’s Liquid Phase Technology (LPDME). Modest additions of DME into diesel fuel (2 wt.% oxygen) showed reductions in particulate matter emissions, but the previous data reported by the author from a multicylinder Navistar 7.3L Turbodiesel engine were scattered. In this study, experiments were performed on a multi-cylinder Navistar 7.3L Turbodiesel engine to repeatably confirm and extend the observations from the earlier studies. This is an important step in not only showing that the fuel does perform well in an engine with minor modifications to the fuel system, but also showing that DME can give consistent, significant results in lowering emissions. The DME and diesel blends tested were to achieve a net addition of 5 and 10 wt. % oxygen in the blended fuel. The data confirms that the addition of DME can reduce the particulate emissions from a compression ignition engine. However, the NOx emissions were not favorable for all conditions. It is believed that through further modification of injection timing, NOx emissions can be effectively reduced.

INTRODUCTION

Demand for cleaner burning diesel fuels is growing worldwide, as governmental regulations make emissions reductions necessary. In the U.S., future regulations that take effect in 2004 and 2007 will require diesel engine and vehicle manufacturers to review all aspects of the vehicle system design [1]. To achieve substantial reductions in emissions, it is thought that reformulated diesel fuels will play an important role. The reformulation of diesel fuels could include lowering the sulfur content, lowering the aromatic content, or potentially the addition of oxygen within the fuel.

A solution for reducing emissions from future and current diesel vehicles is to modify the fuel, without the need to modify the engine hardware. It has been shown that many oxygenates are effective at reducing particulate emissions from diesel engines [2-21]. Therefore, much research has focused on screening of oxygenated fuel additives, including alcohols, esters and ethers. Of particular interest are the glycol ethers, which have been shown to be very effective as blends and as neat fuel. This study focuses on the use of dimethyl ether, which has the chemical formula: CH3-O-CH3.

Dimethyl ether is a common chemical used as an aerosol propellant [22]. The properties of DME are given in Table 3, and are compared to the diesel fuel used for the baseline testing for this experiment. DME is a liquid at low pressure and standard temperature, and is relatively easy to handle. Over the past ten years, researchers have begun to consider the use of DME as a fuel. Because the cetane number and ignition temperature are close to that of diesel fuel, DME was thought to be an excellent substitute for use in compression ignition engines. However, there were some drawbacks to using the fuel, including the reduced viscosity and lubricity of the fuel in neat form, as well as fuel compressibility effects [23,24].

To potentially overcome the fuel property effects of DME, as well as reduce emissions, the experiments for this study focus on mixing dimethyl ether with diesel fuel. The initial goal is to determine the effect of the oxygen concentration on the emissions, with minimal engine modifications. In this part of the work, no changes have
been made to the fuel injection timing, fuel injectors, or engine programming. Changes to the fuel system have been made to allow for the fuel to be delivered to the common rail as a liquid by maintaining the DME-diesel blend at over 100 psi.

Over the last ten years, many researchers have begun to evaluate the performance and emissions effects of neat dimethyl ether. Sorensen and Mikkelsen [25] found that for a fixed speed and across various loads, the particulate and NOx emissions from a .273 Liter direct injection single cylinder engine fueled with neat dimethyl ether could be significantly reduced as compared to emissions when fueled with diesel. In the same study, the HC and CO emissions showed little to no change. Later, Sorensen and Mikkelsen [26] further studied the HC emissions from this same engine, and found that there was an increase in the HC emissions when using neat DME, with more methane found than in a typical diesel engine, and less light hydrocarbons. With another engine, Christensen and Sorenson [27] looked at various effects on the suite of emissions when using neat DME. Of particular interest, the NOx emissions were significantly reduced when the injection timing was retarded towards Top Dead Center (TDC). However, there was an increase in the CO emissions, and little effect on the HC emissions. Other effects tested determined that lower injector opening pressure reduces NOx, and nozzle types did not seem to influence NOx emissions. Experiments completed by Kajitani and coworkers [28] also supported effects of injection timing on reducing NOx, and having little effect on HC emissions, from a single cylinder Yanmar engine fueled with neat DME.

However, in the work completed by Hupperich and coworkers [29] with a 1.75 liter single cylinder engine for the ECE R49 13-mode test, the cumulative emissions show some differing results. With the use of neat DME, HC emissions are reduced. The trends with the other emissions are similar to what had been determined with previous studies. One difference to note is the change in injection nozzle size, which may have affected the emissions results in allowing for more complete combustion of all fuels tested in an effort to maintain consistent conditions.

Recently, experiments completed by Ikeda and coworkers [30] with a single cylinder engine using a binary fuel injection method, showed similar NOx emissions between diesel fuel and 40% DME mixed with diesel fuel, as injection timing was retarded. Also, HC emissions increased and smoke emissions were reduced as injection timing was retarded. In addition, comparisons were made as a function of BMEP. NOx was reduced, HC remained constant and smoke increased with increasing Brake Mean Effective Pressure (BMEP). The experiments also included % DME fractions, but no comparisons were made to the baseline diesel fuel.

Many researchers have been evaluating the performance of other oxygenates including blends of glycol ethers with diesel fuel, and have observed decreases in particulate matter emissions with increasing oxygenate concentration. Most recently, Hallgren and Heywood [31] prepared a review of the collection of work which showed that as the oxygen content of the fuel increases, the particulate matter is reduced, suggesting that this occurs regardless of chemical structure or molecular weight. However, their actual testing showed that the oxygenate structure did impact particulate emissions. Studies completed by Hess et al. [10] as well as by Litzinger and coworkers [11,12] have shown that higher molecular weight glycol ethers are also effective in reducing particulate matter emissions, although to a lesser extent than monoglyme or diglyme.

Although it has been shown that glycol ethers effectively reduce particulate emissions, the fundamental mechanisms of the reduction have not been clearly identified. There has been some work in simulating the ignition and rate mechanism behavior of dimethyl ether in comparison to dimethoxymethane [19]. Also, oxidation mechanisms have been proposed for gaseous forms of DME [32-34]. Limited data is available for many of the liquid oxygenates under consideration.

For this experimental work, an Emissions Certification Diesel-LS, provided by Specified: Fuels & Chemicals, LLC., used in combination with dimethyl ether, was evaluated in a multi-cylinder direct injection (DI) engine. In-cylinder pressure measurements provided information about the impact of the oxygenated fuel on the combustion process. In addition, fuel property tests were performed on the base fuels, as well as for the blended fuels. These measurements were used to understand and describe the combustion behavior. Since dimethyl ether is a vapor at 1 atm, the fuel property tests for the fuel blends were performed under pressure to maintain the DME in a liquid state. Because not all tests could be performed, data for DME reported in literature is used for most values. Therefore, the fuel system of the engine was redesigned to accommodate the pressurized fuel delivery.

**EXPERIMENT**

**TEST ENGINE:** For the purpose of studying the effects of fuel additives on light-medium duty diesel combustion, a Navistar T444E 7.3L Turbodiesel engine was coupled to a 450 horsepower Eaton (Model AD-1802) eddy-current dynamometer. The specifications for the engine are given in Table 1. A Pentium PC with Keithley Metrabyte DAS-1800 data acquisition card was connected to the engine to log real-time engine parameters. These parameters included engine speed, torque, and power from the engine. A Modicon PLC was used to record temperatures from the engine, as well as, for the entire experimental system. Intake airflow rates were determined via an electronic flow sensor, which
was calibrated using a laminar flow element. Fuel consumption was monitored using a precision Sartorius scale (Model EA60EDE-1), with an accuracy of ± 2 grams. Figure 1 shows the test cell set up, and additional equipment used for emissions monitoring.

Table 1. Characteristics of the 1998 Navistar T444E 7.3L Turbodiesel engine

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Value</th>
</tr>
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<tbody>
<tr>
<td>Displacement</td>
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</tr>
<tr>
<td>Bore</td>
<td>4.11 inch (104.39mm)</td>
</tr>
<tr>
<td>Stroke</td>
<td>4.18 inch (106.20mm)</td>
</tr>
<tr>
<td>Rated Power</td>
<td>190 HP @2300 RPM</td>
</tr>
<tr>
<td>Peak Torque</td>
<td>485 lbf-ft @ 1500 RPM</td>
</tr>
<tr>
<td>Configuration</td>
<td>Turbo charged, Intercooled (Air-to-Air), Direct Injection</td>
</tr>
<tr>
<td>Injection Scheme</td>
<td>HEUI- Hydraulically actuated, electronically controlled unit injectors</td>
</tr>
<tr>
<td>Low Idle Speed</td>
<td>700 RPM</td>
</tr>
<tr>
<td>Features</td>
<td>Split- shot injection</td>
</tr>
<tr>
<td>Compression Ratio</td>
<td>17.5:1</td>
</tr>
</tbody>
</table>

Table 2. AVL 8-Mode Test for the Navistar T444E 7.3L Turbodiesel engine

<table>
<thead>
<tr>
<th>Mode</th>
<th>Speed (rpm)</th>
<th>Load (ft-lb)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>700</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>876</td>
<td>84</td>
</tr>
<tr>
<td>3</td>
<td>1036</td>
<td>224</td>
</tr>
<tr>
<td>4</td>
<td>1212</td>
<td>357</td>
</tr>
<tr>
<td>5</td>
<td>2300</td>
<td>77</td>
</tr>
<tr>
<td>6</td>
<td>2220</td>
<td>178</td>
</tr>
<tr>
<td>7</td>
<td>2220</td>
<td>307</td>
</tr>
<tr>
<td>8</td>
<td>2124</td>
<td>409</td>
</tr>
</tbody>
</table>

Figure 1. Multicylinder Test Cell, Navistar T444E Turbodiesel

TEST PROCEDURE- In this work, an AVL 8-mode test procedure has been utilized as a model for diesel emissions tests. The AVL 8-mode test was designed to correlate to the U.S. Federal Heavy-Duty Transient Test procedure through a weighted 8- mode steady state test procedure. The 8 modes are a combination of speeds and loads, that produce the same emissions output as would be recorded for a transient cycle [35]. For this engine, the test procedure included the speed and load settings, shown in Table 2.

EMISSIONS EQUIPMENT- An extended warm-up period was used to prepare the engine for testing. The sampling and measurements during each mode commenced when the exhaust temperatures reached steady state. During this time, RPM and torque were maintained within 1-2% of the target test conditions. Once steady-state operation was achieved, a portion of the exhaust gas was passed through a Sierra Instruments BG-1 micro-dilution test stand with a constant dilution air / sample flow ratio of 8:1 and a total flow of 150 liters/min. These settings were chosen in order to maintain the filter temperature below the EPA specification of 52°C. Particulate collection occurred on Pallflex 90mm filters (Type EMFAB TX40HI20-WW), conditioned in an environmental chamber at 25°C and 45% relative humidity before and after sampling. Five particulate samples were taken for each fuel at each test mode.

Exhaust gas analyses were completed using a Nicolet Magna 550 Fourier Transform Infrared (FTIR) Spectrometer. For each mode, five gas samples were analyzed for CO2, CO, NO and NO2. Also, a Rosemont Analytical on-line O2 analyzer was used to monitor the percent oxygen in the exhaust gas. The oxygen readings were used in conjunction with the mass flow sensor to determine and verify the air / fuel ratio. Additionally, total hydrocarbon emissions were monitored using a California Analytical Instruments Model 300 Heated Flame Ionization Detector (HFID) Total
Hydrocarbon Gas Analyzer. For the total hydrocarbon measurements, undiluted exhaust gas was collected via a heated sample line, which was maintained to 190°C. Calibration of all equipment was completed prior to each day of testing.

PRESSURE TRACE ANALYSIS- In order to observe the impact of the oxygenated blends on combustion and heat release, the combustion chamber of cylinder 1 of the engine was fitted with a Kistler 6125A pressure probe. The pressure sensor was used with a Kistler 2612 optical crank angle encoder to provide time resolved in-cylinder pressure traces of the combustion event. Pressure, crank angle, and TDC trigger signals were acquired with a Keithley DAS-1800 data acquisition card operating in a “burst” mode. The pressure traces were analyzed with PtrAn V.02, a software product designed by Optimum Power.

TEST FUELS- Previous work has been completed comparing the increasing percentage of oxygenate mixed with diesel fuels within several types of engines [9-18,36]. For this testing, comparisons are made between a 5 wt.% and 10 wt. % oxygen via blending of DME in diesel fuel. The baseline diesel fuel properties, as well as test fuel properties are given below in Table 3. Because of the difficulty in obtaining experimentally the fuel blend properties for DME as a liquid, the properties available in the literature for neat DME are represented, as well as linear calculation of the blends.

PRESSURIZED FUEL DELIVERY SYSTEM FOR DIESEL-DME BLENDS -Dimethyl ether (DME) is a liquefied gas. At room temperature and atmospheric pressure, it is a gas, but changes to a liquid at a moderate pressure. DME is currently manufactured by DuPont Fluorochemicals under the trade name Dymel A. For the purposes of the experimental design, information regarding the vapor pressure and density changes with temperature are available in the Technical Information (ATB-25) bulletin from DuPont.

In tests conducted, DME was found to be miscible with #2 diesel fuel. Miscibility tests were carried out in a pressurized vessel with a glass observation window. The two fuels were introduced taking care not to mix them. Diesel was introduced first into the bottom of the vessel. DME, which has a specific gravity less than diesel fuel, was then introduced on top of the diesel fuel. Thus, initially there were two distinct layers. The two layers were then observed to mix together without physical agitation after a period of 5 to 6 hours to form a homogeneous mixture. The DME was about 60% by mass in this mixture. Furthermore, no separation was observed after standing undisturbed for about 3 days.

A schematic of the modified fuel system is shown in Figure 2. The fuel system on the T444E engine had to be modified to account for the need to deliver fuel at elevated pressure. The fuel rail in the cylinder head of the engine receives fuel at a pressure of about 70 psi. Fuel from this rail is then fed to the injectors.

A study was performed using #2 diesel fuel to measure the temperature rise of the fuel in the fuel rail. This measurement, coupled with the fuel consumption gave an approximate heat transfer rate between the cylinder head and the fuel in the gallery. A maximum target temperature was chosen for the diesel-DME blend based on the vapor pressure curve of DME and the pressure rating of the fuel rail. The required change in fuel recirculation flow rate was then calculated based on the above observations. This recirculated fuel was then cooled down using a water cooled heat exchanger. The fuel delivery pump was sized based on the above calculations.

Table 3. Fuel Properties

<table>
<thead>
<tr>
<th>Fuel Property</th>
<th>ASTM Method</th>
<th>ASTM Spec.</th>
<th>Base Diesel</th>
<th>DME</th>
<th>25 wt% DME in Diesel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Viscosity, 40°C, cSt</td>
<td>D 445</td>
<td>1.39-4.20</td>
<td>2.2</td>
<td>.25</td>
<td>.95[43]</td>
</tr>
<tr>
<td>API Gravity</td>
<td>D 287</td>
<td>API 30</td>
<td>35.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cloud Point (°F)</td>
<td>D 2500</td>
<td>&lt;0</td>
<td>4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pour Point (°F)</td>
<td>D 2500</td>
<td>&lt;0</td>
<td>&lt;0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flash Point (°F)</td>
<td>D 93</td>
<td>125</td>
<td>166</td>
<td>-42</td>
<td></td>
</tr>
<tr>
<td>Calorific Value (BTU/lb)</td>
<td>D 2015</td>
<td>19700</td>
<td>19483</td>
<td>12228</td>
<td>17669*</td>
</tr>
<tr>
<td>Density (kg/m³)</td>
<td>D 4052</td>
<td>.845-.855</td>
<td>.848</td>
<td>.660</td>
<td>.801*</td>
</tr>
<tr>
<td>Cetane Number</td>
<td>D 613</td>
<td>46-48</td>
<td>47.4</td>
<td>&gt;55</td>
<td>&gt;55*</td>
</tr>
</tbody>
</table>

* Projected

Selecting a pump for DME was challenging due to the properties of DME. Gasket material for the pump had to be modified, as common materials such as Viton and buna-N have been found to be unsatisfactory. A fuel filter with a high filter surface area and high pressure capacity was needed. A modified propane filter was selected for the application. The fuel tank consisted of a modified 60 lb capacity LPG cylinder which was pressure tested at 120 psi prior to use.
From the previous studies on this same engine for the 2 wt.% oxygen in diesel, the cooling capacity of the heat exchanger and fuel circuit was determined to be insufficient, based on the fuel temperatures recorded, as well as, observed engine instabilities [37]. Therefore, the system shown in Figure 2 represents the modifications made, which included the addition of a second fuel coil in the cooled bath, and a larger chiller unit for cooling the bath. Additionally, the system was pressurized to 150 psi, which then increases the allowable fuel temperature before the DME becomes vapor.

RESULTS AND DISCUSSION

In this next section, detailed results are provided for the effect of the additive on emissions. Fuel property data is available in Table 3 so that comparisons of the combustion data can be analyzed. Through an uncertainty analysis, based on methods described by Moffat, error bars showing the 95% confidence intervals are presented in each figure [38].

PARTICULATES- As noted previously in the discussion, oxygenates traditionally reduce particulate emissions. The data from this testing indicates that this general trend is for all modes the engine was able to operate, as shown in Figure 3. Using the AVL 8-mode test, the net particulate emission reductions for each mode are found in Table 4. Because the engine was not able to be run for modes 4 and 8 for each additive, the data was presented in a mode by mode comparison. For most modes, shown in Figure 3, particulate reductions were observed on a particulate matter vs. fuel consumed basis. As the engine load is increased, the particulate emission is lowered, except for mode 4. This is true for the low as well as the high engine speeds. The trends follow what would be expected from prior work on oxygenated fuels. Mode 4 operates the engine near the peak torque, and the overall emissions change, possibly due to a change in the injection timing. As has been shown in previous work, this particulate reduction is due to a reduction in the soot portion of the emission, which would result in a percentage increase in the soluble organic faction (SOF) portion [6]. This has also been confirmed more recently by Sidhu and coworkers [39], with DME giving the highest SOF.

OXIDES OF NITROGEN (NO\textsubscript{x})-Table 4 reports the brake specific NO\textsubscript{x} (BSNO\textsubscript{x}) emissions. Figure 4 shows that at lower loads NO\textsubscript{x} decreases. At higher loads, NO\textsubscript{x} increases. The oxygenated additive leads to reduced NO\textsubscript{x} at the lower load conditions, possibly due to the injection timing of the engine. The NO\textsubscript{x} emission decreases for the 5 wt. % oxygen, and then begins to increase with additional oxygen content, as shown in modes 1 and 2. The Hydraulically actuated Electronically controlled Unit Injector (HEUI) use a split shot injection at the lower engine speeds, which may be providing better mixing to the fuel and air, and thus reducing NO\textsubscript{x}. At higher engine speeds, NO\textsubscript{x} is lower, and NO\textsubscript{x} increases as load increases. There are conflicting reports in the literature as to whether oxygenates increase or decrease NO\textsubscript{x} emissions [3,5,6,12,16]. The data from this engine, however, shows a reduction in NO\textsubscript{x} emissions on a brake specific basis, for most modes. Choi and Reitz [5] observed that there is a small penalty on the NO\textsubscript{x} emissions when using a split injection strategy (two fuel pulses) with an oxygenated fuel, which could be affecting the results for modes 1 and 2 for this particular engine. Because the unique multiple fuel injection strategy of the Navistar T444E is especially predominant at lower speeds, the NO\textsubscript{x} reduction could occur due to better improved mixing effect in the cylinder during the combustion event.
Table 4. AVL 8-mode Emissions Results per mode, Brake Specific Basis

### a. Particulate Emissions Per Mode

<table>
<thead>
<tr>
<th>Mode</th>
<th>Baseline Diesel (g/bhp-hr)</th>
<th>5wt. % oxygen via DME (g/bhp-hr)</th>
<th>10wt. % oxygen via DME (g/bhp-hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3.36</td>
<td>3.44</td>
<td>2.87</td>
</tr>
<tr>
<td>2</td>
<td>.224</td>
<td>.149</td>
<td>.118</td>
</tr>
<tr>
<td>3</td>
<td>.091</td>
<td>.069</td>
<td>.082</td>
</tr>
<tr>
<td>4</td>
<td>.209</td>
<td>.095</td>
<td>NA</td>
</tr>
<tr>
<td>5</td>
<td>.339</td>
<td>.255</td>
<td>.214</td>
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<td>6</td>
<td>.137</td>
<td>.128</td>
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<td>7</td>
<td>.078</td>
<td>.086</td>
<td>.057</td>
</tr>
<tr>
<td>8</td>
<td>.068</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

### b. NOx Per Mode

<table>
<thead>
<tr>
<th>Mode</th>
<th>Baseline Diesel (g/bhp-hr)</th>
<th>5wt. % oxygen via DME (g/bhp-hr)</th>
<th>10wt. % oxygen via DME (g/bhp-hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>43.91</td>
<td>24.74</td>
<td>36.54</td>
</tr>
<tr>
<td>2</td>
<td>3.31</td>
<td>3.21</td>
<td>4.14</td>
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<tr>
<td>3</td>
<td>3.52</td>
<td>3.85</td>
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<td>3.35</td>
<td>4.01</td>
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<tr>
<td>5</td>
<td>3.18</td>
<td>2.36</td>
<td>2.40</td>
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<tr>
<td>6</td>
<td>2.44</td>
<td>2.05</td>
<td>2.11</td>
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<tr>
<td>7</td>
<td>2.37</td>
<td>2.67</td>
<td>2.99</td>
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<tr>
<td>8</td>
<td>3.42</td>
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</table>

### c. CO Per Mode

<table>
<thead>
<tr>
<th>Mode</th>
<th>Baseline Diesel (g/bhp-hr)</th>
<th>5wt. % oxygen via DME (g/bhp-hr)</th>
<th>10wt. % oxygen via DME (g/bhp-hr)</th>
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<tbody>
<tr>
<td>1</td>
<td>40.0</td>
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<td>3</td>
<td>.560</td>
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<td>.671</td>
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<td>8.12</td>
<td>7.63</td>
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<tr>
<td>5</td>
<td>2.33</td>
<td>4.16</td>
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<tr>
<td>6</td>
<td>.913</td>
<td>1.344</td>
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<tr>
<td>7</td>
<td>.691</td>
<td>.875</td>
<td>.916</td>
</tr>
<tr>
<td>8</td>
<td>.493</td>
<td>NA</td>
<td>NA</td>
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</tbody>
</table>

### d. Hydrocarbon Per Mode

<table>
<thead>
<tr>
<th>Mode</th>
<th>Baseline Diesel (g/bhp-hr)</th>
<th>5wt. % oxygen via DME (g/bhp-hr)</th>
<th>10wt. % oxygen via DME (g/bhp-hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>34.3</td>
<td>51.4</td>
<td>48.5</td>
</tr>
<tr>
<td>2</td>
<td>.707</td>
<td>1.78</td>
<td>1.54</td>
</tr>
<tr>
<td>3</td>
<td>.211</td>
<td>.953</td>
<td>1.54</td>
</tr>
<tr>
<td>4</td>
<td>.137</td>
<td>.561</td>
<td>NA</td>
</tr>
<tr>
<td>5</td>
<td>.549</td>
<td>3.07</td>
<td>3.18</td>
</tr>
<tr>
<td>6</td>
<td>.216</td>
<td>1.22</td>
<td>1.28</td>
</tr>
<tr>
<td>7</td>
<td>.127</td>
<td>.777</td>
<td>.430</td>
</tr>
<tr>
<td>8</td>
<td>.094</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>
Figure 4. NOx Emission Results per unit fuel consumed, g/kg fuel

Figure 5 presents the particulate matter vs. NO\textsubscript{x} tradeoff per mode. As can be seen for modes with lower loads, as particulates are reduced, NO\textsubscript{x} is reduced. However, for modes 3, 4 and 7, an increase in NO\textsubscript{x} with decreasing particulates is observed. In some cases, the PM-NO\textsubscript{x} emissions point shifts toward the origin, which demonstrates that oxygen addition is a viable means of reducing diesel engine emissions. However, this is not true for all cases, and could possibly be due to changes in injection timing by the engine controls, and not necessarily a function of the changing fuel properties.

Figure 5. Brake-Specific Particulate Emission vs. NO\textsubscript{x} Emission Tradeoff

CARBON MONOXIDE (CO)- Table 4 reports the brake specific CO (BSCO) emissions. On a CO per unit of fuel basis, CO emissions increase as the wt. % oxygen is increased, with a decreasing effect as the load increases. In general, CO decreases as load increases, as shown in Figure 6. This may again support the idea that during the low speed and low load conditions, CO formed during early reaction of the fuel is impeded from conversion to CO\textsubscript{2}. This was postulated by Litzinger and coworkers [12]. As explained by Glassman, the conversion of CO to CO\textsubscript{2} would be a function of the size of the hydroxyl radical pool, which does not grow until most all the original fuel and hydrocarbons have been consumed [40]. Since the concentration of hydroxyl radicals is important in the rate of CO oxidation, the additional molecules of oxygen from DME may be playing a role in providing excess CO and CO\textsubscript{2} which continue the creation of the hydroxyl radical pool. In addition, Flynn and coworkers show through kinetic simulations that the addition of the oxygen in the fuel leads to reduced amounts of soot precursors, and larger amounts of carbon leaving the fuel rich premixed combustion zone as CO [41].

Figure 6. CO Emission Results per unit fuel consumed, g/ kg fuel

HC- The brake specific emissions for each mode are reported in Table 4. In general, the HC emissions decrease with higher engine loads, as the engine combustion efficiency increases. For all modes, HC emissions increase with oxygen addition, and decrease as engine load increases, as seen in Figure 7. For the lower engine speeds, as the oxygen addition increases, the HC emissions decrease. However, because very few data points are involved with this figure, it would be important to repeat the study. This data is inconsistent with what has been observed by some previous engine studies.
Figure 7. Hydrocarbon Emission Results per unit fuel consumed, g/ kg fuel

FUEL CONSUMPTION- Figure 8 reports the brake specific fuel consumption (BSFC) for the DME addition. The general trend shows an increase in the amount of fuel required to maintain the same speed and load. This is due to the slightly lower calorific value of the fuel blend, as shown in the Fuel Properties of Table 3. However, when fuel consumption is calculated on an energy basis, the energy consumption results are not significantly different.

Figure 8. Brake Specific Fuel Consumption (BSFC)

CONCLUSION

On-going research continues in testing and reviewing the effects of oxygenates on the composition of emissions from diesel engines. The results in this paper lead to the following conclusions:

- DME has been shown to reduce particulate emissions from a DI diesel engine. Through the addition of oxygen contained within the hydrocarbon structure of the fuel, additional CO and CO2 are present in the combustion process. This prevents the formation of soot precursors, which reduces the formation of particulates.

- Higher amounts of CO have been observed for each mode. This can be explained by the presence of the fuel born oxygen creating a larger pool of CO and CO2, which aids in creating hydroxyl radicals, thus quenching the combustion process and leaving CO not oxidized.

- It has also been observed that along with a reduction in particulate emissions per mode, there is a small NOx reduction for some modes. It is unclear what may be causing this, but several possible reasons are given. Multiple injections at lower loads and speeds may explain the reason for this data. Also, there may be some affect of the increased fueling required by the engine, as the fuel oxygen content changes. The net effect is some change in the ignition timing of the fuel, as a result of the engine control adjusting injection timings differently for each fuel.

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REFERENCES


DEVELOPMENT OF A DIMETHYL ETHER-FUELED SHUTTLE BUS
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Francis J. Waller
Air Products and Chemicals, Inc.
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Introduction
Dimethyl ether has been considered a potential ultra-clean replacement fuel for diesel engines [1]. Dimethyl ether (DME) burns “smokeless”, permits high levels of EGR for in-cylinder NOx control and can be produced from synthesis gas derived from fossil fuel or biomass resources. These potential benefits of DME have motivated studies of the physical properties, the lubricity concerns and the combustion performance of DME [2,3]. In the present work, we seek to operate a laboratory engine and a campus shuttle bus on DME. To overcome the low lubricity and low viscosity of DME so as to be able to operate a conventional, common rail, DI diesel engine on DME, we have chosen to blend DME and diesel fuel. The conversion of the laboratory engine and the shuttle bus required development of a pressurized fuel delivery system to maintain the DME-diesel fuel blend above the vapor pressure of DME. This paper summarizes the outcomes from analyses of fuel properties, the laboratory engine studies and the conversion of the shuttle bus.

Experimental
Viscosity of DME-Diesel Blends. Quantitative measurements of the viscosity of blends of DME in a federal low sulfur fuel were obtained using a high pressure viscometer, using capillary tubes that provided optimal measurement accuracy depending on the viscosity of the fuel mixture [4]. Measurements were obtained over a range of pressures with the viscometer housing immersed in a constant temperature bath at 100°F (38°C). Results obtained at three different levels of chamber pressure are plotted in Figure 1 to show the impact of DME content on viscosity.

Figure 1. Blend response of viscosity to DME addition at various pressures [4].

Laboratory Engine Studies. The fuel system on the Navistar 7.3L V-8 “T444E” turbodiesel engine had to be modified to permit delivery of the fuel blend at elevated pressure [5]. The fuel rail in the cylinder head of the engine receives fuel at a pressure of about 70 psi. Fuel from this rail is then fed to the injectors.

A study was performed using #2 diesel fuel to measure the temperature rise of the fuel in the fuel rail. This measurement, coupled with the fuel consumption gave an approximate heat transfer rate between the cylinder head and the fuel in the gallery. A maximum target temperature was chosen for the diesel-DME blend based on the vapor pressure curve of DME and the pressure rating of the fuel rail. The required change in fuel recirculation flow rate was then calculated based on the above observations. This recirculated fuel was then cooled down using a water cooled heat exchanger. The fuel delivery pump was sized based on the above calculations.

Selecting a pump for DME was challenging due to the properties of DME. Gasket material for the pump had to be modified, as common materials such as Viton and buna-N have been found to be unsatisfactory. A fuel filter with a high filter surface area and high pressure capacity was needed. A modified propane filter was selected for the application. The fuel tank consisted of a modified 60 lb capacity LPG cylinder which was pressure tested at 120 psi prior to use.

From previous studies on this same engine for 2 wt.% oxygen in the diesel fuel, the cooling capacity of the heat exchanger and fuel circuit was found to be insufficient, based on the fuel temperatures recorded, as well as, observed engine instabilities. Therefore, the system shown in Figure 2 presents the modifications made, which included the addition of a second fuel coil in the cooled bath, and a larger chiller unit for cooling the bath. Additionally, the system was pressurized to 150 psi, which then increases the allowable fuel temperature before the DME becomes vapor.

Operation of the laboratory engine on blends of DME and diesel fuel resulted in reduction of particulate matter emissions. As shown in Figure 3, blending at up to 25 wt.% DME in diesel, corresponding

Figure 2. Diagram of the pressurized fuel system [5].
to 10 wt.% oxygen addition resulted in significant reduction in particulate matter emissions.

**Shuttle Bus Conversion**

The final stage of this project was the conversion of a campus shuttle bus and operation of the bus on the DME-diesel fuel blends. To accomplish this goal, the pressurized fuel system on the laboratory engine was adapted for application to a Champion Motorcoach “Defender” model bus with the same model engine (Navistar T444E) as was used in the laboratory study. To simplify the requirements for the fueling station for the shuttle bus, mixing of the DME and diesel fuel is performed onboard the bus. Figure 4 shows the shuttle bus used in this study. The final design and the operational procedures were reviewed and modified in a detailed “HAZOP” or failure modes effects analysis (FMEA). The outcome of the HAZOP analysis was to increase the number of check valves, manual valves and redundancy in the system.

The system on the bus consists of a transfer pump that delivers diesel fuel from the existing diesel fuel tank to the tank for the blended fuels, a propane tank for a recreational vehicle. This LPG tank was modified only by replacing o-ring materials with Kalrez™ o-rings. The connections on the LPG tank permit diesel fuel to be transferred to the tank, while the tank is vented to the atmosphere. Then, DME is transferred into the LPG tank. During the refueling processes, a handheld controller notifies the operator of the fill level and transfer the fuel blend to the fuel rails in the cylinder heads of the engine. A backpressure regulator maintains the pressure in the LPG tank so that the desired proportions of fuel are transferred. Finally, a compressed cylinder of helium is connected to the LPG tank to provide a blanket of inert gas to maintain a minimum of 120 psig in the LPG tank and keep the DME in the liquid phase. A magnetically coupled gear pump serves to draw fuel from the LPG tank and transfer the fuel blend to the fuel rails in the cylinder heads of the engine. A backpressure regulator maintains the pressure in the fuel rails at a minimum of 150 psig, although during operation the rail pressure is typically near 200 psig. Fuel rejected from the rails passes through a pair of fuel coolers mounted in front of the radiator to keep the fuel temperature from rising above a bulk temperature of 50°C.

Upon completion of the majority of the conversion process, the bus was operated with the pressurized fueling system on diesel fuel but without DME blending. The bus was operated over several days at the Pennsylvania Transportation Institute’s test track near the Penn State University Park campus. During this shakedown process, emissions measurements were obtained in collaboration with Clean Air Technologies (Buffalo, NY) using their portable diesel emissions analyzer (XXX). Among the chief challenges faced during the conversion and shakedown tasks was the power requirements and stability of operation of the gear pump. However, in-field adjustments and modifications resulted in consistent and stable operation of the components of the pressurized fueling system. The shuttle bus will operate on the DME-diesel fuel blend, at 25 wt.% DME, through Fall 2002 during which time periodic performance, emissions and system integrity tests will be performed.

**Conclusions**

Operation of a commercial diesel engine on DME-diesel blends has been accomplished with minimal modification to the engine, apart from addition of a pressurized fuel delivery system. This technique permits operation of vehicles, in part, on DME without jeopardizing the long term durability of the engine. Consistent with operation of diesel engines on oxygenated fuels, particulate emissions with the DME-diesel fuel blends are substantially reduced.

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**References**

Development of a Dimethyl Ether (DME)-Fueled Shuttle Bus

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ABSTRACT

Dimethyl Ether (DME) is a potential ultra-clean diesel fuel. Its unique characteristics require special handling and accommodation of its low viscosity and low lubricity. In this project, DME was blended with diesel fuel to provide sufficient viscosity and lubricity to permit operation of a 7.3 liter turbodiesel engine in a campus shuttle bus with minimal modification of the fuel injection system. A pressurized fuel delivery system was added to the existing common rail injection system on the engine, allowing the DME-diesel fuel blend to be circulated through the rail at pressures above 200 psig keeping the DME in the liquid state. Fuel exiting the rail is cooled by finned tubed heat exchangers and recirculated to the rail using a gear pump. A modified LPG tank (for use on recreational vehicles) stores the DME-diesel fuel blend onboard the shuttle bus. A small cylinder of helium is used to provide a blanket of inert gas above the fuel mixture to keep the DME in the liquid state and to push the mixture to the fuel rails. A significant challenge is posed by the rapid increase in DME vapor pressure with increasing fuel temperature. The target for maximum fuel rail temperature was set at 50°C, which corresponds to a DME vapor pressure of 150 psig. Refueling was accomplished by mixing the diesel fuel and DME onboard the bus, with diesel fuel delivered from the existing diesel tank and DME delivered by 1000 lb cylinders at a small refueling station. The shuttle bus operates on the Faculty/Staff loop at the University Park campus of the Pennsylvania State University.

INTRODUCTION

The motivation for researching new fuels stems from several factors. Primarily governments worldwide, including the United States, are setting stricter standards for emissions from new engines and vehicles. This is being done for the benefit of human health and the environment, via the reduction of particulate matter and smog-causing NOx [1]. The final ruling in December 2000 from the U.S. EPA sets the new 2007 standards for heavy duty diesel engines to the following:

- Particulate Matter (PM) emissions to 0.01 grams per brake-horsepower-hour (g/bhp-hr) to take full effect in 2007, and
- NOx and non-methane hydrocarbons (NMHC) to 0.20 g/bhp-hr and 0.14 g/bhp-hr, respectively. These standards will be phased in after 2007 [2].

Table 1 shows the trend in the tightening of emission standards.

<table>
<thead>
<tr>
<th>Emissions Type</th>
<th>Current Regulation (g/bhp-hr)</th>
<th>2004 (g/bhp-hr)</th>
<th>2007 (g/bhp-hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NOx</td>
<td>4</td>
<td>.2</td>
<td></td>
</tr>
<tr>
<td>HC</td>
<td>1.3</td>
<td></td>
<td>.14 (NMHC)</td>
</tr>
<tr>
<td>NOx and HC</td>
<td>2.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PM</td>
<td>.1</td>
<td></td>
<td>.01</td>
</tr>
</tbody>
</table>

*Required on-board diagnostics (OBD) systems for vehicles between 8500 and 14000 lbs to be phased-in, beginning in 2005

NOx- Nitrogen Oxides; HC- Hydrocarbons; PM- Particulate Matter

Table 1. U.S. Heavy-Duty Diesel Engine Emission Standards [1, 2]

If fuels can be synthesized from domestic resources and engineered to meet the upcoming emissions standards, there will be additional benefits for the energy security of the United States [3]. These new fuels, must then undergo substantial testing to demonstrate their
compatibility with engine and vehicle technologies, or to demonstrate the extent to which engine and vehicle technologies need to be altered to make use of the fuel. This means that fuel development projects require both experimental “proof of concept” and field vehicle testing.

Dimethyl ether is a potential ultra clean diesel fuel that could be synthesized from a variety of feedstocks, which can support the use of alternative energy resources [10, 11]. DME is a compound that has been targeted for future use as a fuel in several countries around the world [4-6]. Motivation to use DME exists for several reasons. There has been confirmation that the fuel yields low particulate emissions and possibly lower NOx emissions [7-9].

The following review of the literature will cover a summary of the testing completed to examine the potential use of DME as a fuel. Many technical challenges have been discovered during this testing, shedding light on new approaches for using DME as a fuel. Researchers have started to study the fundamental nature of the combustion process, and to understand the mechanisms of DME combustion [12, 13]. Although use of DME has been demonstrated in compression ignition engines, technical challenges exist in transferring that knowledge to field vehicles.

PROPERTIES OF DIMETHYL ETHER

Compounds in which two hydrocarbon groups are bonded to one oxygen, represented as R-O-R', are called ethers. The organic groups bonded to the ether may be alkyl, aryl, or vinylic, and the molecule can either be an open chain or ring configuration [14]. Ethers commonly observed in long chain structures are referred to as linear ethers. As compared to alkanes of similar structure, where the CH2 group replaced the O atom, the boiling points of ethers are higher [14]. This class of oxygenated compounds have high cetane numbers and excellent cold flow properties [15].

Simply stated, dimethyl ether is an ether with two methyl groups on each side of an oxygen atom. Today, DME is predominantly used as an aerosol propellant because, in contrast to other aerosol propellants used previously, it is not harmful to the ozone layer [10]. Also, it is virtually non-toxic and is easily degraded in the upper atmosphere. It can be represented by the structure: CH3-O-CH3. The physical properties of DME are shown in Table 2 along with diesel fuel and propane for comparison [7, 9, 16, 17].

The properties that are significant for the use of DME as a fuel are cetane number, boiling point, and ignition temperature. However, the properties of concern are viscosity, heating value, and vapor pressure.

The cetane number describes the ignition quality of the fuel. The shorter the ignition delay the better the ignition quality of the fuel, and thus, the higher the cetane number. Since DME has a higher cetane number than conventional diesel fuel, it will ignite readily and burn more completely.

<table>
<thead>
<tr>
<th>Property</th>
<th>DME</th>
<th>Diesel</th>
<th>Propane</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical Formula</td>
<td>C2H6O</td>
<td>C10.8H18.7</td>
<td>C3H8</td>
</tr>
<tr>
<td>Mole Weight</td>
<td>46.07</td>
<td>148.6</td>
<td>44.11</td>
</tr>
<tr>
<td>Critical Temperature- °C</td>
<td>127</td>
<td>-</td>
<td>95.6</td>
</tr>
<tr>
<td>Boiling Point- °C</td>
<td>-24.9</td>
<td>71-193</td>
<td>-42.1</td>
</tr>
<tr>
<td>Vapor Pressure at 20 °C-kg/m²</td>
<td>5.1</td>
<td>&lt;0.01</td>
<td>8.4</td>
</tr>
<tr>
<td>Critical Pressure-bar</td>
<td>53.7</td>
<td>-</td>
<td>43</td>
</tr>
<tr>
<td>Liquid Viscosity- cP</td>
<td>.15</td>
<td>2-4</td>
<td>.10</td>
</tr>
<tr>
<td>Liquid Density at 20 °C-Kg/m³</td>
<td>668</td>
<td>800-840</td>
<td>501</td>
</tr>
<tr>
<td>Bulk Modulus (N/m²)</td>
<td>6.37E+0</td>
<td>1.49E+0</td>
<td>9</td>
</tr>
<tr>
<td>Specific Density,gas</td>
<td>1.59</td>
<td>-</td>
<td>1.52</td>
</tr>
<tr>
<td>Solubility in H2O at 20 °C g/l</td>
<td>70</td>
<td>Negligible</td>
<td>.12</td>
</tr>
<tr>
<td>Lower Heating Value-kJ/kg</td>
<td>28430</td>
<td>42500</td>
<td>46360</td>
</tr>
<tr>
<td>Heat of vaporization-kJ/kg 20°C</td>
<td>410</td>
<td>233</td>
<td>426</td>
</tr>
<tr>
<td>Explosion limit in air-vol%</td>
<td>3.4-17</td>
<td>1.0-6.0</td>
<td>2.1-9.4</td>
</tr>
<tr>
<td>Ignition temperature at 1 atm- °C</td>
<td>235</td>
<td>250</td>
<td>470</td>
</tr>
<tr>
<td>Cetane Number</td>
<td>55-60</td>
<td>40-55</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 2. Properties of Dimethyl Ether  [7, 9, 16, 17]

The viscosity of DME is much lower than that of diesel fuel. This offers an advantage in that the fuel will be easier to deliver into the engine cylinder during cold weather conditions. However, some studies have shown that the DME leaks from the injectors [18, 19]. In addition, using neat DME (100% DME as the fuel) within an engine creates some lubricating problems because of the low viscosity. Researchers are now understanding that the inherent lubricating traits of fuels used in automotive fuel injection systems, are also a very significant factor, especially when additives and alternative fuels are being considered [20-23].

The low boiling point of DME is another important advantage for its use as a fuel. Even in cold starting conditions the DME vaporizes in the cylinder yielding better atomization and hence improving combustion. The vapor pressure of DME is a concern since the fuel is a gas at atmospheric pressure and temperature. A pressurized fuel system is required so that the fuels can be mixed and injected as a liquid. This leads to other complications with fuel delivery, although the technology
to do this is similar in nature to LPG (Liquid Propane Gas) because it is also moderately pressurized to keep the fuel in a liquid state [24].

Another important aspect of combustion emissions from a compression ignition engine fueled on DME versus diesel fuel, is the reduction and elimination of particulate emissions. Particulate emissions are also commonly known as “soot” or black smoke. The oxygen content of diesel fuel blended with DME (at roughly 40 to 100 wt.%), allows for the emissions to be smokeless, as shown in the literature [7, 9, 16, 25-28]. Nabi and coworkers have shown “smokeless” engine operation from a diesel fuel mixture with an oxygen content at around 38 wt. % [25]. However, the work by Chen and coworkers confirms that even with 80 wt.% DME addition to diesel fuel, some smoke will be produced at high engine loads, although it is a small amount [28].

The heating value of DME is a concern, because it is just 60% that of diesel. This requires a larger volume of fuel to produce the same output from combustion. By altering injection amounts to the cylinders, the amount can be compensated to counteract the decreased heating value and prevent “de-rating” of engine output.

Other issues will need to be addressed in future work regarding the understanding of DME fuel properties, including, the lubricity of the fuel and the combustion mechanism. Because of the need for the fuel to be tested while in the liquid phase under pressure, further analysis outside of combustion studies may be impractical or require development of highly specialized instrumentation.

RESEARCH WITH DME IN ENGINE APPLICATIONS

Dimethyl ether is a common chemical used as a aerosol propellant [29]. The properties of DME are given in Table 2, and are compared to the diesel fuel used for the baseline testing for this experiment. DME is a liquid when contained under moderate pressure, with a vapor pressure of 5.1 bar at 20°C, and is relatively easy to handle. Over the past ten years, researchers have started to consider the use of DME as a fuel. Because the cetane number and ignition temperature are close to that of diesel fuel, DME was thought to be an excellent substitute for use in compression ignition engines. However, there were some drawbacks to using the fuel, including the reduced viscosity and lubricity of the fuel in neat form, as well as fuel compressibility effects [10].

To potentially overcome the fuel property effects of DME, as well as, reduce emissions, the experiments for this study focus on mixing dimethyl ether with diesel fuel. The initial goal is to determine the effect of the oxygen concentration on the emissions, with minimal engine modifications. In this part of the work, no changes have been made to the fuel injection timing, fuel injectors, or engine programming. Changes to the fuel system have been made to allow the fuel to be delivered to the common rail as a liquid by maintaining the DME-diesel blend at a pressure greater than 100 psi.

Over the last ten years, many researchers have begun to evaluate the performance and emissions effects of neat dimethyl ether. Sorenson and Mikkelsen [7] found that for a fixed speed and across various loads, the particulate and NOx emissions from a .273 Liter direct injection single cylinder engine fueled with neat dimethyl ether could be significantly reduced as compared to emissions with diesel fuel. In the same study, the HC and CO emissions showed little or no change. Later, Sorenson and Mikkelsen [30] further studied the HC emissions from this same engine, and found that there was an increase in the HC emissions when using neat DME, with more methane found than in a typical diesel engine exhaust, and less light hydrocarbons. With another engine, Christensen and Sorenson [31] looked at various effects on the suite of emissions when using neat DME. Of particular interest, the NOx emissions were significantly reduced when the injection timing was retarded towards Top Dead Center (TDC). However, there was an increase in the CO emissions, and little effect on the HC emissions. Other tests determined that lower injector opening pressure reduces NOx, and nozzle types did not seem to influence NOx emissions. Experiments by Kajitani and coworkers [26] also showed the effects of injection timing on reducing NOx, which had little effect on HC emissions, from a single cylinder Yanmar engine fueled with neat DME.

However, in the work completed by Hupperich and coworkers [32] with a 1.75 liter single cylinder engine for the ECE R49 13-mode test, the cumulative emissions data displayed some differing results. With the use of neat DME, HC emissions are reduced and the trends with the other emissions are similar to what had been determined with previous studies. One difference to note is the change in injection nozzle size, which may have affected the emission results by allowing for more complete combustion of all fuels tested in an effort to maintain consistent test conditions.

Recent experiments by Ikeda and coworkers [27] with a single cylinder engine using a binary fuel injection method showed similar NOx emissions between diesel fuel and 40 volume % DME mixed with diesel fuel, as injection timing was retarded. Also, HC emissions increased and smoke emissions were reduced as injection timing was retarded. In addition, comparisons were made as a function of BMEP Brake Mean Effective Pressure (BMEP). NOx was reduced, HC remained constant and smoke increased with increasing BMEP. The experiments also included % DME fractions mixed with diesel fuel up to 60 volume % addition, with comparisons made to the baseline diesel fuel. The smoke level, indicating presence of soot in the exhaust stream, showed a slight increase between 0 and 20 % DME addition, and then returned to zero for DME.
addition over 20%. NOx emissions decreased slightly, then increased slightly to the original point for diesel fuel. HC emissions increased slightly up to 45% DME addition, and increased sharply above this point [27].

Many researchers have been evaluating the performance of other oxygenates including blends of glycol ethers with diesel fuel, and have observed decreases in particulate matter emissions with increasing oxygenate concentration. Most recently, Hallgren and Heywood [33] prepared a review of the collection of work which showed that as the oxygen content of the fuel increases, the particulate matter is reduced, suggesting that this occurs regardless of chemical structure or molecular weight. However, their actual testing showed that the oxygenate structure did impact particulate emissions. Studies completed by Hess and coworkers [34] as well as by Litzinger and coworkers [35, 36] have shown that higher molecular weight glycol ethers are also effective in reducing particulate matter emissions, although to a lesser extent than monoglyme or diglyme.

Although it has been shown that glycol ethers effectively reduce particulate emissions, the fundamental mechanisms of the reduction have not been clearly identified. There has been some work in simulating the ignition and rate mechanism behavior of dimethyl ether in comparison to dimethoxymethane [37]. Also, oxidation mechanisms have been proposed for gaseous forms of DME [9, 38, 39]. More recently, the modeling of DME oxidation has proven consistent with experimental results from jet stirred reactor theory and shock-tube conditions, providing confidence in the proposed reaction mechanisms [40].

RESEARCH WITH DME IN VEHICLE APPLICATIONS

Over the years, there have been various attempts to operate vehicles on neat DME. One successful vehicle demonstration involved the Volvo B10BLE bus. Even though there were some technical issues related to the use of DME in the fuel system, the data showed that a DME fueled vehicle with an oxidation catalytic converter could meet the EURO 4 Standards [41]. While the deployment of the Volvo bus was prevented by difficulties with reliability of the fueling system, this project demonstrated chassis dynamometer emissions that were very low. More recently, successful demonstrations of vehicles have occurred in Japan, with DME powering 2 ton trucks [42].

SHUTTLE BUS CONVERSION

The objective of this research project was to demonstrate operation of a campus shuttle bus on dimethyl ether (DME). To accomplish this goal, DME was blended with diesel fuel to provide sufficient viscosity and lubricity to the fuel blend to prevent damage to the fuel injection system of the engine in the shuttle bus. This involved development of a conversion strategy. The ultimate goal of the conversion process was to build a fueling system that allowed for delivery of pressurized fuel with as little modification to the engine as possible, with relatively simple controls and a reasonably convenient re-fueling process. The system layout was based on the design developed by Chapman and Bhide [46, 47] for the same Navistar T444E diesel engine.

In the work by Chapman and Bhide, a T444E engine was converted in a laboratory test cell to operate on blends of DME and diesel fuel. A schematic of the modified fuel system used in the laboratory test cell is shown in Figure 1. The fuel system on the T444E engine had to be modified to account for the need to deliver fuel at an elevated pressure. The fuel rail in the cylinder head of the engine normally receives fuel at a pressure of about 70 psi. Fuel from this rail is then fed to the injectors. To prevent boiling of the DME in the fuel blend, fuel from the rail was kept at a pressure greater than 100 psig and recirculated through a chilled bath to keep the fuel temperature below 50°C before returning to the rail by use of a gear pump. The fuel tank consisted of a modified 60 lb capacity LPG cylinder.

Also in work by Bhide, Perez and Boehman, the viscosity of blends of DME and diesel fuel were measured [48]. The viscosity of a conventional diesel fuel drops below the ASTM specification of 1.3 cSt when the blend ratio approaches 25 wt.% DME. Thus, to protect the fuel injectors from excessive wear due to viscosity-related wear, the target DME concentration for the vehicle tests as set to a maximum of 25 wt.% DME.

---

**Figure 1. Pressurized Fueling System Developed in Laboratory Studies on DME-diesel Blends [46, 47]**
The development of the conversion strategy for the shuttle bus was subjected to a detailed failure mode effects analysis (FMEA), as prescribed by SAE J1739 [49]. A similar procedure, used by Air Products and Chemicals, Inc., is referred to as a Hazardous Operation Process Analysis, or “Hazop”. Such an analysis was performed to ensure safe operation of the converted bus. The objective of the Hazop analysis was to review the fuel handling system and fueling procedures in detail and to attempt to foresee any potential problems that could occur. The initial system design was divided arbitrarily into four nodes. This division allowed attention to be focused on a specific subsystem of the design. Each segment of the design was scrutinized for conditions of fluid flow (no flow, high flow, low flow, reverse flow), loss of containment (full and partial), pressure (high pressure and low pressure), temperature (high and low), level (high level, low level, and no level), composition, mixing, reactions, operational procedure, phase, startup, shutdown, erosion, maintenance, and fire. The completion of this analysis increased the complexity of the design considerably but was required to ensure that the vehicle met safety standards of the National Fire Protection Agency (NFPA) and the Pennsylvania Department of Transportation.

COMPONENT SELECTION

Before any hardware could be purchased a significant amount of research needed to be completed. Since DME will attack most polymers, and therefore most o-rings or seals, material compatibility was a top priority. Stainless steel tubing was used for any fuel line that would contain pure DME and was grounded to eliminate hazard from electrostatic discharge. In addition, all of the components had to be durable enough to withstand exposure to road debris and vehicle vibration.

A 33.2 gallon water capacity liquid propane (LP) tank from Manchester Tank (Lynwood, CA) for use on recreational vehicles serves as the primary fuel tank to store and deliver the blend of DME and diesel fuel. The tank size was calculated for the amount of fuel required to run one full day on the Faculty/Staff Shuttle loop at the University Park campus of the Pennsylvania State University. All valves that came installed in the tank were removed and new valves were installed to ensure material compatibility with DME. As many suppliers have various materials to select, Kalrez™, a product of Dupont-Dow Elastomer, was selected when available. To maintain tank certification, an 80% fill valve is required on the LP tank, but a modified fill valve was installed with the desired threaded connection for filling the tank with DME and diesel fuel.

A positive displacement gear pump from Liquiflo Equipment Company (Garwood, NJ) transfers diesel fuel from the existing diesel tank to the liquid propane tank. The pump was ordered to be compatible with diesel only, since under normal operation it would not be exposed to DME.

A magnetically coupled gear pump from Tuthill Pump (Concord, CA) circulates the blended fuel to the cylinder heads and then through two transmission oil coolers. Cooling the fuel is required to maintain the DME in liquid state and hence to avoid two-phase flow in the fueling system.

The fuel was filtered through two types of filters, provided by Parker Racor Filtration. The fuel was passed through a water separator and a high pressure LNG filter. Initially, the high pressure filter which had a 5 micron mesh filter element clogged repeatedly. This fouling of the filter blocked fuel flow and disrupted operation of the engine. The fouling appeared to be due to the solvent action of DME as it cleaned the fuel system and mixture fuel tank, depositing an oily residue on the filter. Eventually, through the technical assistance from Parker Racor, all fuel filtration issues were resolved.

“Fail in position” pneumatically operated ball valves were used throughout the fuel handling system. Solenoid valves delivered compressed air to actuate the ball valves. Since the solenoid valves were not compatible with road elements a cabinet was purchased to house the solenoid valves inside the driver/passenger compartment. To ensure safety and correct flow direction, pressure relief valves and check valves were installed in the fuel handling system to supplement the pneumatic ball valves.

COMPONENT INSTALLATION

The first step of the construction was to drain and modify the existing diesel tank. The tank was removed from the bus and two couplers were welded onto the tank. These couplers serve as the connection points for fuel supply and fuel return from the engine compartment. Two additional vents including vapor traps were also added to the diesel tank to prevent over pressurization of the tank and release of flammable vapors. The existing tank was then remounted on the bus. Next the liquid propane tank, here after referred to as mix tank, was mounted on the bus. This was done using the tank manufacturer’s suggested hardware and instructions. Finally, the rest of the fuel handling system hardware (pumps, valves, tubing, fittings, heat exchangers) was installed on the bus. A full diagram of the completed system is shown in Figure 2.

The standard configuration of the T444E engine is for fuel to enter from the backside of the cylinder head (side closest to the firewall) and remain in the fuel rail until needed by the injectors. This configuration, referred to as a “dead heading,” causes the fuel to be heated to engine coolant temperatures. When the engine is fueled...
Figure 2. Schematic Diagram of the DME-Diesel Fueling System
on diesel fuel this is not a problem. However, when running on a blend of DME and diesel fuel, this heat soak becomes an issue. To combat the heating of the fuel in the rail, the fuel flow path is changed from a “dead heading” arrangement to a circulation loop. Fuel enters the engine at the same place as in the original configuration and exits at the front of the engine. The fuel from each side of the engine is then combined and routed through four transmission oil coolers (referred to as “fuel coolers”) that are mounted just in front of the factory radiator and charge air cooler. This configuration allows the fuel to be cooled and fresh makeup fuel added to the loop before the mixture is re-circulated to the engine. Figure 3 shows a photograph of the fuel handling system installed under the hood of the shuttle bus.

TEST FIRING

Once construction of the fuel handling system was complete, the vehicle was test fired on diesel fuel. This allowed for a full operational check of the new system and a fully pressurized leak check. The vehicle was then operated at the Pennsylvania Transportation Institute Test Track facility to verify vehicle performance, operation of the vehicle with the converted fueling system and refueling procedures. The vehicle was fueled on a conservative mixture of 10 vol.% DME in diesel fuel for initial testing after operation problems had been addressed.

OPERATIONAL ISSUES

With the complexity of the design, several operational issues became evident once construction was completed. These problems involved formation of two-phase flow in the fuel handling system that caused rough engine operation and re-fueling difficulties.

CIRCULATION PUMP

As described above, a magnetically coupled gear pump is used to circulate fuel through the cylinder heads through fuel coolers and back to the engine. Since DME is a vapor at room temperature and pressure, the fuel must be kept pressurized to avoid formation of two-phase flow within the system. When the fuel is exposed to engine temperatures, the amount of pressure needed to keep the DME in the liquid phase increases. The vapor pressure of DME as a function of temperature is shown in Figure 5 for reference, provided by Dupont Technical Bulletin for Dymel A [50]. Since the system pressure is set, the fuel temperature must be lowered via heat exchange through the fuel coolers. Thus, if for some reason the circulation pump does not work, fuel is on.

Several scenarios for pump failure were encountered. The pump, which runs off 110 AC power provided by a DC-to-AC converter, was demanding increasing amounts of power due to: inadequate power supply, inadequate flow of fluid through the system due to blockage, jamming of the gears due to debris, thermal shutoff, and swelling of the gears.

Initially, after construction was complete, the bus was fueled on diesel fuel. During this time no problems with the pump were noticed. However, after long periods of operation, the engine ran roughly and the amount of blow-by increased, as observed by an increase in smoke emitted by the crankcase vent and a noticeable increase in odor. When the bus was fueled on DME, the current draw by the electric motor that drives the circulation pump gradually increased over time. The current demand by the pump motor would increase to the maximum rated current for the motor and trip an internal...
protection circuit in the motor. Hence, the pump would shut off.

The manufacturer was contacted and a larger motor and a motor adapter were purchased and installed. After installation the operation of the pump seemed to become worse and the motor and motor adapter were removed. It was found that the magnetic coupler from the motor shaft to the cap of the pump was destroyed. The pump, motor mate, and coupler were taken to the attention of the regional sales representative who replaced the destroyed coupler and correctly installed a new one. The pump was then reincorporated into the system.

With the motor now working correctly, it became evident that a larger power source was required. Since the pump runs on 110 AC but only 12V DC is available on the vehicle an inverter is used. To obtain the size inverter needed a recreational vehicle dealer was contacted. Once the inverter was in place the pump performed flawlessly on diesel fuel for 8 days. When the vehicle was switched to DME-diesel blends, problems again became evident.

A flow meter was installed to determine if the pump was getting adequate fluid or if it was running dry. The meter was installed in the system with as little changes as necessary to ensure a true reading. The flow was found to be in the acceptable range for the motor rating. This ruled out insufficient flow due to blockage in the system. It was postulated that the flow restriction could be on the outlet side of the pump so the filter element was examined.

The filter was found to have disintegrated at the top and bottom of the element where rubber caps were used in the manufacturing. A new filter element with a completely metal housing and a more resilient rubber was chosen and installed. The pump was then removed and the gears checked for any debris that could be preventing their rotation. Very little material was found in the gears and the pump was reassembled and reinstalled.

By the manufacturer's suggestion, a surface temperature thermocouple was installed on the skin of the pump. It was found that there was inadequate airflow to the motor of the pump so a 12V DC accessory fan was installed to direct air toward the pump. After the installation of the fan, the skin temperature was found to be in the acceptable range, thus ruling out thermal shutoff. With all of these test and modifications, problems were still evident after fueling with DME.

It was thus apparent that gear swell due to prolonged exposure to DME was the source of the problem. The manufacturer was again contacted and a new pump was obtained. Pump shutoff and rough engine operation ceased to be a problem from this point forward.

**FUEL LEVEL GAUGE**

The level of fuel in the mix tank is monitored by a float gauge inside the tank. The signal is then sent to two individual readouts. One readout is mounted above the driver's seat and the other is attached to a five-pin connector on the side of the bus during fueling.

**VEHICLE EMISSIONS TESTING**

Emission testing is done by use of on board analysis equipment owned and operated by Clean Air Inc. The equipment consists of two units, which are approx. 24 inches wide by 22 inches deep, and several laptop computers that provide interface and control ability. The equipment measures CO, CO2, NOx, particulate matter, as a function of vehicle speed and load. Fuel consumption is calculated from the CO and CO2 measurements.

**DRIVING CYCLES**

For the purpose of ensuring and verifying repeatability, a specific testing procedure was adhered to for all testing runs. All testing was performed at The Pennsylvania State Transportation Institute Test Track facility on a one-mile oval track. Four different driving cycles were used for the testing: P20, P40, Orange County, and Manhattan Cycles.

A P20 consists of starting at a specific point on the track and accelerating to 20 mph. The vehicle is held at 20 mph until the first stopping point is reached. The vehicle is then brought to a stop for seven seconds and the driver then accelerates back to 20 mph. There are 8 stops performed in one loop. A complete P20 consists of 2 loops.

A P40 consists of starting again at a specific point on the track and accelerating to 40 mph. The vehicle's speed is then held at 40 mph until the first stopping point is reached. Then vehicle is then brought to a stop for seven seconds and the driver accelerates back to 40 mph. The first stop is at half-track the second is at the original starting point. A complete P40 consists of 2 loops.

An Orange County cycle is a reproduction of a typical bus cycle in Orange County California. The cycle is displayed on a laptop computer running Driver's Aid software® provided by Clean Air Technologies International, Inc. The software also displays the vehicle's speed and the driver matches the speed trace to the prescribed cycle being displayed. The Orange County cycle is broken into two parts to minimize datafile length and can be seen in Figures 6 and 7. A complete Orange County set consists of four complete cycles.

A Manhattan cycle is a reproduction of a typical bus cycle in Manhattan, New York. Again the cycle is...
displayed by the Driver’s Aid® software and the driver matches the vehicle speed trace to the prescribed cycle. A complete Manhattan set consists of four cycles. The Manhattan cycle is shown in Figure 8.

VEHICLE TESTING PROCEDURES

Since the Driver’s Aid® software display is difficult to see during the day the Orange County and Manhattan cycles are run at night and the P20 and P40 cycles are run during the day.

A typical day of testing begins with warming up the equipment. This process takes about one hour to complete during which the vehicle is at idle. At the completion of this time period, several test laps are made to check the equipment status. If redundant instruments indicate conflicting measurements the laps are aborted and the equipment checked. Once consistent results are obtained, actual testing begins.

Using the onboard equipment as a live monitoring system, three laps at 30 mph are made to burn off particulates that may have accumulated during the warm-up time. If particulate levels continue to trend downward in the last lap, additional laps are done until stable readings are seen.

At the completion of these “burn off” laps, the vehicle is brought to a stop at the P20 starting point. A period of 90 seconds is allowed to transpire allowing for the equipment to be reset. A complete P20 is then run followed by a complete P40, this pattern is then continued alternating back and forth with 90 seconds between each complete set.

An identical pattern is followed for the Orange County and Manhattan cycles. After equipment warm up, “burn off” laps are run, pausing 90 seconds then beginning the desired cycle. The testing pattern consists of running blocks of Manhattan cycles before switching to Orange County cycles.

RESULTS AND DISCUSSION

Testing was performed for 3 different fuel mixtures. The first test served as a baseline run and the vehicle was fueled on pure diesel. The second test was a fuel mixture of 14% by volume DME and diesel; the third test was 25% by volume DME and diesel. While emissions data were monitored for the P20, P40, Orange County and Manhattan test cycles, only data from the Manhattan cycle is presented here.

Total particulate results for all three-fuel compositions for the Manhattan cycle can be seen in Figure 9. As is expected with the addition of an oxygenate as the amount of DME in the fuel mixture increases the measured particulate level drops. For 14 vol.% DME, the PM emissions drop by 60%. For 25 vo.% DME, the PM emissions drop by 80%.
Similar results to those seen by Chapman et al. [47] were seen for hydrocarbons, NOx, and CO in tests with the laboratory engine when operated on DME blends. For a Manhattan cycle, the emissions of hydrocarbons, NOx, and CO are shown in Figures 10, 11, and 12, respectively.

While the emissions of particulate were substantially reduced, emissions of hydrocarbons, NOx, and CO were significantly increased with DME addition. While these trends were also observed in the laboratory engine [46,47], they point to a need to optimize the injection characteristics of the fuel blend and to the need to add an oxidation catalyst to a vehicle fueled in this manner. A similar observation was made in the Volvo bus demonstration, that addition of an oxidation catalyst to a DME-fueled vehicle provided very low emissions [41].

TECHNICAL ISSUES WITH THE UTILIZATION OF DME AS A FUEL

Much of the early work with DME utilization has been done at the AVL labs in Graz, Austria. The testing showed that DME reduces particulate emissions to zero, and also showed that the typical diesel fuel injection system does not tolerate DME [18]. The present work
shows that with DME-diesel blends, substantial particulate emissions reduction can be obtained while preserving the integrity of the fuel injection system. Figure 13 shows a picture of the DME-Fueled Shuttle bus in operation at Penn State.

Figure 13. DME-Fueled Shuttle Bus Operating on the Faculty/Staff Loop at the University Park campus of the Pennsylvania State University, August, 2002.

Overall, this demonstration project has shown that a vehicle can be successfully operated on DME, but at restricted DME blend levels below 25 wt.% (30 vol.%). The emissions and performance data presented here are restricted to 21 wt.% (25 vol.%) DME addition. There are remaining concerns and challenges, particularly because of the low viscosity of DME. Below is a combined list of concerns not only from the present work, but also from other groups who are now working through the fuel property, design and technology issues for utilization of DME:

- DME was leaking past clearances on the injectors and seals. This caused the need for the camshaft housing and crankcase of the engine to be vented [18].
- At high vapor pressure, the DME was cavitating, which caused difficulties in maintaining stable fuel injection [18].
- While DME is more compressible than diesel fuel, it was found that the compressibility changed with temperature and pressure. Therefore, this made it difficult to inject the maximum fuel quantity at high temperatures and during full load operation using traditional diesel equipment [18].
- DME chemically attacked some seals [18].
- Not much effort has been put towards understanding the environmental impacts of the compound itself or the emissions from the fuel combustion, as compared to other fuels [18].
- A larger fuel tank will be required, as compared to diesel fuel, because of the lower density and heating value of DME [19].
- Since the vapor pressure of DME is low, the fuel vaporizes immediately upon injection into the cylinder. This may or may not be an issue, but further study may confirm how the combustion reaction takes place after the vaporization occurs [19].
- Injection via some fuel pumps causes uncontrollable pressure waves in the entire system [43].
- Predictability of spray behavior and characteristics is important in repeatability of combustion [44].
- Turbulence within the cylinder is important for mixing of the fuel, which in turn reduces emissions [45].

CONCLUSIONS

This work has demonstrated that a conventional diesel vehicle can be converted to operate on blends of DME and diesel fuel. Significant reductions in particulate emissions were observed when the converted shuttle bus was operated on DME – diesel blends, while there were increases in unburned hydrocarbons, NOx and CO. By blending DME with diesel fuel, an acceptable viscosity and lubricity could be obtained in the fuel mixture to provide reliable operation of the fuel injection system. Nonetheless, operation of the vehicle was not without challenges during the conversion and debugging phases, as is documented in this paper. In addition, the vehicle could not meet peak accelerations required during the Manhattan driving cycle, indicating that the engine was de-rated at higher DME concentrations.

DME holds great promise as a fuel for the future. But, many technical challenges remain to be overcome before neat DME-fueled vehicles can be a commercially viable option. Among them are the low viscosity, low lubricity and material incompatibilities.

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EMISSION CHARACTERISTICS OF A NAVISTAR 7.3L TURBODIESEL
OPERATED WITH BLENDS OF DIMETHYL ETHER (DME) AND DIESEL
FUEL

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by
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Several oxygenates have been proposed and tested for use with diesel fuel as a means of reducing exhaust emissions. This paper examines dimethyl ether (DME), which is a potential ultra clean fuel, in blends with diesel fuel. Modest additions of DME into diesel fuel (2 wt.% oxygen) showed reductions in particulate matter emissions, but the previously reported data by the author from a multicylinder Navistar 7.3L Turbodiesel engine were scattered. In this study, experiments were performed on a multi-cylinder Navistar 7.3L Turbodiesel engine to repeatably confirm and extend the observations from the earlier studies. This is an important step in not only showing that the fuel does perform well in an engine with minor modifications to the fuel system, but also in showing that DME can give consistent, significant results in lowering emissions. The Dimethyl ether and diesel fuel were blended to achieve a net addition of 5 and 10 wt. % oxygen in the blended fuel. The data confirms that the addition of DME can reduce the particulate emissions from a compression ignition engine. However, the NO_x emissions were not favorable for all conditions. It is believed that through further modification of injection timing, NO_x emissions can be effectively reduced. Additionally, pressure trace analyses showed that the fuel affects the ignition delay of the combustion process, and that optimization of the engine controls would be needed to further improve performance and reduce exhaust emissions.
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Chapter 1

Introduction

1.1 Background

In 1892, it was the intention of Rudolf Diesel to provide a new type of internal combustion engine operating with higher efficiency than conventional steam and Otto cycle engines, and that could be operated on many types of fuels [1]. The engine cycle developed by Diesel involves injection of fuel into a volume of air heated by compression. Over the course of history, the durable and efficient diesel engine has replaced other less efficient modes of power production, including steam engines in the railroad industry.

As our world and our environment have changed, concern about the exhaust emissions from internal combustion engines has driven improvements in both types of widely used engines, spark ignition and compression ignition. These improvements include many types of aftertreatment technologies to combat the specific emissions from each type of engine. For the spark ignition engine, aftertreatment technologies include three-way catalysts, utilizing electronics for fuel and air control, and exhaust gas recirculation. Conversely, the emissions from diesel engines have been managed, up until now, primarily by subtle improvements in the fuel injection timing, turbocharging and
intercooling, and higher fuel injection pressure, without the need for significant exhaust aftertreatment [2]. The challenges which the automotive industry now faces are the more stringent emissions standards promulgated by governing bodies worldwide, and increasing costs of petroleum products. Specifically within the diesel engine sector, improvements in emissions are necessary for future sales, and improvements in efficiency have been made necessary by the rising cost of diesel fuel.

In the U.S., future regulations that take effect in 2004 and 2007 will require diesel engine and vehicle manufacturers to review all aspects of the vehicle system design [3, 4]. Demand for cleaner burning diesel fuels is growing worldwide, as governmental regulations make emissions reductions necessary. To achieve substantial reductions in emissions, it is thought that reformulated diesel fuels will play an important role [5]. The reformulation of diesel fuels could include lowering the sulfur content, lowering the aromatic content, or potentially adding the oxygen to the fuel [6].

One solution for reducing emissions from future and current diesel vehicles is to modify the fuel, without modifying the engine hardware. It has been shown that many oxygenates are effective at reducing particulate emissions from diesel engines [7-27]. Therefore, much research has focused on the screening of oxygenated fuel additives, including alcohols, esters, and ethers [6, 28]. Of particular interest are the glycol ethers, which have been shown to be very effective as blends and as neat fuel. This study focuses on the use of a short chain ether, dimethyl ether (DME).
1.2 Statement of Purpose

The present study involves the use of a light-medium duty diesel engine, for use in class 2 through 8 trucks and buses as suggested by Hower and co-workers, to study the ability of using blends of diesel and dimethyl ether with minimal adjustments to the engine hardware [29]. Various studies with neat dimethyl ether have shown that significant emissions reductions are possible [30]. However, when neat DME was used by Sorenson and co-workers, the durability of the fuel system was significantly reduced. Therefore, this suggests the need for a lubricity improver in the fuel. In this study, it is hypothesized that diesel fuel can provide the lubricating qualities needed, while using DME blended with diesel in a quantity that provides an emissions reduction.

Very little information and data exists regarding the use of dimethyl ether and diesel fuel blended together. Since dimethyl ether has been shown to significantly reduce the durability of fuel system hardware [31], blending DME with diesel fuel will improve the lubricating quality of the fluid. The purpose of this study is to perform initial assessments of emissions reductions with the use of the fuel blends.

1.3 Research Questions and Hypothesis

The motivation for this work is the potential of using DME blended with diesel fuel to reduce emissions, specifically particulate emissions. Since it has been shown that
neat DME eliminates particulate emissions, mixtures of DME and diesel are expected to yield a decrease in particulate emissions. However, what is not known is if this decrease will follow the same trend as observed with other oxygenates mixed with diesel with similar oxygen content.

It is hypothesized that a diesel engine can be operated with modest blends of DME-diesel fuel, with minimal modification to the engine and that diesel fuel can provide the lubricating qualities needed, while DME provides an emissions reduction. The hypothesis is being tested by a three step approach: develop an engine test facility, modify a fuel system to operate with DME-diesel fuel blends, and collect data with regard to total particulate mass emissions. Additional emissions and engine data will be collected to complement and explain the particulate emissions reductions. Some researchers have demonstrated that there is a relationship between oxygen addition to diesel fuel, particulate emission reductions, and carbon monoxide increases [28]. Also, because of the physical properties of DME, which is a liquified gas with high vapor pressure, it is important to observe the temperature and pressure of the fuel as it passes through the engine. Observations of basic engine operation will be discussed in the results. Pressure history traces of the combustion process will be used to understand the changes occurring in the ignition delay, start of combustion and rate of heat release between the diesel and DME-diesel fuel blends.
1.4 Limitations and Assumptions

The following limitations were recognized and assumptions were formulated about the experiment in order to proceed with the research. The purpose of acknowledging these limitations is to establish factors which could be influencing the results and support conclusions from the data. These limitations and assumptions will be discussed later as to how they relate to the results.

- The fuel is a homogenous mixture going into the engine.

- If the DME in the fuel rail is below 50°C, DME will be in the liquid state because the pressure will overcome the vapor pressure of DME.

- The engine will be able to operate with the fuel blend, while reducing the durability of the engine in an unknown way and amount.

- The engine operates per the factory calibration of the electronic controls for the engine. No changes have been made to engine control algorithms or parameters.

- Limitations in the comparisons that can be made and the conclusions that can be drawn are the result of the variability of engine operation as the engine responds to the differences in the physical properties of the fuel.
Chapter 2

Literature Review

2.1 Background

As with any new fuel being developed for use in the transportation industry, a comprehensive study of various aspects and impacts of the fuel are important to review and analyze before moving ahead with implementation. The most important aspect, which is the focus of this research, involves the benefits associated with using the fuel in a compression ignition engine. After confirming its potential use, inquiry shifts to other important aspects such as production cost of the fuel, health concerns, and long term use of the fuel.

The motivation for researching new fuels stems from several factors. Primarily, governments worldwide, including the United States, are setting new standards for emissions from new engines and vehicles. This is being done for the benefit of human health and the environment, via the reduction of particulate matter and smog-causing NOx [32]. The final ruling in December 2000 from the U.S. EPA sets the new 2007 standards for heavy duty diesel engines to the following:

- Particulate Matter (PM) emissions to 0.01 grams per brake-horsepower-hour (g/bhp-hr) to take full effect in 2007, and
• NO\textsubscript{x} and non-methane hydrocarbons (NMHC) to 0.20 g/bhp-hr and 0.14 g/bhp-hr, respectively. These standards will be phased in after 2007 [33].

This is shown in Figure 2–1 tracing the history of the tightening emission standards.

<table>
<thead>
<tr>
<th>Emissions Type</th>
<th>Current Regulation</th>
<th>2004 (gm/bhp-hr)</th>
<th>2007(gm/bhp-hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NO\textsubscript{x}</td>
<td>4</td>
<td>.2</td>
<td></td>
</tr>
<tr>
<td>HC</td>
<td>1.3</td>
<td>.14 (NMHC)</td>
<td></td>
</tr>
<tr>
<td>NO\textsubscript{x} and HC</td>
<td>2.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PM</td>
<td>.1</td>
<td>.01</td>
<td></td>
</tr>
</tbody>
</table>

*Required on-board diagnostics (OBD) systems for vehicles between 8500 and 14000 lbs to be phased-in, beginning in 2005

NO\textsubscript{x}- Nitrogen Oxides; HC- Hydrocarbons; PM- Particulate Matter

*Figure 2–1: U.S. Heavy-Duty Diesel Engine Emission Standards [32, 33]*

Additionally, a new fuel type that can be made from several fossil fuel sources would benefit the energy security of the U.S. [34]. Finally, it is necessary to confirm the ability of an engine to operate on a fuel in a manner similar to the operation on the fuel for which it was designed to operate. This means that development projects are necessary to experimentally prove concepts and long term operation.

DME is a compound that has been targeted for future use as a fuel, in several countries around the world [35-37]. Motivation to use DME exists for several reasons.
There has been confirmation that the fuel yields low particulate emissions and possibly lower NO\textsubscript{x} emissions [30, 38, 39]. In addition, DME can be made from a variety of feedstocks, which can support the use of alternative energy resources [40, 41].

The following review of the literature will cover a summary of the testing completed to examine the potential use of DME as a fuel. Many technical challenges have been discovered through this testing, giving light to new approaches to using DME as a fuel. Researchers have begun to study the fundamental nature of the combustion process, and to understand the mechanisms of DME combustion [42, 43]. Production processes, costs and health concerns will also be covered.

2.2 Emissions from Diesel Engines

Diesel engines have been traditionally high emitters of NO\textsubscript{x} and particulate matter. Other pollutants include carbon monoxide (CO) and unburned hydrocarbons (HC), which are typically very low for a diesel engine since the air--fuel mixture is lean of stoichiometric [44, 45]. The following sections will focus on the two major pollutants from a diesel engine NO\textsubscript{x} and PM.

2.2.1 NO\textsubscript{x} Formation

Diesel engines produce significant amounts of NO\textsubscript{x}, depending on the load conditions of the engine. Of the 100 to 2000 ppm which is produced, the majority of this is nitric oxide (NO). There are several mechanisms for the formation of NO\textsubscript{x}, however,
in the case of the diesel engine, the most significant mechanism is the extended Zeldovich mechanism, also known as the thermal mechanism. The extended Zeldovich mechanism is shown in equations 2.1, 2.2, and 2.3:

\[
\begin{align*}
O + N_2 &= NO + N \\
N + O_2 &= NO + O \\
N + OH &= NO + H
\end{align*}
\]

(NO\textsubscript{x} formation is strongly dependent on temperature [44]. It is typically formed during the first 20 degrees of crank angle rotation, after the start of combustion. Therefore, emission reduction strategies target this time period, and aim to reduce combustion temperatures [1].

### 2.2.2 Particulate Formation

Particulate matter can be defined as the combination of soot, condensed hydrocarbons, sulfates, oil, and water that attach to agglomerated soot particles in the exhaust stream [46]. Detailed modeling and experiments have been conducted to try to understand the soot formation process in different types of flame structures and in flames of various fuels [47]. It has been established that several classes of higher hydrocarbons,
namely polycyclic aromatic hydrocarbons (PAH) and acetylene (C$_2$H$_2$), are important precursors to soot formation and are created in fuel-rich conditions [48]. From the initial formation of a benzene ring, the soot particle increases in size through the addition of acetylene [48].

### 2.3 Emission Control Techniques

Emission control strategies can be divided into three categories: pre-combustion, in-cylinder, and post-combustion [2, 49]. A list of these approaches can be found below in Figure 2–2 [50]. For many years, diesel engines have escaped the necessity of post-combustion controls because modifications which could be made to fuel or to the engine combustion and performance to meet new emissions regulations. Also, an important requirement for implementation of many post-combustion technologies is the reduction of the sulfur content of the fuel. Since diesel fuel properties will be discussed later in this section, the following discussion will focus on the in-cylinder and post–combustion strategies of controlling particulate and NO$_x$ emissions.
**Figure 2–2**: Emission control strategies for diesel engines [50]

- **Pre-Combustion (Fuel Modification)**
  - **Particulates**
    - Decrease Sulfur
    - Increase O\textsubscript{2} Content
    - Decrease Aromatics
  - **NO\textsubscript{x}**
    - Increase Cetane No.
    - Add Water

- **In-Cylinder Combustion**
  - **Particulates**
    - Fuel Atomization
    - Common Rail Injection
    - Injection Timing
    - Rate Shaping
    - Split Injection
  - **NO\textsubscript{x}**
    - EGR (Exhaust Gas Recirculation)
    - Water Injection
    - Injection Timing
    - Rate Shaping
    - Split Injection

- **Post-Combustion**
  - **Particulates**
    - Particulate Trap
    - Particulate Oxidizer
  - **NO\textsubscript{x}**
    - SCR
    - SNCR (Thermal & Non-Thermal)
    - SNR

SCR- Selective Catalytic Reduction
SNCR- Selective Non-Catalytic Reduction
2.3.1 Particulate Emissions Control

The following sections cover the in-cylinder and post-combustion methods of controlling particulate emissions.

2.3.1.1 In-Cylinder Combustion

In general, the methods used to control or modify the particulates generated during combustion modify the fuel atomization, injection pressure, injection timing, and fuel injection rate. The goal of providing better atomization and controlling the heat release rate is to allow soot particles to be oxidized through improved air entrainment and utilization [51]. However, this must be carefully determined through design and testing. In changing the heat release rate, overall bulk peak temperatures in the cylinder could be reduced, reducing the rate of reactions and increasing time for soot oxidation [44, 52].

2.3.1.2 Post-Combustion

Two general methods exist to control particulate emissions in post combustion. First, a particulate filter can be used. This is typically a ceramic wall flow filter with alternative plugged channels, as seen in Figure 2-3. As accumulation increases, heat is supplied to incinerate the carbon. Second, an oxidation catalyst can be used. In this method, flow is not obstructed. It is made from a monolithic substrate (cordierite; Mg₂Al₄Si₅O₁₈) with a precious metal washcoat. However, this method only oxidizes the
soluble organic fraction (SOF) of the particulate, but does not address the soot portion (insoluble) [2, 53].

---

**Figure 2–3**: Diesel Particulate Filter [53]

---

### 2.3.2 NO\(_x\) Emission Reduction

The following sections cover the in-cylinder and post-combustion methods of controlling NO\(_x\) emissions.
2.3.2.1 In-Cylinder Combustion

There are several techniques which can aid in reducing NO$_x$ emissions. These include exhaust gas recirculation (EGR), injection timing control, and fuel injection rate shaping. The goal of these three techniques is to reduce the peak flame temperature which leads to thermal NO$_x$ formation. However, these techniques also have drawbacks. In altering the fuel injection timing to the cylinder, power decreases and fuel consumption increases. Also, reducing the peak flame temperature causes an increase in the level of soot emissions, due to the reduced rate of reactions, increasing soot formation [44, 52].

2.3.2.2 Post Combustion

The general post-combustion method which exists for controlling NO$_x$ involves some type of catalytic or chemical reaction. As seen in Figure 4–4 there is Selective Catalytic Reduction (SCR), and Selective Non-Catalytic Reduction (SNCR), both thermal and non-thermal [2]. The method that is most widely talked about for use is the SCR method, also most commonly known as DeNO$_x$ Catalysis. An example of this is shown in Figure 4–4. In this method, a zeolite catalyst is used to adsorb NO$_x$ molecules. A hydrocarbon reagent is injected into the exhaust stream to chemically reduce the NO$_x$. The typical reagent used is the fuel itself, which reduces the efficiency of the vehicle. It should be noted that this process works in a small temperature band (200-250 °C for
precious metal zeolites and 400-450 °C for base metal zeolites), and for vehicles the NO\textsubscript{x} reduction is low and selectivity is poor [53]. Another method of selective catalytic reduction uses ammonia or urea as the reductant [54]. Testing has been completed using SCR, EGR, and intensive engine mapping which resulted in dramatic reduction of NO\textsubscript{x} emissions [55].

---

*Figure 2–4: DeNO\textsubscript{x} Catalyst [53]*
2.4 Compression Ignition Engines

The phrase “compression ignition” refers to the engine classification based on method of fuel ignition. The compression ignition (CI) process is used in conventional diesel engines and in gas engines by pilot injection of fuel oil. In this type of engine, air is inducted into the cylinder, and compressed. Just before the combustion process is to start, the fuel is injected into the engine. For a given engine speed, the air flow is essentially constant, and engine load can be adjusted by increasing the amount of fuel injected for each cycle. There are several variations on the type of CI engine including the working cycle (2 or 4 stroke), method of air preparation (naturally aspirated or turbocharged), and method of fueling (indirect or direct injection) [44].

For a typical 4-stroke CI engine, the compression ratio is between 12 and 24, and depends on whether the intake air is naturally aspirated or turbocharged. On the intake stroke, air is inducted into the cylinder as the piston is moving down. Then, as the piston moves up, the air is compressed. A few degrees before the piston reaches the top of its travel, fuel is injected into the cylinder. As the fuel is injected, the combustion process begins, pushing down on the piston. This is the point at which work is performed in the system through diffusion combustion and release of the energy stored in the fuel. The final stroke of the process is for the exhaust valve to open as the piston moves to the top of the cylinder a second time, pushing the products from the combustion process out of the cylinder [53]. A representation of this process is shown in Figure 2-5 below.
Air is inducted into the cylinder at atmospheric pressure and compressed to about 4 MPa (600 lb/in²) at a temperature of 800 K (1000 °F). A few degrees before Top Dead Center (TDC) of the piston, the fuel is injected into the cylinder. The fuel is atomized as it passes through the injector nozzle ports, and entrains air into the spray plume. As the liquid fuel evaporates, it mixes with the air into proportions which are combustible. Since the air temperature and pressure are above the ignition point of the fuel, there is a short delay period, and then spontaneous ignition of parts of the mixture occurs. This spontaneous ignition can be observed as an increase of pressure in the cylinder [44]. Because the compression ignition engine depends upon spontaneous ignition of the fuel,
a high cetane number fuel is required. The cetane number ensures that the fuel will autoignite when required by the location in the engine cycle.

Combustion in a diesel engine is defined as rapid oxidation generating heat and light. The combustion occurs in a flame mode, encompassing two types of flames: pre-mixed and diffusion flames. In a pre-mixed flame, the fuel and oxidizer are mixed before any rapid chemical reaction occurs. In a diffusion flame, the fuel and oxidizer are separated, and then reaction occurs at the interface between them, as mixing and reaction take place. Diffusion implies that the molecules are diffusing together towards the flame, the fuel and oxidizer moving in from opposite sides. While this molecular diffusion is occurring on a microscopic level to finish the combustion process, turbulent convection in the cylinder mixes the fuel together on a macroscopic basis [56].

Other methods to describe the 4-stroke cycle include using a pressure vs. volume diagram, and a pressure vs. crank angle diagram. The pressure vs. volume diagram shows the relationship inside the combustion chamber during the cycle. This can be seen in Figure 2–6.
The pressure vs. crank angle diagram shows the same process cycle, as a function of the degree location on the crank. From this, the peak pressure location can be determined. This can be seen in Figure 2–7.

*Figure 2–6: Pressure vs. Volume Diagram for a Four Stroke Engine [53]*
2.5 Diesel Fuel Components and Properties: Emission Effect

In 1990, Morris and Wallace of Ethyl Petroleum Additives Limited presented a diagram illustrating the effect of diesel fuel properties on emissions and overall fuel system and vehicle performance, as indicated from their entire research experience [57]. This diagram is found in Figure 2–8. Six of the ten properties directly contribute to emissions: sulfur, aromatics, density, distillation, cetane number, and viscosity. The following discussion will place emphasis on these properties.
Figure 2–8: Diesel Fuel Properties and their significance [57]
2.5.1 Diesel Fuel Sulfur Content

It has been widely established in the literature that fuel sulfur is the most important contributing factor to brake specific particulate emissions [58-60]. Diesel particulates are composed of four different constituents: the insoluble organic fraction, the soluble organic fraction, lubricants oils, and sulfates with bound water [61]. Specifically, the sulfur has been shown to contribute to the soluble portion of the particulate [58]. When there has been an increase in the sulfur content of the fuel, the engine out particulate emissions have increased, regardless of engine type, emissions levels, and combustion systems [58]. Therefore, lowering fuel sulfur will lower engine out emissions [60].

However, reducing fuel sulfur may have additional positive and negative benefits. The primary motivation for reducing fuel sulfur currently is for manufacturers to be able to implement aggressive exhaust aftertreatment technologies [60]. A potentially negative impact of the reduction of sulfur is the reduction of the lubricating quality of the fuel, known as lubricity [62, 63]. The impact of this lower lubricity may be observed in the fuel injection components and system, but it is not clear how significant this impact is [64-66].

2.5.2 Diesel Fuel Aromatic Content

The effects of aromatics, both as total aromatics (as a combination of mono-, di-, tri-, and polyaromatics) and polyaromatics, is unclear. Schmidt and coworkers showed in
their review of literature that some researchers have found no clear outcomes of the influence of aromatics on particulates and NO<sub>x</sub>, where others have found only small reductions in these emissions [60]. Lee and coworkers concluded from their series of experiments across a large platform of engines, that there is a NO<sub>x</sub> reduction benefit with lower aromatics [59]. They hypothesized that this is true via the reduction in flame temperature affecting the kinetics of the reaction by lowering the concentration of oxygen radicals. Ryan and coworkers found in their experiments on a Caterpillar 3176 engine that there was some correlation between NO<sub>x</sub> and HC emissions, and aromatic content of the fuel [67].

### 2.5.3 Diesel Fuel Density

Density is commonly used to describe the energy content of the fuel [49]. Typical diesel engine equipment meters fuel on a volume basis. Therefore, if there is a change in density, there is also a change in power output from the engine for a given throttle position [49]. Ryan and coworkers found in their set of experiments, with a Caterpillar 3176B engine, no obvious relationship between density and emissions [67]. However, Lee and coworkers have found that there is a relationship between density and emissions [59]. They hypothesize that the influence arises because of physical interactions within the fuel injection system, specifically changes in the dynamic timing and mass flow rate. The change in density is said to affect the injection spray dispersion angle and penetration, also referred to as liquid length [68, 69]. Lee and coworkers also describe that the effect of density on emissions seemed to change as the engine emissions
equipment changed to meet reduced emissions standards. The modern engines were utilizing higher injection pressures and sophisticated electronic control of injection timing. This could explain why Ryan and coworkers did not find a correlation, as the Caterpillar 3176B has these features.

2.5.4 Diesel Fuel Volatility

Volutility is the measure of the boiling point of a fuel. It characterizes the ability of a fuel to combust, and influences many other properties of the fuel, including density, autoignition temperature, flash point, viscosity, and cetane number [49]. Lee and coworkers determined that the T90/T95 portion had little effect on emissions. However, Schmidt and coworkers found from the literature that reductions in T90 did reduce the particulate matter and HC emissions [60]. Canaan, Dec, and coworkers showed that T50 affects emissions because there is a change in the spray length penetration into the cylinder, and that the chemical processes dominate the physical ones with respect to autoignition [70]. They also declared that the liquid-length penetration is also affected by other fuel properties, such as viscosity and density (specific gravity), which are influential to the atomization and jet break-up process. However, they also suggest that the liquid length penetration is primarily dependent on the evaporation process and the fuel properties which characterize this, such as vapor pressure and boiling point.

The relationship of volatility, or the boiling point, to density, is established in several ASTM standards, D 86, D976, and D4737 [71-73]. These standards present a correlation chart of T50 volatility and density to an index number for cetane [71-73].
Based on these charts, it seems that T50 has a stronger correlation to density than the other boiling points.

2.5.5 Diesel Fuel Cetane Number

In general cetane number refers to the quality of the fuel based in comparison to a standard that has the same ignition delay as the fuel under test, in terms of degrees of rotation of the crankshaft. Simply, it is a measure of ignitability of the fuel. A more specific definition of cetane can be found in ASTM D4737, and is a function of the boiling point (volatility) at T10, T50, and T90, and the density of the fuel at 15 °C [49].

For a compression ignition engine, it has been traditionally thought that the higher the cetane number, the more complete the combustion and the cleaner the exhaust [49]. This has been confirmed through several studies. Ryan and coworkers found no obvious trend between particulate emissions and cetane number [67]. They hypothesized this was due to the reduced pre-mixed burn phase of combustion. However, Lee and coworkers found that changes in particulate emissions are very engine specific. Also, they determined that higher cetane had a beneficial effect on HC emissions, but this was also engine emission level dependent [59]. Since cetane number affects the autoignition quality of the fuel, the shortened ignition delay of the fuel is reducing the time and size of the pre-mixed burn of the fuel. Schmidt and coworkers also confirmed Lee and coworkers findings that higher cetane also lowers NOx [59., 60].
2.5.6 Diesel Fuel Viscosity

A fluid is defined as a substance that continuously deforms under the action of a shear stress. If there is no shear stress, then there is no deformation. A term used to classify fluids is the viscosity, which is described as the resistance to deformation due to a shearing force. A Newtonian fluid is one in which the shear stress is directly proportional to the rate of deformation. A non-Newtonian fluid is a fluid in which the shear stress is not directly proportional to the rate of deformation [74]. The ratio of the absolute viscosity, $\mu$, and density, $\rho$, of a fluid is commonly referred to as kinematic viscosity, $\nu$ ($\nu = \mu/\rho$).

From a physical effects standpoint, the viscosity of the fluid affects the mass flow rate, injector spray cone angle, fuel distribution and penetration, droplet size of the fuel, and optimum timing of injection. Lower viscosity is characterized as providing better atomization. However, if the viscosity is too low, this could result in leakage through injection equipment clearances. If the viscosity is too high, excessive heat generation in injection equipment could result, because of viscous shear in the clearances between injection plungers and cylinders [49].

2.6 Properties of Dimethyl Ether

Compounds in which two hydrocarbon groups are bonded to one oxygen, represented as R-O-R’, are called ethers. The organic groups bounded to the ether may be alkyl, aryl, or vinylic, and the oxygen can either be an open chain or ring configuration
Ethers commonly observed in long chain structures are referred to as linear ethers. As compared to alkanes of similar structure, for example if the CH₂ group replaced the O atom, the boiling points of ethers are higher [75]. This class of oxygenated compounds have high cetane numbers and excellent cold flow properties [76].

Simply stated, dimethyl ether is an ether with two methyl groups on each side of an oxygen atom. Today, it is predominantly used as an aerosol propellant due to it not being harmful to the ozone layer in contrast to other aerosols used previously [41]. Also, it is virtually non-toxic and is easily degraded in the upper atmosphere. It can be represented by the symbol: CH₃-O-CH₃. The physical properties of DME are shown in Table 2–1 along with some comparison fuels [30, 38, 77, 78]:

Table 2–1: Properties of Dimethyl Ether [30, 38, 77, 78]

<table>
<thead>
<tr>
<th>Property</th>
<th>DME</th>
<th>Diesel (C₁₀H₁₈.₇)</th>
<th>Propane (C₃H₈)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical Formula</td>
<td>C₂H₆O</td>
<td>C₁₀.₈ H₁₈.₇</td>
<td>C₃₃H₈</td>
</tr>
<tr>
<td>Mole Weight</td>
<td>46.07</td>
<td>148.6</td>
<td>44.11</td>
</tr>
<tr>
<td>Critical Temperature- °C</td>
<td>127</td>
<td>-</td>
<td>95.6</td>
</tr>
<tr>
<td>Boiling Point- °C</td>
<td>-24.9</td>
<td>71-193</td>
<td>-42.1</td>
</tr>
<tr>
<td>Vapor Pressure at 20 °C-kg/m²</td>
<td>5.1</td>
<td>&lt;0.01</td>
<td>8.4</td>
</tr>
<tr>
<td>Critical Pressure-bar</td>
<td>53.7</td>
<td>-</td>
<td>43</td>
</tr>
<tr>
<td>Liquid Viscosity- cP</td>
<td>.15</td>
<td>2-4</td>
<td>.10</td>
</tr>
<tr>
<td>Liquid Density at 20 °C-kg/m³</td>
<td>668</td>
<td>800-840</td>
<td>501</td>
</tr>
<tr>
<td>Bulk Modulus (N/m²)</td>
<td>6.37E+08</td>
<td>1.49E+09</td>
<td></td>
</tr>
<tr>
<td>Specific Density,gas</td>
<td>1.59</td>
<td>-</td>
<td>1.52</td>
</tr>
<tr>
<td>Solubility in H2O at 20 °C g/l</td>
<td>70</td>
<td>Negligible</td>
<td>.12</td>
</tr>
<tr>
<td>Lower Heating Value- kJ/kg</td>
<td>28430</td>
<td>42500</td>
<td>46360</td>
</tr>
<tr>
<td>Heat of vaporization- kJ/kg 20°C</td>
<td>410</td>
<td>233</td>
<td>426</td>
</tr>
<tr>
<td>Explosion limit in air- vol%</td>
<td>3.4-17</td>
<td>1.0-6.0</td>
<td>2.1-9.4</td>
</tr>
<tr>
<td>Ignition temperature at 1 atm- °C</td>
<td>235</td>
<td>250</td>
<td>470</td>
</tr>
<tr>
<td>Cetane Number</td>
<td>55-60</td>
<td>40-55</td>
<td>-</td>
</tr>
</tbody>
</table>
The properties that are significant to the use of DME as fuel for combustion are cetane number, boiling point, and ignition temperature. However, the properties of concern are viscosity, heating value, and vapor pressure.

To gain an appreciation for the advantages and disadvantages of dimethyl ether, one must understand the properties required for diesel fuels in compression ignition engines. *Table 2–2* below provides some insight [49]:

*Table 2–2: Fuel Property Description* [49]

<table>
<thead>
<tr>
<th>Property</th>
<th>Rationale</th>
</tr>
</thead>
<tbody>
<tr>
<td>Volatility</td>
<td>High volatility aids cold starting and incomplete combustion.</td>
</tr>
<tr>
<td>Flash Point</td>
<td>Higher flash point provides safety in handling and storage.  Adam j26db.</td>
</tr>
<tr>
<td>Cetane Number</td>
<td>Measure of ignitability. The higher the cetane number, the more complete the combustion.</td>
</tr>
<tr>
<td>Viscosity</td>
<td>Low viscosity leads to good atomization.</td>
</tr>
<tr>
<td>Sulfur</td>
<td>Low sulfur content means low wear in the fuel system and lower total particulate content in the exhaust.</td>
</tr>
<tr>
<td>Density</td>
<td>The higher the density, the greater is the energy content of the fuel.</td>
</tr>
<tr>
<td>Waxing Tendency</td>
<td>Wax precipitation can render cold starting difficult and subsequently stop the engine.</td>
</tr>
</tbody>
</table>
The cetane number describes the ignition quality of the fuel. The shorter the ignition delay the better the ignition quality of the fuel, and thus, the higher the cetane number. Since DME has a higher cetane number than conventional diesel fuel, it will ignite readily and burn more completely.

The viscosity of DME is much lower than that of diesel fuels. This offers an advantage in that the fuel will be easier to deliver into the engine cylinder than diesel fuel during cold weather conditions. However, some studies have shown that the fuel leaks from the injectors [79, 80]. In addition, using neat DME within an engine creates some lubricating problems because of the low viscosity. What researchers are now understanding about the fuels used in automotive fuel injection systems, is that inherent lubricating traits of the fuel are also a very significant factor, especially when additives and alternative fuels are being considered [62, 63, 81, 82].

The boiling point of DME is another important advantage for its use as a fuel. Again, it proves to have better characteristics for cold starting conditions, which is a key factor in engine development. The vapor pressure of DME is a concern. Since the fuel is a gas at atmospheric pressure, and since we need to mix the fuel and inject the fuel as a liquid, the fuel and entire fuel system needs to be pressurized. This leads to other complications with fuel delivery, although the technology to do this is similar in nature to LPG (Liquid Propane Gas) because it is also moderately pressurized to keep the fuel in a liquid state [83].

Another important aspect of combustion emissions from a diesel engine fueled on diesel fuel versus DME, is the reduction and elimination of particulate emissions.
Particulate emissions are also commonly known as “soot” or black smoke. The oxygen content of a fuel blend with DME (at roughly 40 to 100 wt.%), allows for the emissions to be smokeless, as shown in the literature [23, 30, 38, 78, 84-86]. Nabi and coworkers evidence shows “smokeless” engine operation from a diesel fuel with an oxygen content at around 38 wt. % [23]. However, the work by Chen and coworkers confirms that even with 80 wt.% DME addition to diesel fuel, some smoke will be produced at high engine loads, even though it is a small amount [84].

The heating value of DME is a concern, because the heating value of diesel is roughly 1.7 times that of DME. This results in the need for more volume of fuel to produce the same output from combustion. By altering injection amounts to the cylinders, the amount can be compensated to counteract the decreased heating value and prevent “de-rating” of engine output.

Other issues will need to be addressed in future work regarding the understanding of DME fuel properties, including, for example, the lubricity of the fuel. Because of the need for the fuel to be tested while in the liquid phase under pressure, further analysis outside of combustion studies may be impractical or require development of highly specialized instrumentation.

2.6.1 Supercritical behavior of DME

A supercritical fluid is defined as the state of a substance at the point above the critical pressure and temperature of a substance [87]. This is a point at which there is a loss in the distinction between the liquid phase and gas phase. Another way to define the
fluid would be through the use of isotherms from a pressure-volume diagram, when temperature is the critical temperature, \( T \), the following relationship is given in equation \( 2.4 \) is given:

\[
\left( \frac{\partial^2 p}{\partial V^2} \right)_T = \left( \frac{\partial p}{\partial V} \right)_T = 0 \text{ at the critical point} \tag{2.4}
\]

This suggests isothermal compressibility, \( \kappa_T \), defined in equation \( 2.5 \) tends to infinity at the critical point. The symbol \( \kappa_T \) defines the rate of change in volume with pressure at constant temperature.

\[
K_T = -\left( \frac{1}{V} \right) \times \left( \frac{\partial V}{\partial p} \right)_T \tag{2.5}
\]

Therefore, in the critical region, the compressibility is high [87].

Supercritical fluids exhibit characteristics such as compressibility, homogeneity, and continuous changing between gas-like and liquid-like properties, as \( \kappa_T \) tends to infinity at the critical point. [87]. The compressibility of the substance is described by the compression factor, \( Z \), and is defined by equation \( 2.6 \) as:
\[ Z = \left( \frac{pV}{RT} \right) \] where \( R \) is the universal gas constant

and at the critical point

\[ Z_c = \left( \frac{p_c V_c}{RT_c} \right) \]  \hspace{1cm} (2.6)

The critical properties for DME are [88]:

Critical Temperature \( T_c = 400 \text{ K (126.85 } ^\circ \text{C)} \)

Critical Pressure \( P_c = 5370 \text{ kPa (53 atm)} \)

Critical Compression Factor \( Z_c = 0.273 \)

A graphical representation of a supercritical fluid is shown in Figure 2–9.
In this study, the DME may experience changes in fluid properties as it is drawn into the fuel injector and injected into the engine. The DME and diesel fuel are combined into one miscible fluid at 20° C and 100 psi. Therefore, because the fuel is a mixture, it is assumed that the properties of the two individual fluids combine to yield intermediate property values for the mixture. Since fuel injectors are designed to perform optimally with liquid properties similar to diesel fuel, maintaining the fluid close to the ASTM D 975 specification for No. 2 diesel standard is important. Since temperatures and pressures change throughout the injection process, knowing the temperature and pressure dependencies of the properties of the fluid can give some insight into how the properties of the fuel being injected are changing through the injection process.

Figure 2–9: The phase diagram of a single substance [87]
2.7 Production of DME

From the production standpoint, if DME does become a replacement fuel in the future, larger production facilities will be necessary. The processes available today to produce DME deliver very small scale quantities. Via the dehydration of methanol, DME is made through a fixed bed catalyst process. Thus, the cost to make DME is more than the cost to make methanol [41]. For DME to become advantageous as a fuel, it must be made cheap enough to compete with diesel fuel costs.

The proposed process for larger scale production would combine methanol and DME synthesis via a single process step for the direct conversion of synthesis gas to DME. This can happen by allowing the methanol synthesis, water gas shift reaction and DME reactions to occur simultaneously. The synthesis involves a series of reactions, which follow via:

\[
\text{CO}_2 + 3 \text{H}_2 \rightarrow \text{CH}_3\text{OH} + \text{H}_2\text{O} \quad \text{[Reaction 1]} \quad (2.7)
\]

\[
\text{H}_2\text{O} + \text{CO} \rightarrow \text{H}_2 + \text{CO}_2 \quad \text{[Reaction 2]} \quad (2.8)
\]

\[
2 \text{CH}_3\text{OH} \rightarrow \text{CH}_3\text{OCH}_3 + \text{H}_2\text{O} \quad \text{[Reaction 3]} \quad (2.9)
\]

The introduction of the DME reaction, equation 2.9 [Reaction 3], serves to help relieve the thermodynamic constraints on the methanol synthesis. Also, the water formed in equation 2.9 [Reaction 3] aids in providing for the equation 2.8 [Reaction 2], which in
turn provides for the equation \[ 2.7 \] (Reaction 1). Therefore, the process is synergistic with high conversion levels [41].

In an effort to determine the short term and long term options for producing DME in mass quantity, scientists at Haldor Topsøe have proposed several methods to utilize existing facilities, as well as new facilities. For small scale production, some recommended facilities include:

- Direct dehydration of methanol via an adiabatic reactor and catalyst. The reactor would be operated at temperatures between 290 – 400 °C, with the feed rate at 10 bar g. The conversion rate was determined to be 80% for the first pass. Then, DME would be separated via a distillation column.
- Revamp existing methanol plants to make DME as a secondary product by co-producing DME in the methanol synthesis loop.
- Revamp existing methanol plants to make DME as the primary product by using a new type of catalyst and a new distillation section.

For medium sized plants, there are several methods to use, and it is recommended to let capacity determine technology. For very large plants, the autothermal reforming of natural gas to synthesis gas is preferred. \([Figure 2–10]\) below shows how this process works [41]:
Feasibility studies of the financial cost of DME production in comparison to LNG production confirm the viability of DME as a potential fuel [89]. For a similarly sized plant of LNG compared to DME, operating and capital costs are similar, with transportation costs being an issue for DME for greater than 6000 miles. However, the end fuel costs are very similar to each other, which would then force the application issue to the quality of the fuel for the particular application use.

Another avenue of focus regarding DME is the use of LPDME™ (Liquid Phase DME), a new fuel process technology created by Air Products & Chemicals Inc. In this process, synthesis gas is converted to LPDME™ in a single slurry-phase reaction using a

Figure 2–10: Diagram of DME plant based on autothermal reforming [41]
physical mixture of a commercial methanol synthesis catalyst and an APCI proprietary catalyst, slurried in mineral oil [40]. LPDME™, through the use of the Liquid Phase™ technology creates a fuel in a liquid phase process rather than a gaseous process. The advantage of this involves controlling the process through heat removal in the liquid, and allows for higher throughput from the reactor. The fuel from the process, which includes DME and, from an additional separate oxidative coupling process, longer chain linear ethers. These fuels can be used in neat form or as an additive, and are being considered as a cetane improver, and now goes by the name CETANER™. A schematic of this process is shown below in Figure 2–11.

Figure 2–11: Liquid Phase™ Reactor and Reaction Schematic [40]
An important aspect of this process is that it can be incorporated into an Integrated Gasification Combined Cycle (IGCC) facility to fully utilize the capacity of the gasifiers[90]. The Liquid Phase technology has been applied for other fuel processing, including methanol, called the LPMEOH process[90]. A schematic showing a Once-through LPMEOH process design option is shown in Figure 3–12.
Improving DME processing involves work in several areas. First, understanding how DME is formed leads to improved catalysts, and thus, higher conversion rates. Second, the capital costs for investing in a new facility are significant, and mean that a market needs to be created to demand such an establishment. Finally, creating a market
and addressing distribution issues, even though not addressed in most research, is inherent with any new fuel.

2.8 Research with DME in engine applications

Dimethyl ether is a common chemical used as a propellant for spray cans [26]. The properties of DME are given in Table 2–1 and are compared to the diesel fuel used for the baseline testing for this experiment. DME is a liquid when contained under moderate pressure, with a vapor pressure of 5.1 bar at 20°C, and is relatively easy to handle. Over the past ten years, researchers have begun to consider the use of DME as a fuel. Because the cetane number and ignition temperature are close to that of diesel fuel, DME was thought to be an excellent substitute for use in compression ignition engines. However, there were some drawbacks to using the fuel, including the reduced viscosity and lubricity of the fuel in neat form, as well as fuel compressibility effects [41].

To potentially overcome the fuel property effects of DME, as well as, reduce emissions, the experiments for this study focus on mixing dimethyl ether with diesel fuel. The initial goal is to determine the effect of the oxygen concentration on the emissions, with minimal engine modifications. In this part of the work, no changes have been made to the fuel injection timing, fuel injectors, or engine programming. Changes to the fuel system have been made to allow the fuel to be delivered to the common rail as a liquid by maintaining the DME-diesel blend at over 100 psi.
Over the last ten years, many researches have begun to evaluate the performance and emissions effects of neat dimethyl ether. Sorenson and Mikkelsen [30] found that for a fixed speed and across various loads, the particulate and NO\textsubscript{x} emissions from a .273 Liter direct injection single cylinder engine fueled with neat dimethyl ether could be significantly reduced as compared to emissions with diesel fuel. In the same study, the HC and CO emissions showed little or no change. Later, Sorenson and Mikkelsen [91] further studied the HC emissions from this same engine, and found that there was an increase in the HC emissions when using neat DME, with more methane found than in a typical diesel engine, and less light hydrocarbons. With another engine, Christensen and Sorenson [92] looked at various effects on the suite of emissions when using neat DME. Of particular interest, the NO\textsubscript{x} emissions were significantly reduced when the injection timing was retarded towards Top Dead Center (TDC). However, there was an increase in the CO emissions, and little effect on the HC emissions. Other tests determined that lower injector opening pressure reduces NO\textsubscript{x}, and nozzle types did not seem to influence NO\textsubscript{x} emissions. Experiments by Kajitani and coworkers [85] also showed the effects of injection timing on reducing NO\textsubscript{x}, which had little effect on HC emissions, from a single cylinder Yanmar engine fueled with neat DME.

However, in the work completed by Hupperich and coworkers [93] with a 1.75 liter single cylinder engine for the ECE R49 13-mode test, the cumulative emissions show some differing results. With the use of neat DME, HC emissions are reduced. The trends with the other emissions are similar to what had been determined with previous studies. One difference to note is the change in injection nozzle size, which may have
affected the emission results in allowing for more complete combustion of all fuels tested in an effort to maintain consistent test conditions.

Recently, experiments completed by Ikeda and coworkers [86] with a single cylinder engine using a binary fuel injection method, showed similar NO$_x$ emissions between diesel fuel and 40 volume % DME mixed with diesel fuel, as injection timing was retarded. Also, HC emissions increased and smoke emissions were reduced as injection timing was retarded. In addition, comparisons were made as a function of BMEP. NO$_x$ was reduced, HC remained constant and smoke increased with increasing Brake Mean Effective Pressure (BMEP). The experiments also included % DME fractions mixed with diesel fuel up to 60 volume % addition, with comparisons made to the baseline diesel fuel. The smoke level, indicating presence of soot in the exhaust stream, showed a slight increase between 0 and 20 % DME addition, and then returned to zero for DME addition over 20%. NO emissions decreased slightly, then increased slightly to the original point for diesel fuel. HC emissions increased slightly up to 45% DME addition, and increased sharply above this point [86].

Many researchers have been evaluating the performance of other oxygenates including blends of glycol ethers with diesel fuel, and have observed decreases in particulate matter emissions with increasing oxygenate concentration. Most recently, Hallgren and Heywood [6] prepared a review of the collection of work which showed that as the oxygen content of the fuel increases, the particulate matter is reduced, suggesting that this occurs regardless of chemical structure or molecular weight. However, their actual testing showed that the oxygenate structure did impact particulate
emissions. Studies completed by Hess and coworkers [15] as well as by Litzinger and coworkers [16, 17] have shown that higher molecular weight glycol ethers are also effective in reducing particulate matter emissions, although to a lesser extent than monoglyme or diglyme.

Although it has been shown that glycol ethers effectively reduce particulate emissions, the fundamental mechanisms of the reduction have not been clearly identified. There has been some work in simulating the ignition and rate mechanism behavior of dimethyl ether in comparison to dimethoxymethane [25]. Also, oxidation mechanisms have been proposed for gaseous forms of DME [38, 94, 95]. More recently, the modeling of DME oxidation has proven consistent with experimental results from jet stirred reactor theory and shock-tube conditions, providing confidence in reaction mechanisms [96].

2.9 Fundamental Property Research on DME

It almost seems like a mythical story. It is true that everyone studying the use of DME refers to this as the beginning of the journey: “New fuel: The lawnmower’s tale” [97]. One day, an excited scientist for Haldor Topsøe took some DME home to try in his lawnmower, and had great success. Then, he tried it in a diesel fork lift at the plant, and was astounded when the power was turned off and the fork lift kept running. Again, this supports the finding that DME is a very compressible and ignitable fuel.
Another important aspect of DME that is demonstrated through this tale is the
chemical kinetics of the combustion process. It has been demonstrated that the radical
reactions during the propagation phase of the combustion process include OH, H and CH₃
[43]. With the presence of the OH radical, the ignition quality is enhanced by making the
fuel mixture more reactive, thereby shortening the ignition delay and increasing oxidation
rates. When the OH radical was present and DME was provided, the reaction continued.
The proposed reaction channels for the presence of the OH radical are equations 2.10 and
2.11[42]:

\[
\text{CH}_3\text{OCH}_2 + \text{O}_2 + \text{M} \rightarrow \text{CH}_3\text{OCH}_2\text{O}_2 + \text{M} \quad (2.10)
\]

\[
\text{CH}_3\text{OCH}_2 + \text{O}_2 \rightarrow 2\text{CH}_2\text{O}^\bullet + \text{OH}^\bullet \quad (2.11)
\]

From other work, the proposed decomposition reactions include the following
chain reaction in equation 2.12[43]:

\[
\text{CH}_3\text{OCH}_3 \rightarrow \text{CH}_3^\bullet + \text{CH}_3\text{O}^\bullet \quad (2.12)
\]

Then, a CH₃ radical abstracts a hydrogen atom from a second molecule of dimethyl ether
by the equation 2.13
The resulting \( \text{CH}_2\text{OCH}_3 \) radical then decomposes to formaldehyde and methyl radical, and then finally to formaldehyde and methane, according to the following reactions, equations 2.14 and 2.15[43]:

\[
\text{CH}_2\text{OCH}_3 + \text{M} \rightarrow \text{CH}_3 + \text{CH}_2\text{O} + \text{M} \\
(2.14)
\]

\[
\text{CH}_2\text{OCH}_3 + \text{M} \rightarrow \text{CH}_4 + \text{CH}_2\text{O} + \text{M} \\
(2.15)
\]

Within this study, the author proposed and concluded the following competitive reactions, equation 2.16 through an ab initio study:

\[
\text{CH}_3\text{OCH}_3 \rightarrow \text{CH}_3\text{O} + \text{CH}_3 \\
\rightarrow \text{CH}_3\text{OCH}_3 + \text{H} \\
\rightarrow \text{CH}_4 + \text{CH}_2\text{O} \\
\rightarrow \text{CH}_3\text{OH} + \text{CH}_2 \\
\rightarrow \text{CH}_3\text{OCH} + \text{H}_2 \\
(2.16)
\]

Even though there is some experimental and theoretical work for the combustion mechanisms of DME, there is still much work to be done to fully understand the process.
In addition, rate coefficients for the hydroxyl radical reaction have been determined over temperature ranges. This is an important key in preparing a theoretical model of the heat release rate during the combustion of DME. The Arrhenius expression recommended for combustion modelers is in equation 2.17 below:

\[ k(295-650K) = 1.05 \times 10^{-7} T^{2.0} \exp[-328/T] \]  
\[ \text{ (in units of cm}^3\text{ molecule}^{-1}\text{ s}^{-1}) \]  
(2.17)

This study also concluded that the dominant fate of the \( \text{CH}_3\text{OCH}_2 \) radical was the beta bond cleavage between the C-O bond, which finally decomposed to formaldehyde and methyl radicals [98].

Not only has DME been demonstrated theoretically and experimentally to exhibit rapid reaction chemistry, but also DME has been shown to reduce NO and CO emissions [99]. In comparing DME with propane and butane, DME demonstrated reduced CO emissions, and a less striking, NO reduction. It was speculated that this is due to the longer residence time of DME in the flame zone, confirming the speculation about the rapid reaction chemistry.

### 2.10 Technical Issues with Utilization of DME as a fuel

Much of the early work with DME utilization has been done at the AVL labs in Graz, Austria. The testing concluded that DME does indeed reduce the particulate
emission to zero, and also concluded that the typical diesel fuel injection system does not tolerate DME [79]. Below is a combined list of concerns not only from this testing, but also from other groups who are now working through the fuel property, design and technology issues for utilization of DME:

- DME was leaking past clearances on the injectors and seals. This caused the need for the camshaft housing and crankcase of the engine to be vented [79].
- At high vapor pressure, the DME was cavitating, which caused difficulties in maintaining stable fuel injection.
- While DME is more compressible than diesel fuel, it was found that the compressibility changed with temperature and pressure. Therefore, this made it difficult to inject the maximum fuel quantity at high temperatures and during full load operation using traditional diesel equipment [79].
- DME chemically attacked some seals [79].
- Not much effort has been put towards understanding the environmental impacts of the compound itself or the emissions from the fuel combustion, as compared to other fuels [79].
- A larger fuel tank will be required, as compared to diesel fuel, because of the lower density and heating value of DME [80].
- Since the vapor pressure of DME is low, the fuel vaporizes immediately upon injection into the cylinder. This may or may not be an issue, but further study may be
suggested to confirm how the combustion reaction takes place after the vaporization occurs [80].

• Injection via some fuel pumps causes uncontrollable pressure waves in the entire system [100].

• Predictability of spray behavior and characteristics is important in repeatability of combustion [101].

• Turbulence within the cylinder is important for mixing of the fuel, which in turn reduces emissions [102].

2.11 Health and Safety Concerns

As stated earlier, DME is used as an aerosol propellant. DuPont Fluorochemicals markets DME as Dymel A. With regard to flammability, DME has a higher explosion limit than propane, but it should still be considered extremely flammable and proper caution should be used. DME is a very stable compound, as it does not decompose to peroxides under normal conditions encountered in the aerosol industry. Dimethyl ether is highly soluble in polar and non-polar solvents, which means that it is miscible in water. According to the studies reviewed by DuPont Fluorochemicals, DME has low acute and chronic toxicity. In circumstances of high inhalation levels, the main affect to a person would be of that of a weak anesthetic. From a longer term inhalation study on animals, the exposure level of 20% showed no compound related effects. In summary, DME showed no signs of carcinogenicity, and no evidence of mutagenicity or teratogenicity.
Therefore, it had been given approval for use in consumer products and approved by the DuPont company [103].
Chapter 3

Methodology: Experimental Facility and Combustion Measurements

3.1 Introduction

The key feature of the compression ignition engine is to control the ignition of the fuel to occur at the appropriate time in the 4-stroke process. There is an appropriate point for fuel injection into the engine at which engine power and torque output are optimized in conjunction with minimizing the exhaust emissions [1]. Additionally, this optimal point for fuel injection may change as the engine’s speed and load changes. The optimal point for fuel injection is a function of the ignition delay of the fuel, or the time it takes from the moment of injection until a rapid rise in pressure occurs [1]. As changes in fuel composition are made, changes in the ignition delay may occur due to changes in the fuel injection/mixture preparation process, and changes in the autoignition chemistry of the fuel. To address this conflict between fuel economy and low emissions from a modified fuel, it is important to understand the effect of fuel on the timing of the combustion process, as observed by the emissions, cylinder pressure trace, and heat release rate.

There are many ways to assess the effect of the fuel on the combustion process. In this work, a pressurized fuel delivery system was implemented on a turbodiesel engine to permit operation on DME-diesel blends, and thereby to permit examination of the
impact of DME on the diesel combustion process. Later, information gained from this data will be used to determine how to optimize the engine’s operation.

The following discussion provides the specific details of the experimental facility, experimental test procedure, and measurement acquisition. A brief overview of important characteristics of the engine’s fuel system will assist in understanding and explaining the information presented in the Results and Discussion section.

3.2 Experimental Design

An experimental facility was designed to permit operation of a Navistar 7.3L “T444E” V-8, direct injection turbodiesel engine on DME-diesel fuel blends, with minimal changes to the fuel system. The following sections describe the engine, modifications to the fueling system and equipment used to quantify the exhaust emissions and effects on ignition delay.

3.2.1 Test Engine

To study the effects of fuel additives on combustion in light-duty to medium-duty diesel engines, a Navistar T444E 7.3L Turbodiesel engine was coupled to a 450 horsepower Eaton (Model AD-1802) eddy-current dynamometer. The specifications of the engine are given in Table 3–1. A Pentium PC with Keithley Metrabyte DAS-1800 data acquisition card was connected to the engine to log real-time engine outputs. These outputs included engine crank angle, cylinder pressure, mass air flow, oxygen concentration and engine
load (torque). Additionally, a Modicon Programmable Logic Controller (PLC) was interfaced with the same computer connected to the Kiethly DAS acquisition system. A Visual Basic program was written and used to combine both acquisition system’s output into one data collection/storage location, with output streamed into an Excel spreadsheet. The outputs from the PLC included engine speed, torque, and temperatures from the engine. From this data, power was calculated. The Modicon PLC (Programmable Logic Controller) was used to record temperatures from the engine, as well as, from the entire experimental system. Intake airflow rates were measured directly via a Delphi electronic mass air flow sensor (Model No. 25178988; 90.0mm dia. Aluminum inlet and outlet, 74mm dia. throat), which was calibrated using a laminar flow element. Fuel consumption was monitored using a precision Sartorius scale (Model EA60EDE-1), with an accuracy of ± 2 grams. Figure 3–1 shows the test cell set up, and additional equipment used for emissions monitoring.
Table 3–1: Characteristics of the 1998 Navistar T444E 7.3L Turbodiesel engine [104]

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Displacement</td>
<td>444 cu. in. (7.3 Liter)</td>
</tr>
<tr>
<td>Bore</td>
<td>4.11 inch (104.39mm)</td>
</tr>
<tr>
<td>Stroke</td>
<td>4.18 inch (106.20mm)</td>
</tr>
<tr>
<td>Rated Power</td>
<td>190 HP @ 2300 RPM</td>
</tr>
<tr>
<td>Peak Torque</td>
<td>485 lbf-ft @ 1500 RPM</td>
</tr>
<tr>
<td>Configuration</td>
<td>Turbo charged, Charge air cooled (Air to Air), Direct Injection</td>
</tr>
<tr>
<td>Injection Scheme</td>
<td>HEUI- Hydraulically actuated, electronically controlled unit injectors</td>
</tr>
<tr>
<td>Low Idle Speed</td>
<td>700 RPM</td>
</tr>
<tr>
<td>Features</td>
<td>Split-shot injection (low speed only)</td>
</tr>
<tr>
<td>Compression Ratio</td>
<td>17.5:1</td>
</tr>
</tbody>
</table>
3.2.1.1 Fuel Injection

Most of the modern concepts of fuel injection come from the early developments of Robert Bosch [77]. The Navistar T444E fuel system is based on the unit fuel injector concept. In simple terms, this means that the fuel is pumped from the tank to the fuel gallery that is common (common rail) to a series of injectors. The injector is built so that it times, meters, pressurizes, and atomizes the fuel as it enters the combustion chamber. Additionally, typical systems will use this fuel to cool and lubricate the injectors, while
passing a large portion of the fuel back into the tank. However, the Navistar T444E has a dead head style fuel rail, meaning that no fuel is returned to the tank from the fuel injectors. The fuel remains in the rail until the fuel is used [105].

The T444E engine utilizes a unique fuel injector system developed by Caterpillar, Inc. The injector is commonly referred to as a “HEUI”, which means Hydraulically actuated Electronically controlled Unit Injector. A picture of the injector is found below in Figure 3–2. The injector is actuated through the use of high pressure engine oil. The pressure intensification is achieved by the design of the injector, as the area of the piston is seven times the area of the plunger. The fuel is injected at pressures from 19 to 120 MPa through a conventional multi-orifice nozzle [29].

Figure 3–2: Schematic of HEUI Fuel Injector [29]
The injection control pressure is governed by the Electronic Control Module (ECM) through the Injection Pressure Regulator (IPR). As stated above, the pressure is achieved through the use of the engine oil, so the sensors are located in the engine head oil reservoirs. There is a feedback closed loop control on the pressure, accomplished through the use of an Injection Control Pressure Sensor (ICP). The engine control gets information from the ICP on the actual pressure, at all times. The information is then used by the ECM to determine how to adjust the IPR. Figure 3–3 helps to convey how the ECM operates the injection control system [104]. Figure 3–4 shows the closed loop operation of the injection pressure control.
Figure 3–3: Injection Control System [104]

ICP-Injection Control Pressure Sensor

IPR- Injection Pressure Regulator

ECM- Engine Control Module
The IPR operates by pulse width modulation (duty cycle %) at a frequency of 400 Hz. The pulse width is modulated from 8-60% to control the pressure from 3.4 to 20 kPa (500 to 3000 psi). As the engine demand for pressure increases, the ECM increases the pulse width on the IPR solenoid, which in turn increases the oil pressure. Figure 3–5 shows how higher injection pressure is commanded [104]. Notice the pulse width in the upper portion of the figure.
Figure 3–6 shows how lower injection pressure is commanded. Notice the pulse width duty in the upper portions of the figures. It is the “on time” of the pulse which controls the operation of the armature on the solenoid windings, which in turn opens and closes the spool valve to keep the required pressure. This is the pressure of the injector, in the upper potion above the intensifier plunger. Therefore, this pressure is critical to the injector’s operation.

Figure 3–5: IPR Higher Injection Pressure [104]
The operation of the HEUI fuel injector is divided into three stages: fill, injection, end of injection. During the fill stage, the solenoid is de-energized and the poppet valve is in the closed position, preventing oil from entering the intensifier piston area. Without oil in the intensifier piston area, the piston locates itself in an upper open position, which allows for fuel to flow into the nozzle of the injector. When, the ECM commands injection, the solenoid is energized, which starts the injection stage. The poppet valve is lifted, and high pressure oil enters the intensifier piston area. The pressure is transmitted from the

*Figure 3–6: IPR Lower Injection Pressure [104]*
oil to the fuel and intensified by a factor of 7, creating pressures up to 21,000 psi. This is accomplished by the area of the lower portion being 7 times that of the upper portion. Once the pressure is high enough to lift the needle valve, high pressure fuel is sprayed through the nozzle and atomized into the cylinder. After the injection is complete, the solenoid is de-energized and the high pressure oil leaves the intensifier piston area. The fuel injector returns to the fill stage. Figure 3–7 shows a pictorial view of the injector and how the stages work [104].
3.2.1.2 Turbocharging

The end result of what is known as turbocharging is to increase the mass of air brought into the cylinders of the engine by raising the density of the intake air, and thus allow more fuel to be burnt. The power output of the engine is increased for a given swept
volume of the cylinders. On an engine, a compressor is used to achieve this increase in air density. The system may have either a turbocharger or a supercharger. In a supercharged system, the compressor is driven from the crankshaft of the engine, and thus called “mechanically driven supercharging”, or just supercharging [1]. In a turbocharged system, the compressor is driven by a turbine, which is driven by the exhaust gas.

Since the process of compression raises cylinder or charge temperature and pressure, a charge air cooler is used to cool the air between the turbocharger and the intake manifold. This assures that a maximum rise in density is achieved with the pressure increase. Figure 3–8 shows a comparison between an ideal combustion cycle and a supercharged cycle on a P-V diagram (Pressure-Volume). Supercharing gives the same result as turbocharging, utilizing a different method of compressing the gas. As can be seen, turbocharging causes the inlet and exhaust temperatures to be above ambient, with a total pressure increase throughout the cycle [1]. The figure shows this comparison with what is called a dual combustion cycle. This cycle represents a combination of a constant pressure and constant volume cycle, which is a closer approximation to an actual diesel cycle.
3.2.1.3 Intercooling

The term intercooling is often used to describe the temperature reduction of the turbocharged charge air. As described above in the turbocharger section, the cooling is necessitated by the increase in temperature associated with the increase in pressure and density of air achieved by the compression effect in the turbocharger. Again, the net goal is to increase the density of intake air, thus providing for more fuel to be combusted [1].
3.2.2 Pressurized Fuel Delivery System for Diesel-DME Blends

Dimethyl ether (DME) is a liquefied gas. At room temperature and atmospheric pressure, it is a gas, but changes to a liquid at a moderate pressure. Dimethyl ether is currently manufactured by DuPont Fluorochemicals under the trade name Dymel A. For the purposes of effective operation of the experimental design, information regarding the vapor pressure, density changes with temperature, viscosity, and miscibility was needed. A portion of this information is reported in Figure 3–9 from the DuPont Technical Information (ATB-25) bulletin [103]. However, no experimental data exists for DME-diesel fuel blend properties, which had to be determined experimentally.

Dimethyl ether was found to be miscible with #2 diesel fuel. Miscibility tests were carried out in a pressurized vessel with a glass observation window [106]. The two fuels were introduced taking care not to physically mix them. Diesel was introduced first into the bottom of the vessel. DME, which has a specific gravity less than diesel fuel, was then introduced on top of the diesel fuel. Thus, initially there were two distinct layers. The two layers were then observed to mix together without physical agitation after a period of 5 to 6 hours to form a homogeneous mixture, at up to 60% DME by mass in the mixture. Furthermore, no separation was observed after standing undisturbed for about 3 days [107].
Also, *Figure 3–10* shows the changes in density of DME as a function of temperature. This information was used to determine the energy density of the fuel.
A schematic of the modified fuel system is shown in Figure 3–11. The fuel system on the T444E engine had to be modified to account for the need to deliver fuel at elevated pressure, because of the vapor pressure of DME at higher temperatures. For the unmodified Navistar T444E engine, the fuel rail in the cylinder head of the engine receives fuel at a pressure of about 70 psi. Fuel from this rail is then fed to the injectors. A study was performed using #2 diesel fuel to measure the temperature rise of the fuel in the fuel rail. This measurement, coupled with the fuel consumption, gave an approximate heat transfer rate between the cylinder head and the fuel in the gallery. A maximum target

Figure 3–10: Density changes of Dymel A as a function of temperature [103]
temperature of 50°C was chosen for the diesel-DME blend based on the vapor pressure curve of DME and the pressure rating of the fuel rail. This required a change in fuel recirculation flow rate was then calculated based on the above observations. This recirculated fuel was then cooled using a water cooled heat exchanger. The fuel delivery pump was sized based on the required flow that was calculated. Selecting a pump for DME was challenging due to the properties of DME. Gasket material for the pump had to be modified, as common materials such as Viton and buna-N have been found to be unsatisfactory [108]. A fuel filter with a high filter surface area and high pressure capacity was needed. A modified propane filter was selected for the application. The fuel tank consisted of a modified 100 lb capacity LPG cylinder which was pressure tested at 120 psi prior to use.

The fuel delivery system works as follows:

1. The fuel tank is pressurized to between 90 and 100 psi using helium to pressurize the space above the fuel. Therefore, this pressure is the force to drive the fuel from the tank into the engine fuel system which eliminates the need for a fuel transfer pump. The overpressure is necessary to keep the DME in the liquid state. Any inert gas can be used. In this case, helium was chosen because it has a lower solubility in DME than nitrogen.
2. The pressure of the fuel is then boosted by a gear pump to between 120 to 150 psi. A higher pressure allows for the fuel to absorb more heat and still remain in the liquid state. The unmodified rail pressure is 70 psi, provided by the low pressure fuel pump. The fuel injectors fill through a pressure difference created by the rail pressure, regardless of what the pump outlet pressure to the fuel rail is. This higher pressure fuel may be increasing the force present on the fuel o-rings located around the fuel injector. This is important to note because this higher pressure around the sleeve of the injector could cause fuel to leak into the oil portion of the fuel injector line, if after repeated use the o-ring begins to fail.

3. The fuel return loop pressure is held at about 120 to 150 psi, depending on the pressure setting of the back pressure regulator. The regulator is a simple spring loaded valve that regulates the flow to keep the back pressure at the desired setting. A pressure gage was placed upstream and close to the back pressure regulator to verify system pressure.

4. The return fuel then passes through a heat exchanger, where cooling of the fuel after exiting the rail is necessary to keep the temperature of the fuel in the rail below 45 °C.
5. After cooling, the fuel is then fed to a tee in the fuel line, prior to the inlet of the pump. All fuel in this portion of the line is at 100 psi, which is the pressure of the fuel tank.
71

Figure 3–11: Diagram on the pressurized fuel system for the Navistar T444E Turbodiesel engine
3.2.2.1 Pressure, Temperature, and Flow Requirements

At 20°C, the vapor pressure of DME is about 0.52 MPa (75 psia), and is a strong function of temperature. Keeping DME in a liquefied state calls for pressurizing the entire fuel system from the fuel tank up to the fuel injectors. The vapor pressure also changes strongly with temperature. The required pressure of the fuel system is hence dictated by the fuel temperature. The pressure, however, is limited by the pressure rating of the fuel rail. The engine used in the study has a common rail injection system. Both cylinder heads have a fuel rail running along their length, which is the source of fuel for the pressure intensifier in the fuel injectors. In the original fuel system of the engine, the pressure in the rail is maintained at 0.48 MPa (70 psig). This facilitates proper filling of the pressure intensifiers. The fuel rails in the cylinder head form a dead head system. This means that there is no fuel return once the fuel enters the fuel rail. Because of this configuration, the fuel temperature in the rail approaches the engine coolant temperature in the head. This layout of the fuel system was modified to accommodate a fuel return from the cylinder heads.

In designing the experimental facility, a study was performed in which the temperature of the fuel in the fuel rail was recorded in conjunction with the fuel consumption of the engine, for the “AVL 8-mode” test protocol [38]. Approximating values for heat capacities for diesel and DME, a minimum flow rate value was calculated so as to keep the temperature of the fuel in the rail below 50°C. The vapor pressure of DME at this temperature is about 1.0 MPa (150 psi). This pressure, more or less, dictated the allowable temperature rise. The fuel delivery pump was sized based on the above
calculations. Because the cooling provided by the fuel flow rate was not sufficient to keep the fuel in the liquid state, cooling of the returned fuel was necessary to maintain the required fuel temperature. In these tests, a 500 W capacity chiller was used to chill a bath through which the fuel was passed within stainless steel coils. This fuel cooling was insufficient to maintain the fuel temperature below 50°C under some operating conditions, particularly during Mode 8 which is close to rated speed and maximum load of the engine.

The final system has an operating pressure of 150 psi at the back pressure regulator, and the flow rate of the fuel is fixed as it moves through the system. The fuel cooling was insufficient for high load conditions (at the peak load of the engine) at medium and high engine speeds. This indicated further optimization of the system was required.

3.2.2.2 Design Considerations for the Pressurized Fuel System

Dimethyl ether is known to be incompatible with common gasket materials such as Viton and buna-N, used in diesel service, as shown in the Chemical Resistance Guide available at the Dupont-Dow Elastomer website. Data provided by DuPont-Dow, Inc. indicated Kalrez™, a perflouropolymer, (designed and manufactured by DuPont–Dow Inc.) to be the best material for DME. For economic considerations, however, this material was used sparingly for the engine fuel system modifications. Other materials such as butyl rubber, Teflon and neoprene have also been found to be compatible, though
not to the same degree as Kalrez™. Stainless steel was used for the fuel lines as a safeguard against corrosion. All the other components such as valves and regulators were also made of stainless steel.

Selecting a pump for circulating DME was challenging due to the low lubricity and low viscosity of DME. Due to the vapor pressure of DME, the pump housing was required to handle pressures up to 1.7 MPag (250 psig). Positive displacement pumps such as vane pumps, diaphragm pumps and gear pumps were considered. Gear pumps were found to be economical, as well as convenient to operate. With these considerations, a gear pump made by Tuthill Pump Co., California (Model #TXS2.6PPPT3WN00000) was selected. This pump has a magnetically coupled AC motor. This configuration does not have a driveshaft going through the pump housing, which circumvents the need for seals, a potential source of leakage. The gear material is Ryton (Polyphenylene sulphide), which was found to be compatible with diesel and DME as per the data provided by DuPont Fluorochemicals, Inc. The pump body seals are made of Teflon.

The original fuel filters on the engine could not be used because of the high pressure of the modified fuel system. The minimum pressure in the fuel lines was 0.62 MPag (90 psig). This required the use of special filters, which would withstand higher pressure. A diesel water separator was used as a primary filter. This is rated at 0.69 MPag (100 psig). The final filter was an LPG filter rated at 3.4 MPag (500 psig). The mesh size of the filter was 2 micron, very near to the engine specification.
The fuel tank was made out of a modified 45 kg (100 lb) capacity LPG cylinder which was pressure tested prior to use. This tank was fitted with a 1/2” NPT fitting at the bottom for liquid exchange.

### 3.2.2.3 Redesign of the Modified Fuel System

From previous studies on this same engine for the DME-diesel blend providing 2 wt.% oxygen in diesel, the cooling capacity of the heat exchanger and fuel circuit was determined to be insufficient, based on the fuel temperatures recorded, as well as observed engine instabilities [109]. In response to this shortcoming, the system, shown in Figure 3–11, was modified by the addition of a second fuel coil in the cooling bath, and a larger chiller unit for cooling the bath. Additionally, the system was pressurized to 150 psi, which then increases the allowable fuel temperature before the DME vaporizes.

Testing with this modified system has continued so as to determine the flow rate and pressure drop through the entire fuel loop. Because the flow conditions in the fuel system are critical, sizing a pump appropriate for maintaining the fuel in a liquid state is essential.

### 3.2.3 Test Procedure

In this work, an AVL 8-mode test procedure has been utilized as a model for diesel emissions tests. The AVL 8-mode test was designed to correlate to the U.S.
Federal Heavy-Duty Transient Test procedure through a weighted 8-mode steady state test procedure. The 8 modes are a combination of speeds and loads, which when combined with the weighting factors, reports the same emissions output as would be recorded for a transient cycle [38]. For this engine, the test procedure included the speed and load settings shown in Table 3–2

Table 3–2: AVL 8-Mode Test for the Navistar T444E Turbodiesel engine

<table>
<thead>
<tr>
<th>Mode</th>
<th>Speed (RPM)</th>
<th>Load (ft-lb)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>700</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>876</td>
<td>84</td>
</tr>
<tr>
<td>3</td>
<td>1036</td>
<td>224</td>
</tr>
<tr>
<td>4</td>
<td>1212</td>
<td>357</td>
</tr>
<tr>
<td>5</td>
<td>2300</td>
<td>77</td>
</tr>
<tr>
<td>6</td>
<td>2220</td>
<td>178</td>
</tr>
<tr>
<td>7</td>
<td>2220</td>
<td>307</td>
</tr>
<tr>
<td>8</td>
<td>2124</td>
<td>409</td>
</tr>
</tbody>
</table>

Another perspective of the testing can be seen in Figure 3–12 showing Modes 1-4 within the low speed region and defined by increasing load, and Modes 5-8 within the high speed region of the engine and defined by increasing load.
An extended warm-up period was used to prepare the engine for testing. The sampling and measurements during each mode commenced when the exhaust temperature reached steady state. During this time, speed and torque were maintained within 2% of the target test conditions. Once steady-state operation was achieved,
portion of the exhaust gas was passed through a Sierra Instruments BG-1 micro-dilution test stand with a constant dilution air / sample flow ratio of 8:1 and a total flow of 150 liters/min. These settings were chosen in order to maintain the temperature of the particulate sampling below the International Standard (ISO/DIS 8178-1.2) of 52°C [46]. Particulate were collected on Pallflex 90mm filters (Type EMFAB TX40HI20-WW), conditioned in an environmental chamber at 25°C and 45% relative humidity before and after sampling. For reproducibility, five particulate samples were taken for each fuel at each test mode.

Exhaust gas analyses were completed using a Nicolet Magna 550 Fourier Transform Infrared (FTIR) Spectrometer. For each mode, five gas samples were analyzed for CO₂, CO, NO and NO₂. The FTIR has a 2 meter cell, and operating conditions include keeping the cell temperature at 150°C and the cell pressure at 680 mm Hg absolute. The FTIR was calibrated in specific frequency ranges so as to minimize other molecular interferences, especially from water. The species are calibrated for a particular ppm range in the following wave numbers: CO₂ (732.8-766.0), NO₂ (1580.0-1632.0), NO (1850.6-1953.5), and CO (2002.0-2058.8). Also, a Rosemont Analytical on-line O₂ analyzer was used to monitor the oxygen concentration in the exhaust gas. The oxygen readings were used in conjunction with the mass flow sensor to determine and verify the air / fuel ratio. Additionally, total hydrocarbon emissions were monitored using a California Analytical Instruments Model 300 HFID Heated Total Hydrocarbon Gas Analyzer. For the total hydrocarbon measurements, undiluted exhaust gas was collected via a heated sample line, which was maintained at 190°C. Calibration of the
HFID and the Rosemont Analytical on-line O₂ analyzer was completed prior to each day of testing. All other equipment calibrations were checked on a periodic basis.

3.2.5 Pressure Trace Analysis

In order to observe the impact of the oxygenated blends on combustion and heat release, the combustion chamber of Cylinder # 1 of the engine was fitted with a Kistler 6125A pressure probe. The pressure sensor was used with a Kistler 2612 optical crank angle encoder to provide time resolved in-cylinder pressure traces of the combustion event. Pressure, crank angle, and TDC trigger signals were acquired with a Keithley DAS-1800 data acquisition card operating in a “burst” mode. The pressure traces were analyzed with PTrAn V.02, a software product designed by Optimum Power. The techniques used for the analysis are based on the Rassweiler and Withrow methods, established in 1938 [110]. The methods assess burn rate from measured pressure data by estimating the apparent pressure rise as a result of combustion.

3.2.6 Navistar Engine Control Data

In the process of debugging the engine, it was necessary to purchase the Service Technician Tool which would allow the failure modes of the engine to be observed. The tool allows for communication across the ATA (American Trucking Association) communication lines through one of the established communication languages of the
electronic controls. The tool does allow additional engine parameters to be monitored and for small portions of that information to be logged into the tool itself. The tool has served a great purpose in allowing for the engine to be easily debugged during the start up phase and while operating. It has allowed for understanding of which common failure modes the engine experiences. The tool allows for understanding of the engine operation, so that observations can be made of what may be occurring when the engine will not operate.

3.2.6.1 Automotive Electronic Communication Protocols

In the past, there have been many different types of communication protocols used by automotive and truck manufacturers. Over the past 10 years, a conscientious effort has been made to standardize these methods so as to reduce the proliferation of Service Technician tools, and the amount of software that needs to be programmed into the ECM for the manufacturing facility’s test equipment. This in turn helped to reduce the overall cost and complexity of the product by using the least memory necessary. The protocol which is gaining wide acceptance in the industry is called CAN (Controller Area Network). The important aspect of the protocol is that with the correct IC (Integrated Circuit) within the ECM, software can be transmitted serially to the product through the ECM connector. In comparison, in the past, if an ECM needed to be reprogrammed, this was accomplished by sending it back to the factory and programming occurred by probing on the microprosessor pins.
3.2.6.2 Engine Control Module Computer Interfacing

After gaining a brief understanding of the engine control module (ECM) electronics, the complexity of being able to make changes to software calibrations became obvious. Further, after numerous communications with Navistar, the possibility to modify the electronic controls calibration for the DME/diesel fuel blend optimization would cause Navistar a great deal of product liability. Additionally, this would create an open electronic architecture structure, which requires years of research to comprehend. The potential calibration method would potentially cause significant struggle with procuring and building a great deal of expensive hardware to modify only a few parameters. Another option was required.

A compromise was suggested by Navistar which would allow for the minor monitoring and data collection on the engine, while allowing for Navistar to be involved with monitoring modifications and assisting in the development. The suggested course involves using the CAN communication protocol as described earlier. Navistar suggested the purchase of a National Instruments CAN instrumentation board and Labview software. The CAN board allows communication via the ATA data protocol with the ECM. Labview is National Instrument’s data acquisition software. It is useful for creating visual instruments, and as one programs the software it operates as if one were assembling simple electrical circuits with inputs and outputs.

Using these two products together, the following was accomplished:
Engine Parameter Programming

The CAN Card together with the modified ECM calibration allows for specific parameters to be modified as the engine is operating. Additionally, the engine control software sends out specific information regarding how the engine is operating.

Parameters Needed for Pressure Trace Analysis

The injection timing is a little more difficult to measure since it is calculated from several tables and is a combination of an internal variable in the ECM as well as the hydraulic delay of the pressurized oil. It cannot be measured directly because the injector firing as given by the ECM lags the initiation of the injector firing due to delays within the fuel injection system that depend on oil temperature and injection pressure. The variable will be logged, but further data will need to be collected to confirm the actual time of the injector firing. The value recorded from the ECM can be used for simple relative comparison.

3.2.6.3 Engine Control Data Acquisition

Because of the throughput on the existing computer at the test cell, another laboratory computer was reallocated to the test cell to serve as the support for the
National Instruments CAN card. A Labview data acquisition program was written with the support of Navistar to send to and to receive data from the ECM.

To gain further insight into the operation of the engine, instrumentation was developed to enable access the engine control signals, and to eventually enable modification of some of them. The following are the signals acquired from the engine control module:

- Volume Fuel desired
- Dynamic Injection timing desired
- Injection Control Pressure
- Manifold Absolute Pressure
- Barometric Absolute Pressure
- Air Intake Temperature
- Engine Speed
- Engine Oil Temperature
- Engine Coolant Temperature

The observation of these signals allowed for an electronic recording of the steady state operation of the engine. Additionally, the CAN system will allow Injection Control Pressure and Dynamic Injection Timing to be adjusted, as programmed by engineers at Navistar. These two parameters are important, because they control the time the fuel injection begins, and the pressure that the injector must build up to force the required fuel
out of the nozzle. Once the correlation between injection pressure, duration and timing are understood, the CAN system can be used to modify the injection for optimization of performance. At this time, the correlation which exists between these parameters is unclear, and will require further work [104, 111, 112].

3.2.7 Test Fuels

Previous work has examined the effects of increasing the percentage of oxygenate mixed with diesel fuels within several types of engines [14-23, 95]. For this work, comparisons are made between a 5 wt. % and 10 wt. % oxygen via blending of DME in diesel fuel. The baseline diesel fuel properties, as well as test fuel properties are given below in Table 3–3. Because of the difficulty in obtaining the fuel blend properties for DME as a liquid, the properties available in the literature for neat DME are presented.
3.3 Data Analysis

After all testing was completed, the data were analyzed through the use of several methods. As described above, there are three data collection mechanisms, each handled independently and differently. The following section will briefly describe how each data set is handled and what observations are expected.
3.3.1 Steady State Engine Data

To confirm that the engine was operating in a steady state condition, a variety of data were recorded while the engine was running. These data include:

- Speed
- Torque
- Power, (calculated from speed and torque)
- Ambient temperature
- Coolant Temperature
- Oil Temperature
- Exhaust Temperature
- Dyno Water Temperature
- Charge Air Cooler Temperature
- Fuel In Temperature
- Fuel Out Temperature
- Mass Air Flow

The primary use of these data is to confirm that the engine conditions were similar for each fuel blend tested, or within a range of about 2%. These data can also be used to explain emissions results and how they may be related to changes in fuel properties. Additionally, the fuel temperatures are useful in diagnosing the DME behavior in the injector. Charge Air Cooler exit temperature is monitored before the air goes into the
cylinder, after the turbocharger, which indicates air density into the engine. Other temperatures collected can be compared to the data collected from the engine control, as an indication of the calibrated data algorithm. Of these, the engine oil temperature would be most important to note, as it directly affects the injection timing and pressure.

3.3.2 Engine Control Data from the CAN Communication Line

The primary use of these data is to flight record the engine’s steady state operation, as the engine is operated on the different fuel blends. The most important values are injection timing and pressure, and volume of fuel desired. These show how engine operation was changing based on the fuel energy density change. What should also be noted is that changes in oil temperature can also affect the injection timing and pressure. Therefore, it is important to note these and compare at each mode tested. A secondary use for these data will be to determine how to optimize the engine to improve emissions reductions and performance. A final use was to flight record the engine operation when changes were noticed in engine behavior, such as metallic sounds from the injector, and erratic speed changes.

3.3.3 Pressure Trace Analysis

The primary use of pressure trace data is to assess the changes in the combustion process due to the change in fuel type. The changes to be observed include: heat release rate, ignition delay, pressure rise as fuel is injected, and any indication that the fuel may
be leaking past the injector causing pressure pulses where they should not be. The PTrAn software, provided by Optimum Power Technology, will process the data to calculate IMEP (indicated mean effective pressure) and mass fraction burned. The output includes pressure trace and burn rate curves for each cycle of the process, or as an average. Since the processing is for a diesel engine, a double Wiebe function is used to fit a mathematical expression for the mass fraction burned for the two phases of combustion, pre-mixed and diffusion [44, 113]. The general form of the Wiebe function is given in equation 3.1 as:

\[
x(\theta) = 1 - e^{-\left[\frac{\theta - \theta_o}{\Delta\theta}\right]^{am}}
\]

where \( x(\theta) \) is the mass fraction burnt at crank angle \( \theta \)
\( \theta_o \) is the crank angle at the start of combustion
\( \Delta\theta \) is the duration of combustion
and \( a \) and \( m \) are constants that can be varied so that a computed P-V diagram can be matched to a particular engine

Heat release analysis is accomplished through the use of the equation 3.2 given below from Heywood [44]:

\[
\frac{dQ_n}{dt} = \frac{\gamma}{\gamma - 1} p \frac{dV}{dt} + \frac{1}{\gamma - 1} V \frac{dp}{dt}
\]

where \( \gamma \) is the ratio of specific heats \( c_p/c_v \)
The contents of the cylinder are modeled as an ideal gas to arrive at this equation. This calculation is performed separate from PTrAn, as the software is not currently capable of this computation.
Chapter 4

Results and Discussion

4.1 Introduction

The objective of the research was to demonstrate that a Navistar T444E engine could be operated with blends of DME in diesel fuel with minimal engine changes. In this study, the engine was shown to operate with blends of 5 and 10 wt. % oxygen, with an appropriate fuel system design. In this section, detailed results are provided for the effect of the oxygen addition on emissions. Through an uncertainty analysis, based on methods described by Moffat, error bars showing the 95% confidence intervals are presented in each figure showing exhaust emissions [114]. A repeatability study was performed on the steady state operation of the engine for a particular mode to confirm the deviations within the data. Fuel injection timing and fuel injection pressure collected from the engine computer are used to explain the emissions data, and complications during engine operation. Additionally, pressure traces and heat release analysis for one mode are used to show how the ignition timing has changed with the addition of DME to the diesel fuel. Finally, general observations of engine operation will be detailed and explained.
4.2 Fuel Property Data

Fuel property data are available in Table 4–1 to permit comparative analysis of the combustion data. Several data points have been projected as a linear combination because the fuel properties are not able to be determined with available instrumentation.

Table 4–1: Fuel Property Data [30, 41, 92]

<table>
<thead>
<tr>
<th>Fuel Property</th>
<th>ASTM Method</th>
<th>ASTM Spec.</th>
<th>Base Diesel</th>
<th>Neat DME</th>
<th>25 wt. % DME in Diesel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Viscosity, 40°C (cSt)</td>
<td>D445</td>
<td>1.39-4.2</td>
<td>2.2</td>
<td>.25</td>
<td>.92[106]</td>
</tr>
<tr>
<td>API Gravity</td>
<td>D287</td>
<td>API 30</td>
<td>35.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cloud Point (°F)</td>
<td>D2500</td>
<td>&lt;0</td>
<td>4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pour Point (°F)</td>
<td>D2500</td>
<td>&lt;0</td>
<td>&lt;0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flash Point (°F)</td>
<td>D93</td>
<td>125</td>
<td>166</td>
<td></td>
<td>-42</td>
</tr>
<tr>
<td>Calorific Value (BTU/lb)</td>
<td>D2015</td>
<td>19700</td>
<td>19483</td>
<td>12228</td>
<td>17669 *</td>
</tr>
<tr>
<td>Boiling Point (°C)</td>
<td></td>
<td>180-370</td>
<td>-24.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Critical Pressure (bar)</td>
<td></td>
<td>---</td>
<td>53.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Critical Temperature (°C)</td>
<td></td>
<td>---</td>
<td>127</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ignition Temp. (°C)</td>
<td></td>
<td>250</td>
<td>235</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vapor Pressure – bar (°C)</td>
<td></td>
<td>&lt;.001</td>
<td>5.1 (20°C)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bulk Modulus (N/m²)</td>
<td></td>
<td>1.49E+9</td>
<td>6.37E+8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Centane Number</td>
<td>D613</td>
<td>46-48</td>
<td>47</td>
<td>&gt;55</td>
<td></td>
</tr>
<tr>
<td>Density (gm/ml) (20 °C, various P)</td>
<td>D4052</td>
<td>.845-.855</td>
<td>.84</td>
<td>.66</td>
<td>.79 *</td>
</tr>
</tbody>
</table>

* projected
4.3 Particulates

As noted previously, oxygenates traditionally reduce particulate emissions, supposedly through reducing the overall temperatures in the engine, and through chemically changing the composition of the radical pool, thereby inhibiting the formation

---

**Figure 4–1**: Particulate Matter per unit fuel consumed, g/kg fuel

The data indicate a particulate matter reduction for all test modes, as shown in *Figure 4–1*. Using the AVL 8-mode test protocol, the net particulate emission reductions for each mode are found in *Table 4–2*. Because the engine was not able to be operated for Modes 4 and 8 for each additive, the data is presented in a mode by mode
comparison. For most modes, shown in Figure 4–1 particulate reductions were observed on the basis of grams particulate matter emitted per kilogram fuel consumed. As the engine load was increased with increasing engine speed, as in Modes 1 to 4 and Modes 5 to 8, the particulate emission was lowered for each fuel blend as compared to the original baseline diesel fuel, except for Mode 7.

The effect seen in Mode 7 by comparing the baseline diesel to the DME-diesel blends may be due to a change in the injection timing. As shown in Figure 4–2 and Figure 4–3, the injection timing and injection pressure were changing as commanded by the engine.

Table 4–2: AVL 8-mode Weighted Emissions Results per mode, Brake Specific Basis

<table>
<thead>
<tr>
<th>Particulate Emissions Per Mode</th>
<th>Baseline Diesel (g/bhp-hr)</th>
<th>5wt. % oxygen via DME (g/bhp-hr)</th>
<th>10wt. % oxygen via DME (g/bhp-hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3.36</td>
<td>3.44</td>
<td>2.87</td>
</tr>
<tr>
<td>2</td>
<td>.224</td>
<td>.149</td>
<td>.118</td>
</tr>
<tr>
<td>3</td>
<td>.091</td>
<td>.069</td>
<td>.082</td>
</tr>
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<td>4</td>
<td>.209</td>
<td>.095</td>
<td>NA</td>
</tr>
<tr>
<td>5</td>
<td>.339</td>
<td>.255</td>
<td>.214</td>
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<td>6</td>
<td>.137</td>
<td>.128</td>
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</tr>
<tr>
<td>7</td>
<td>.078</td>
<td>.086</td>
<td>.057</td>
</tr>
<tr>
<td>8</td>
<td>.068</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>
control for changing speed and load. This is true for the low as well as the high engine speeds. The trend does not follow what would be expected based on the work by Kajitani and coworkers [85]. Their work showed that as the mean effective pressure increases, which correlates with increasing load for an engine speed of 960 RPM combusting diesel fuel, so do soot emissions based on the Bosch Smoke Number. In their work, the DME emissions were close to zero over the same mean effective pressure range. In the research performed by Ikeda and coworkers on a single cylinder engine operated at 75% of the rated speed, the same pattern was observed for the baseline diesel fuel as in Kajitani’s work. For a DME-diesel fuel blend (40% DME), the Bosch Smoke emissions were shown to increase at a faster rate than the diesel fuel, as the mean effective pressure increased [86]. From an additional set of experiments performed by Kajitani and coworkers, again, the Bosch Smoke number increases as mean effective pressure increases [84]. Additionally, the smoke number decreases with increasing oxygen addition via DME [84]. The study was performed with several blend levels of DME and diesel fuel, and is consistent with the trend of particulate reduction in this thesis.

In the research work presented in this thesis, with increasing content of DME, generally, the soot emissions decrease over a range of speed and load conditions. The trend in particulate emissions reduction using the AVL 8-mode test is similar to the trend observed by Hess and coworkers for each mode, even though the research was conducted with a different multicylinder engine [115].
As has been shown in previous work by Liotta and co-workers, this particulate reduction is due to a reduction in the soot portion of the emission, and would result in a percentage increase in the soluble organic fraction (SOF) portion [11]. This has also been confirmed more recently by Sidhu and coworkers [116], with DME giving the highest SOF.

*Injection timing is relative to the number of degrees before 0 degrees: TDC (Top Dead Center)*
In this research work, the addition of DME into diesel fuel provided a dilution effect of the sulfur content of the final fuel blend. Therefore, it should be noted that the reduction in PM content may also be due to a reduction in the SOF portion of the total PM.

*Figure 4–3:* Engine Injection Pressure

*MFigure 4–4* shows the injection pressure versus engine load. A difference in the injection pressure for a specific engine speed and load can be observed. At higher engine loads, the injection pressure reaches its maximum for this specific engine configuration, regardless of the engine speed.
**Figure 4–4**: Injection Pressure (MPa) versus engine load (ft-lbs)
4.4 Oxides of Nitrogen (NO\textsubscript{x})

Table 4–3 reports the weighted brake specific NO\textsubscript{x} (BSNO\textsubscript{x}) emissions. Figure 4–5 shows that at lower engine loads, the DME-diesel fuel blend causes a decrease in NO\textsubscript{x}. This is seen in Modes 1, 5 and 6, but not for all fuel blends in Mode 2. What is observed in Mode 2 may be the result of an increase in injection pressure without a change in injection timing, causing a change in the fuel plume as it exits the nozzle opening. At higher engine loads (Modes 3, 4 and 7), NO\textsubscript{x} increases. Heywood suggests that NO and NO\textsubscript{2} are a function of equivalence ratio [44]. As equivalence ratio approaches 1 from the lean side, the excess of combustion air decreases, combustion

Figure 4–5: NO\textsubscript{x} Emission Results per unit fuel consumed, g/kg fuel
temperature increases, and NO\textsubscript{2} and NO emissions increase. Above an equivalence ratio of 1, the mixture becomes fuel rich, the pool of O and N decreases and the destruction by CH\textsubscript{x} increases, thus the NO\textsubscript{x} emissions are lower. Equivalence ratio is defined by equation 4.1 below:

\[ \phi = \frac{(F/A)_{\text{actual}}}{(F/A)_{S}} \]

where \( F \) is fuel

\( A \) is air

\( S \) is stochiometric ratio

Also, \( \lambda = \frac{1}{\phi} \)

However, NO\textsubscript{x} is also a function of temperature and time. It is unclear in the data whether the increase or decrease in NO\textsubscript{x} is a function of equivalence ratio, time, temperature, or injection timing.
As can be seen in Figure 4–2, the injection timing of the engine was changing so as to increase the amount of fuel to meet the speed and load conditions. Mode 3 was the only mode where the injection timing did not change, and shows that the NO$_x$ does increase with DME addition. However, injection pressure was increasing so that the required fuel energy could be injected into the cylinder over the same crank angle timing. This increase in injection pressure, as well as the longer fuel spray duration may explain the increase in NO$_x$. Because of DME’s vapor pressure, as the fuel is injected into the engine, the DME may be acting to break apart the diesel fuel into a finer spray, and entraining more air into the edge of the fuel plume. The blended fuel has a lower density
than the diesel fuel, and the viscosity of the fuel blend has also changed. This may be increasing the premixed phase of combustion, causing less of the combustion process to be diffusion-controlled because the fuel is vaporizing and igniting so quickly, with minimal time for mixing. Additionally, more fuel volume is being injected to maintain speed and load conditions. So, there could be some small increase in NO\textsubscript{x} emissions for this reason. In Figure 4–10 the brake specific energy consumption shows that the same amount of fuel on an energy basis is used for each mode, except for Mode 1. Since the fuel is less dense and has lower viscosity, this may be affecting the fuel leaving the injector port and modifying the air entrainment into the fuel jet. Kajitani and coworkers’ data support the scattered increase or decrease in NO\textsubscript{x} emissions depending on engine load [85]. In their work, a comparison of the emissions for various fuel blends was made based on two injection timing points from the engine and showed a significant effect of timing on the emissions for various DME-diesel fuel blends as well as neat DME [85].

However, the data reported here show an interesting phenomenon. The NO\textsubscript{x} emission decreases for the 5 wt. % oxygen, and then begins to increase with additional oxygen content, as shown in Modes 1 and 2. The HEUI fuel injectors use a split shot injection at the lower engine speeds, creating two reaction zones and lowering the overall peak temperature, thus reducing the NO\textsubscript{x}. Choi and Reitz observed that there is a small penalty on the NO\textsubscript{x} emissions when using a split injection strategy (two fuel pulses) with an oxygenated fuel, which could be affecting the results for Mode 1 and 2 for this particular engine [10]. Because the unique multiple fuel injection strategy of the Navistar T444E is especially predominant at lower speeds, the NO\textsubscript{x} reduction could occur due to
improved mixing effect in the cylinder during the combustion event. At higher engine speeds, the data show that NO\textsubscript{x} is lower for DME-diesel blends at light loads, and NO\textsubscript{x} increases more rapidly for the DME-diesel blends as load increases. There are conflicting reports in the literature as to whether oxygenates increase or decrease NO\textsubscript{x} emissions [8, 10, 11, 17, 18]. It is also unclear from these data whether there is a true reduction in NO\textsubscript{x}, because the injection timing and pressure were changing, both of which have been shown to affect NO\textsubscript{x} emissions. Further experiments are needed to determine precisely the effect on NO\textsubscript{x} emissions, through maintaining the injection timing or injection pressure constant while varying the other parameter.

*Figure 4–6* presents the particulate matter vs. NO\textsubscript{x} tradeoff at each test mode. Typically, as PM is reduced, NO\textsubscript{x} is increased, thus making the reduction of either emission a tradeoff for the other. As can be seen for modes with lower loads (Modes 2, 5 and 6), as particulates are reduced, NO\textsubscript{x} is reduced. However, for Modes 3, 4 and 7, an increase in NO\textsubscript{x} with decreasing particulates is observed. In some cases, the PM-NO\textsubscript{x} emissions point shifts toward the origin, which may demonstrate that oxygen addition is a viable means of simultaneously reducing diesel engine NO\textsubscript{x} and particulate matter emissions. However, the changing of the injection timing and injection pressure may also be affecting the reduction or increase in NO\textsubscript{x}. This shift in injection parameters is a confounding effect making interpretation of the emissions more difficult.
4.5 Carbon Monoxide (CO)

Table 4–4 reports the weighted brake specific CO (BSCO) emissions. On a CO per unit of fuel basis, CO emissions increase as the wt. % oxygen is increased, for each mode, except for Mode 3 for the 5 wt.% and 10 wt.% oxygen by DME-diesel blends. The exception for Mode 3 may be due to the observed change in injection pressure, and a combined effect of the change in the fuel blend properties.
In general, CO decreases as load increases, as shown in Figure 4–7. Also, the amount of CO increases with oxygen addition, except for the Modes 3 and 7, which seem to be fairly equal. For Mode 4, there is a decrease in the amount of CO in the exhaust. Oxygenates decompose to form extra CO which “freezes” at elevated levels due to bulk gas quenching during the expansion stroke. This may again support the idea that during the low speed and low load conditions, CO formed during early reaction of the fuel is impeded from conversion to CO₂. The mechanism was postulated by Litzinger and coworkers through the following reactions (equations 4.2 and 4.3), scavenging radicals from the soot creation process [17]:

<table>
<thead>
<tr>
<th>CO Emissions Per Mode</th>
<th>Baseline Diesel (g/bhp-hr)</th>
<th>5wt. % oxygen via DME (g/bhp-hr)</th>
<th>10wt. % oxygen via DME (g/bhp-hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>40.0</td>
<td>51.8</td>
<td>75.93</td>
</tr>
<tr>
<td>2</td>
<td>1.60</td>
<td>1.97</td>
<td>2.75</td>
</tr>
<tr>
<td>3</td>
<td>.560</td>
<td>.689</td>
<td>.671</td>
</tr>
<tr>
<td>4</td>
<td>8.12</td>
<td>7.63</td>
<td>NA</td>
</tr>
<tr>
<td>5</td>
<td>2.33</td>
<td>4.16</td>
<td>6.59</td>
</tr>
<tr>
<td>6</td>
<td>.913</td>
<td>1.344</td>
<td>2.10</td>
</tr>
<tr>
<td>7</td>
<td>.691</td>
<td>.875</td>
<td>.916</td>
</tr>
<tr>
<td>8</td>
<td>.493</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>
\[ H + CO + M \rightarrow HCO + M \quad (4.2) \]

and

\[ H + CO_2 \rightarrow CO + OH \quad (4.3) \]

As explained by Glassman, the conversion of CO to CO\(_2\) would be a function of the size of the hydroxyl radical pool, which does not grow until after all the original fuel and hydrocarbons have been consumed [52]. Since the concentration of hydroxyl radicals is important in the rate of CO oxidation, the additional oxygen atoms from DME may be playing a role in providing excess CO and CO\(_2\), which continue the creation of the hydroxyl radical pool.

In addition, Flynn and coworkers show through kinetic simulations that the addition of the oxygen in the fuel leads to reduced amounts of soot precursors, and larger amounts of carbon leaving the fuel rich premixed combustion zone as CO [117]. Additionally, the work by Flynn and coworkers considers a model of a mixture of 40\% \(n\)-heptane and 60\% DME (molar percentages), which indicates smaller amounts of C\(_2\)H\(_2\), C\(_2\)H\(_4\), and C\(_3\)H\(_3\) formation, and this larger amount of CO leaving the premixed zone. At these molar percentages of fuel, the soot precursors almost disappear.

As seen in the present data, the amount of CO increases with oxygen addition, except for Modes 3 and 7, which seem to be fairly equal. For Mode 4, there is a decrease in the amount of CO in the exhaust, which may not be statistically significant. It seems that above a certain oxygen content, there could be a shift in the kinetic mechanisms,
demonstrated through a drop off in CO. For higher engine loads, regardless of speed, this is shown in the data, which indicates high temperatures to oxidize the CO, and the possibility of an extended heat release time. This is an interesting phenomenon which is also demonstrated in the work by Hess and coworkers [28].

Figure 4–7: CO Emissions per unit fuel consumed, g/kg fuel

The increase in CO has been shown to be a signature of the change in chemical reactions involved in the combustion of an oxygenated fuel. This set of data also shows this to be true. Again, however, changes in the injection timing may be affecting the amount of CO, and skewing the true meaning of the signature. The work by Kajitani shows that the CO emissions actually decrease from the baseline diesel fuel for
increasing amounts of DME in diesel, given an engine speed of 960 RPM and increasing MEP [85].

### 4.6 Hydrocarbons (HC)

The weighted brake specific hydrocarbon emissions for each mode are reported in Table 4–5

<table>
<thead>
<tr>
<th>Hydrocarbon Emissions Per Mode</th>
<th>Baseline Diesel (g/bhp-hr)</th>
<th>5wt. % oxygen via DME (g/bhp-hr)</th>
<th>10wt. % oxygen via DME (g/bhp-hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>34.3</td>
<td>51.4</td>
<td>48.5</td>
</tr>
<tr>
<td>2</td>
<td>.707</td>
<td>1.78</td>
<td>1.54</td>
</tr>
<tr>
<td>3</td>
<td>.211</td>
<td>.953</td>
<td>1.54</td>
</tr>
<tr>
<td>4</td>
<td>.137</td>
<td>.561</td>
<td>NA</td>
</tr>
<tr>
<td>5</td>
<td>.549</td>
<td>3.07</td>
<td>3.18</td>
</tr>
<tr>
<td>6</td>
<td>.216</td>
<td>1.22</td>
<td>1.28</td>
</tr>
<tr>
<td>7</td>
<td>.127</td>
<td>.777</td>
<td>.430</td>
</tr>
<tr>
<td>8</td>
<td>.094</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>
In general, the HC emissions decrease with higher engine loads, as the engine combustion efficiency increases. For all modes, HC emissions increase with oxygen addition, and decrease as engine load increases, as seen in Figure 4–8. For the lower engine speeds, as the oxygen addition increases, the HC emissions increase. However, because very few data points are involved with this figure, these measurements may not be accurate. These data are inconsistent with what has been observed by some previous engine studies. Ikeda and coworkers testing showed that the total hydrocarbon emissions from the DME fuel tests remained roughly the same as the results from the diesel tests [86]. In this work, the two fuels were mixed in the fuel line prior to injection, which ensured a homogeneous mixture. Ikeda’s work also shows that as the DME concentration increases, the exhaust hydrocarbon concentration increases. Kajitani and coworkers testing showed that the HC emissions were roughly the same as the diesel HC emissions, over a range of increasing load and for several injection timings [85]. Therefore, the data in Figure 4–8 may suggest that there are other fuel parameters and injection effects occurring within the system affecting the hydrocarbon emissions.
4.7 Fuel Consumption

Figure 4–9 reports the brake specific fuel consumption (BSFC) for the DME addition. The general trend shows an increase in the amount of fuel required to maintain the same speed and load. This is due to the slightly lower calorific value of the fuel blend, as shown in the Fuel Properties of Table 4–1. However, when fuel consumption is calculated on an energy basis, the energy consumption results are not significantly different.
Below in Figure 4–10 the brake specific energy consumption is shown. The energy content of the fuel was calculated based on the Fuel Property Data in Table 4-1. The DME-diesel blends were calculated based on a linear combination of the fuels.
As described before, the engine was commanding more fuel, which caused the fuel injection pressure, as well as the fuel injection timing to shift. The fuel volume required is also confirmed by the engine control signals, shown in Figure 4–11.
Figure 4–11: Engine control commanded fuel volume desired, mm$^3$
4.8 Pressure Trace and Heat Release Analysis

Cylinder pressure histories of the combustion cycles were collected for each of the fuel blends. This information was compiled by the analysis software as an average of roughly 50 cycles, and plotted in comparison with other fuel blends at the same mode to determine the effect of the fuel on ignition delay in the combustion process. Figure 4–12 shows this comparison for Mode 3, which shows that for the 5 wt.% oxygen by DME one observes a very small premixed combustion phase, with a lower pressure rise, and an early heat release.
Additionally, the peak pressure for 5 wt.% oxygen is slightly lower than the baseline diesel fuel. Looking at an expanded view of this pressure trace in Figure 4–13, the change in the pre-mixed phase of combustion is noted for the 5% oxygen by DME case.

* 0 degrees is TDC (Top Dead Center)
However, the 10 wt.% oxygen by DME curve follows the diesel curve very closely. This could show some relationship to the fuel pressure, since for this curve, the fuel injection pressure was significantly increased. Also, since there is more DME in the blend, this could be the result of the compressibility effect on the injection process, with a delay in the fuel being delivered into the cylinder.

* 0 degrees is TDC (Top Dead Center)
Mode 3 is analyzed for this part of the discussion because it was the only mode that had similar injection timing for diesel fuel and the DME-diesel blends, thereby permitting direct comparison. For all other modes injection timing and injection pressure changed as DME was added, making it difficult to determine and attribute the effect of any single mechanical or chemical change specifically.

Additionally, the heat release for these pressure traces was calculated according to equation 4.4 given by Heywood [44].

$$\frac{dQ_n}{dt} = \frac{\gamma}{\gamma - 1} p \frac{dV}{dt} + \frac{1}{\gamma - 1} V \frac{dp}{dt}$$  (4.4)

where $\gamma$ is the ratio of specific heats $c_p/c_v$

Figure 4–14 shows the following result of this calculation. An interesting observation is the steep peak associated with the 10% oxygen by DME. This again suggests the injection delay-compressibility effect of the DME by about 1 crank angle degree, and then a quick and steep heat release. Additionally, the heating during the delay period is shown on the 10% oxygen by DME, as the curve drops below the axis due to the heat transfer to the walls and to fuel vaporization and heating [44]. Figure 4–2 indicated the injection timing for Mode 3 to be approximately 7 degrees before TDC. Because of other actions occurring in the injector mechanism, specifically the lag time involved with the electronics and the lag time involved with the hydraulic action, it is understood that this is an acceptable number for data analysis and comparison, but not exact. It gives a point of comparison for the chart, in that the ignition delay for the 5 wt.% oxygen by DME
mixture is shorter and the ignition delay for the 10 wt.% oxygen by DME mixture is longer than for the baseline diesel fuel.

Westbrook describes the ignition process of a fuel being dependent on a dominant chain branching reaction that drives the chemical combustion [118]. The ignition process depends on physical, chemical, and mixing and transport dynamics of the situation. To understand the ignition kinetics, it is important to understand the chain branching
mechanisms for the conditions being studied. The most important high temperature reaction for chain branching is the following in equation 4.5:

\[ H + O_2 \rightarrow O + OH \]  

(4.5)

It is important to note that the specific reactions that provide chain branching are dependent on temperature, pressure, and the composition of the reactants over time. At the beginning of a chemical reaction, sufficient time is needed for a system to progress through low and intermediate temperature kinetic pathways until a critical temperature of the system is reached, and a large pool of OH radicals is produced in the system and the system ignites. From laser diagnostic studies performed by Dec and coworkers, a schematic of diesel ignition and combustion has been provided [117]. After the start of injection, a jet of diesel fuel mixes with hot and compressed air. As the air and fuel mix, the fuel-air equivalence ratio is reduced while at the same time the mixture temperature is increasing. Dec and coworkers determined that the mixture starts to ignite when the equivalence ratio is near \( \phi = 4 \). At this point, soot is produced in this fuel-rich pre-mixed ignition flame, and then it is consumed into the diffusion flame. Per the schematic, the diffusion flame is established at the end of the liquid fuel jet. This research supports a direct correlation of soot production to ignition kinetics. Therefore, lowering the post-ignition soot promotion species is a way to reduce soot production.
4.9 General Engine Observations

The engine and fuel system were constructed in such a way as to make sure that the engine would operate as close to steady state as possible. This was important for the comparison of the emission and pressure trace data. The engine seemed to operate, from observed sound and ease of starting, as if pure diesel fuel was being used. However, there were some complications.

From a fuel consumption standpoint, as DME content increased the volume of fuel being commanded from the engine control increased, but reached the maximum limit, as expressed in Figure 4–11. This could be the result of the maximum limit, the fuel injection pressure limit and possibly the injection timing limit, within the engine control program.

Figure 4–15 shows the engine oil temperature. It was important to keep this constant so that the injection timing commands would be constant, since the temperature of the oil determined the hydraulic effect during operation of the fuel injector. Injection timing is a function of several factors including the oil temperature, energized solenoid operation, and timing location commanded by the engine control.
Additionally, the oil temperature and the engine coolant temperature, in Figure 4–16, represent the bulk temperature of the engine. This represents the steady state temperature at each operating mode, and the heat rejection by the engine to the engine coolant and the fuel in the rail.
Further, it was noted after the testing was completed, that the engine would possibly be using the air temperature information to determine the point on the speed and load map to determine injection timing. The sensor was installed in the wiring harness and recording room air measurements, to confirm the engine control was receiving the required voltage input for proper engine operation. However, the sensor was not installed on the engine in the standard location. Figure 4–16 shows the test cell ambient air temperature, which is the data the engine control was receiving.
Figure 4–18 shows the actual intake air temperature, measured on the outlet side of the charge air cooler. This would be the temperature most important to the engine control. It would be important to confirm through the engine manufacturer the specific location and requirement for the sensor.
Finally, a metallic knocking noise was observed from the engine, upon reduction from a high engine speed to a low engine speed, regardless of load. Also, the engine seemed to be close to stall. This could be due to reduced flow rate of fuel in the rail, during a period of high heat rejection from the engine, which could be causing the DME to vaporize in the injectors. Additionally, the viscosity and lubricity of the fuel could be dropping below the ASTM limit, producing a metallic wear “clapping” sound.
4.10 Engine Repeatability Study

A series of data was collected for Mode 4 over the course of several days. The purpose was to confirm that the engine data comparisons were consistent within some error range and to quantify day-to-day variability. Figure 4–19 and Figure 4–20 show the particulates and NO\textsubscript{x} for Mode 4. The data are consistent within 30\%, from center point to center point of the largest and smallest data point.

![Figure 4–19: Particulates, Mode 4 Comparison, g/kg fuel](image)

Along with the above particulate comparison, all filters were analyzed to determine how much error existed in the filter weighting procedures and the use of the humidity chamber. This statistical analysis is presented in Appendix B. There seems to
be some variability in this system, which yields errors above the fixed errors in the system. The cause was believed to be a function of the unstable humidity in the filter weighing chamber, and led to changes in laboratory procedures for filter weighing.

In Figure 4–20, the data are consistent within 3 %, based on the higher point of an error bar to the lowest point of another error bar. From center point to center point of the highest and lowest data point, the error is within 9.5 %.

Variations in the above data can be explained by the higher air consumption for particular modes, and increased exhaust temperature, as shown in Figure 4–21.
Figure 4–21: Exhaust Temperature, Mode 4 Comparison, °C
Chapter 5
Summary, Conclusions, and Recommendations for Future Research

5.1 Summary and Conclusions

Manufacturers of diesel engines and vehicles are facing continued pressure due to governmental regulations to reduce the engine and vehicle emissions. Regulations set for 2004 and 2007 are currently out of reach with current technology [4, 119]. Therefore, manufacturers are resorting to adapting a total system approach for emissions reductions, implementing pre-combustion, in-cylinder, and post-combustion technologies to meet these emission levels. The focus of this research involved the study of a pre-combustion approach, via fuel modification.

Dimethyl ether (DME) has been shown to reduce particulate emissions when combusted in a compression ignition engine [86]. Dimethyl ether is considered an oxygenated compound that can be blended with or used as a fuel. However, when using DME as a fuel, there are some challenges that need to be overcome to achieve optimal engine operation. These include changes in fuel injection quantity and timing, as well as reduction in the lubricating properties of the fuel in the injectors.

In order to accommodate the need for lubricity in the fuel injectors, it was theorized that the DME could be mixed with diesel fuel in order to maintain the quality of the fuel
needed by the fuel system. The minimal quality of fuel is established by the ASTM specification for No. 2 diesel fuel. However, what is not known is what the minimal amount of lubricating quality that the fuel injectors, in particular, can accommodate and for what duration of time.

Mixing the DME with the diesel fuel, provides what is typically referred to as an oxygenated fuel blend. Dimethyl ether contains oxygen within its molecular structure. Many researchers have shown that mixing an oxygenate with a diesel fuel will provide reduction of particulate matter in the exhaust [7, 8, 11, 95]. Since DME has been shown to reduce particulate emissions when used in neat form, it was theorized that there would be some emissions reductions when mixed with diesel fuel.

A Navistar 7.3L Turbodiesel engine and fuel system were modified for the purpose of utilizing pressurized DME and diesel fuel blends. Instruments were placed in the engine and fuel system so that data could be flight recorded for real-time observation, as well as, for future analyses. The objective of the data collection was to provide information which could be used to modify the engine from it’s production configuration to improve performance and emissions with the various fuel blends.

Research continues in testing and reviewing the effects of oxygenates, specifically DME diesel fuel blends, on the composition of emissions from diesel engines. The results from this research lead to the following conclusions:

- DME-diesel blends can be used to fuel a compression ignition engine, using of a pressurized fuel system. The engine operates in a similar manner with the DME-
diesel fuel as with the diesel fuel, as reviewed in the engine stability data. Changes in combustion peak pressure were not audibly noticed, but were shown using pressure trace analysis.

- DME has been shown to reduce particulate emissions from a DI diesel engine. With the addition of oxygen contained within the fuel, additional CO and CO\textsubscript{2} is present in the combustion process. This has lead researchers to conclude this higher level of CO prevents the formation of soot precursors in the pre-mixed flame, which then reduces the formation of particulates.

- It has also been observed that small NO\textsubscript{x} reductions or increases occur over the engine operation map. Multiple injections at lower loads and speeds may explain this data. For higher speeds and loads, this may be explained due to advanced injection timing and injection pressure. No conclusion can be drawn from this data as to the effect of oxygen on the formation of NO\textsubscript{x}.

- The engine control module and the calibration for the Navistar 7.3L T444E engine are sensitive to changes in fuel properties, resulting in inconsistent injection timing and injection pressure for each of the fuel blends tested. This can be explained by the changing energy content and density of the fuel blends. The end result makes comparisons in emission results extremely difficult.

- Upon this initial testing, no determinations of the effect of the DME on the durability of the engine can be addressed. None of the data collected suggests that the testing of
the engine has degraded the injector operation. The engine has been operated for about 100 total hours, with at least 5% DME content in the fuel for half of that time.

- Because the pressurized system requires the fuel to enter the injector at a lower temperature than would the production style engine, the effect of the lower temperature fuel may result in a change in the emissions.

- The pressure trace analysis for Mode 3 revealed that there is some effect of DME on the combustion peak pressure. Additionally, the heat release analysis showed potentially some effect of the compressibility of the fuel, as a result of the blends or the temperature of the blends as they enter the cylinder.

The conclusions from the research completed thus far lead to recommendations for future work in order to improve the engine operation, by further separating the variables affecting the performance and emissions of the engine.

5.2 Recommendations for future research

Although it has been shown that DME can be used in combination with diesel fuel in a compression ignition engine, the fuel blend and engine operation can be further optimized for improved emissions and long term durability of the engine. This involves several factors including, increasing the DME content in the blend, up to the point where the engine can still produce acceptable speed and load, with improved emissions for an acceptable lifetime of the engine and fuel injectors. It should be noted that since the fuel
is being used within an engine, there exists not only chemical effects of the fuel and changes in the fuel as the blend is altered, but also mechanical effects by the engine, some which are a result of changing fuel blends. Therefore, further study of the engine emissions are a function of understanding the various chemical and mechanical effects.

This research work aided in identifying many of the chemical and mechanical effects. Some of the mechanical effects are inherent to the engine and cannot be directly altered. The optimization of the engine is influenced by the following factors:

- DME content in diesel fuel blend
- Ignition delay
- Injection pressure and timing for the fuel blend
- Engine emissions
- Internal injector wear
- Injector o-rings
- Compressibility of the fuel blends
- Vaporization of DME in the cylinder/liquid length of the DME-diesel fuel blend
- Injector nozzle size and flow rate
- Cooling required to maintain the fuel in a liquid state

Changes in the fuel blends relate to changes in the fuel blend properties. Therefore, above a certain blend level, there is a requirement to adjust the engine control system to achieve improved ignition delay in conjunction with the correct engine timing.
To perform this optimization, it would be important to know how the fuel properties are changing that affect the ignition delay. Since the cetane number is increasing with increasing DME content, this would be one factor. Also, and more significantly, the compressibility of the fuel is changing with increasing DME content, which has an unknown relationship to ignition delay. Additionally, the fuel blends have an unknown effect on the engine and injector wear and wear rate, including both metallic and polymeric materials. Therefore, the future work should be divided into several sections:

1. **Fuel Property Studies**

   To aid in explaining the changes in heat release of the DME-diesel blends, it is important to understand how the DME-diesel fuel blends change in compressibility as the DME content is increased over a range of temperatures. Additionally, it is unclear whether a change in viscosity or lubricity with increasing DME content is affecting the wear and wear rate in the injectors.

2. **Engine Optimization for Performance and Emissions**

   To determine the optimal conditions for engine operation, an optimal network of conditions should be established to direct the optimization of the engine for performance and emissions. This would be accomplished both experimentally and through computational analysis, and confirmed experimentally.
3. Engine Optimization for Durability

To determine the ability of the engine and fuel system to be durable and label it as durable, two specific types of data should be collected. One, injector o-rings should be tested in the fuel to determine if they are able to withstand the effect of DME. Two, an injector test stand should be used to operate the injectors for an extended period with the DME-diesel fuel blend and inspect them after the testing. Such testing would be comparable to ASTM methods.


Appendix A

Statistical Control and Analysis of Performance and Emissions Experimental Data

A.1 Introduction

The following section describes the statistical controls and statistical analysis of the data collected for this research. A basic overview of experimental error will be discussed, followed by a generic description of statistical analysis concepts. Finally, a discussion of the experimental analysis techniques specific to this research will be discussed, divided into two parts. The first part will describe the data sample collection from the pieces of equipment used, and how they were programmed to collect samples to provide a statistical normal distribution of the data. The second part will describe the spreadsheet used to produce the graphs of the emissions data, including the error bars for the data. Some assumptions that were made to compile this data will be explained.

A.2 Causes and Types of Experimental Error

Some form of experimental analysis must be performed on all data. The analysis may be simplistic, or a more complex analysis of the errors involved, in an effort to match results with fundamental physical principles. In the process of setting up any
experimental equipment, an analysis of the data is important for several reasons. These reasons include:

- to determine and quantify the errors and establish areas where maximum reductions can be achieved,
- to establish an appropriate number of samples needed to reduce standard deviations to the acceptable confidence level and gain precision, and
- to give validity to experimental measurements [120].

Typical types of errors which cause uncertainty in experiments include apparatus or instrumentation error, fixed errors (systematic errors), human error, and random errors. Instrumentation errors are those involved with setup and design mistakes. Fixed errors are considered errors which occur for unknown reasons, with the error being roughly the same amount for each repeated reading. Human error would involve a mistake being made in reading and recording a direct observation, and could be considered a reproducibility issue if various people are involved in data collection. Random errors involve random events, which may or may not be attributed to the other three types of errors [120].

### A.3 Basic Statistical Analysis Concepts

To be able to draw conclusions from a data set, a quantitative description of the data is needed. The first step is to determine the uncertainty involved with the data, as a function of the results and independent variables which make up the data set. This
method is also known as Root Sum Square analysis. Next, these results would be used, in addition to the standard deviations and means for the individual data sets, to create the error bars describing the error in the experimental results. The following section will further explain the equations involved.

A.3.1 Uncertainty Analysis

When trying to determine the uncertainty in a set of measurements, the odds for the particular uncertainty would also be an important factor to add further specification to the uncertainty estimate. For example, knowing the calibration history of a device versus not knowing the calibration history may change the precision odds for the measurement. If a set of measurements is made and the odds for the uncertainty in the primary measurements are known, uncertainties in the calculated results can be estimated. This result is sometimes referred to as the Root Sum Square method. The result $R$ is a given function of the independent variables $x_1, x_2, x_3, \ldots x_n$. Then, the following is true in equation [A.1]

$$R=R(x_1, x_2, x_3, \ldots x_n) \quad (A.1)$$

If $w_r$ is the uncertainty in the result and $w_1, w_2, w_3, \ldots w_n$ is the uncertainty in the independent variables, given that the uncertainties in the independent variables are given with the same odds, then the uncertainty in the result having these odds would be given in
equation [A.2] The relationship from this equation can be used to calculate the percent uncertainty [120] [114].

\[
w_r = \left[ w_1 \frac{\partial R}{\partial x_1}\right]^2 + \left( w_2 \frac{\partial R}{\partial x_2}\right)^2 + \left( w_n \frac{\partial R}{\partial x_n}\right)^2 \right]^{1/2}
\]  

(A.2)

A.3.2 Statistical Analysis and Confidence Interval

Typically, a 95% confidence level is utilized during error analysis testing. This means that the \( \alpha/2 \) (\( \alpha \), the confidence interval) equals .025 and is used in the hypothesis test. Depending on the number of data points collected, the Central Limit Theorem may or may not be valid. In the case where it is not valid (sample size less than 30 samples), a “t” test approximation is used. The “t” test establishes a bell shaped approximation of the data, based on the degrees of freedom present in the data. Using the “t” test yields equations to calculate the confidence intervals and the error, given in [A.3] The error involves one side of the interval divided by the sample mean.

\[
\bar{x} \pm t_{\alpha/2, n-1} \cdot s / \sqrt{n}
\]  

(A.3)

Where:
\( t_{\alpha/2, n-1} \) is the number on the measurement axis for which the area under the t curve with \( \nu \) degree of freedom to the right of \( t_{\alpha/2, n-1} \) is \( \alpha \), and \( \nu \) is \( n-1 \).

\( \nu \) is the degrees of freedom, \( n-1 \)

\( s \) is the standard deviation of the sample population

\( n \) is the number of data points

\( \bar{x} \) is the average sample population

The value of the t test varies depending on the value of \( \alpha \), the confidence interval, and the degrees of freedom, \( \nu \) [121]. A typical t curve is shown below in Figure A–1.

![t-curve](image)

*Figure A–1: t-curve*

If several data groups have been collected, an analysis of variance (ANOVA) can be used to determine the qualitative response from the data. A Single-Factor ANOVA is used to compare more than two populations or treatment means. If the comparison yields nothing significant, a multiple comparison procedure is then used [121]. This will
become important when performing the comparison on the Mode 4 engine data found in Appendix C.

A.4 Summary of Experimental Error Analysis

The list of equipment used in this experiment is given in Table A–1 with the manufacturer’s listed error, or allowed minimum calibrated error. Additionally, the table shows how the data is collected and how many samples are collected for each variable monitored. The data collected by the Windows-based computer connected to the Modicon PLC sampled the data 33 times, and used the average of the samples for each data point. This allowed for the assumption of a normal distribution for a sample size above 30 samples, and the use of the standard normal curves for determining confidence intervals. The data points were collected throughout the entire engine operation for each mode, which was usually 60 to 80 minutes of data per mode. From this group of data points, the average of every 30 points were used in the excel spreadsheet, giving 5 points from which the standard deviation and 95% confidence interval was calculated.
**Table A–1: Engine Laboratory equipment**

<table>
<thead>
<tr>
<th>Equipment Type</th>
<th>Variables Collected</th>
<th>Error in Data Point</th>
<th>Number of data points per time</th>
<th>Assumptions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sierra Instruments BG-1 Micro Dilution Test Stand</td>
<td>Particulate Filters</td>
<td>Standard Deviation between Filter samples; causes unknown</td>
<td>1 sample each 5 minutes; 5 samples total each engine mode</td>
<td>Dilution flow is constant, Temperature difference between filters and modes is negligible; Temp. maintained below 52°C</td>
</tr>
<tr>
<td>Sartorius M3P Precisions Microbalance (located in environmental chamber)</td>
<td>Weight Particulate Filters</td>
<td>± 0.005 gm</td>
<td>Filters weighted 3 times pre- and post collection</td>
<td>Humidity chamber is constant 45% relative humidity</td>
</tr>
<tr>
<td>Nicolet Magna 550 FTIR Spectrometer</td>
<td>NO,NO₂,CO,CO₂</td>
<td>Calibration curve for 1% or less error</td>
<td>5 samples throughout engine test run</td>
<td>Stable gas flow rate, clean spectrometer windows; negligible interferences at wavelengths; cell temp. and pressure constant</td>
</tr>
<tr>
<td>California Analytical HFID 300 THC analyzer</td>
<td>Total Hydrocarbons</td>
<td>Resolution 0.01ppm, Repeatability ± 0.05% of full scale</td>
<td>2 per engine mode</td>
<td>Constant clean sample flow</td>
</tr>
<tr>
<td>Rosemont Analytical Oxygen analyzer</td>
<td>Percent Oxygen</td>
<td>± 1% per daily calibration; Resolution 0.01%</td>
<td>5 per engine mode</td>
<td>Constant flow through system</td>
</tr>
<tr>
<td>450 hp Eaton eddy current dynamometer</td>
<td>Engine Load control</td>
<td>Zero:1% of rated Repeatability:0.05% of FS; temp.effect 0.002% of FS/°F</td>
<td></td>
<td>Coolant temp. causes no variability in load; Non-linear 0.03% of FS* calib.; Hysteresis 0.04% of FS</td>
</tr>
<tr>
<td>Navistar Engine Computer connected to National Instruments CAN (Controller Area Network) board</td>
<td>Injection timing and pressure, temperatures, volume fuel desired</td>
<td>Unknown</td>
<td>800 per 10 seconds; using average of 800 points</td>
<td>Engine is steady state, so that data is steady state</td>
</tr>
<tr>
<td>Kistler 6125 Pressure sensor, charge amplifier, Visual Basic program</td>
<td>Pressure, crank angle and TDC trigger</td>
<td>Sensitivity :=15 pC/bar; Linearity: &lt;0.5%FSO</td>
<td>1 data point per crank angle, average of 50</td>
<td>Load created by water flowing over dyno at zero load factor is negligible.</td>
</tr>
<tr>
<td>Torque meter</td>
<td>Torque load on engine</td>
<td>Repeatability 0.05% of full scale</td>
<td>Average of 33 cycles/data point, appx. 1 point/min</td>
<td></td>
</tr>
<tr>
<td>Modicon (PLC) Programmable Logic Controller; Visual Basic program by DDE (Dynamic Data Exchange)</td>
<td>Engine and fuel temperatures, speed, load, mass air flow</td>
<td>Resolution of 0.25°C; Error of 0.4% over the range of 0 - 60 °C</td>
<td>Average of 33 cycles per data point, appx. 1 data point per minute</td>
<td>PLC is adequately providing temperature reference</td>
</tr>
<tr>
<td>MPSI Prolink Service tool</td>
<td>Turbo Boost Pressure</td>
<td>Unknown sampling rate</td>
<td>5 per engine mode test</td>
<td></td>
</tr>
<tr>
<td>Delphi Mass Air Flow (MAF) meter calibrated with a Marion Laminar Flow Element</td>
<td>Voltage signal converted to MAF</td>
<td>Calibration curve has R² of 1</td>
<td>5 per engine mode</td>
<td></td>
</tr>
<tr>
<td>Omega K type Thermocouples</td>
<td>Temperature</td>
<td>2.2°C Error or 0.75% above 0°C</td>
<td>Average of 33 cycles/data point, appx. 1 point/min</td>
<td></td>
</tr>
<tr>
<td>Sartorius Scale</td>
<td>Fuel</td>
<td>Resolution 0.2gm</td>
<td>5 /engine mode, for 5 minutes each</td>
<td>Fuel reading taken when scale reading changed</td>
</tr>
</tbody>
</table>

* FS- Full Scale
A.5 Summary of Excel Spreadsheet Calculations

The following section will describe the equations used to analyze the emissions data, including any assumptions used. Additionally, a description of how the error bars for the data sets were created will be discussed.

To aid in simplifying the analysis process, some assumptions are made regarding the exhaust. First, since the dilution ratio for the particulate sampling with the BG-1 was 8:1 at 150 SLPM (standard liter per minute) total flow rate, the true flow of exhaust particulate was 150 divided by 9, or 16.667 l/min of exhaust. This factor, which represents the flow of exhaust for the particulate measurement at that mode, was the divisor of the total exhaust flow in l/min, and was used to obtain a multiplier to calculate the total particulate in grams per hour. All other emissions measurements used the density of the measured gas divided by the total exhaust flow for the mode to get the measured gas in grams per hour. The density for each measured gas was calculated using the exhaust gas temperature. The density of the exhaust gas was assumed to be 1.2 kg/m\(^3\) at 300 K, and corrected for the specific measured exhaust temperature for the mode.

Primarily, the data sets establishing the steady state operation of the engine for each mode consist of 5 groups. Within these five groups, which includes engine speed, power, torque, and various temperatures, are 30 or more data points which have been collected from the data acquisition system. Each of these data points is from an average of 33 points collected by the system. The standard deviation across the points for each of the five groups is determined from the set of 30 points. This data is used to create the error bars, and consists of items such as speed, load and air flow. The emissions data is
collected 5 times for each mode. Therefore, the “t” test is set up for a 95% confidence interval, with a sample size of 5. The error interval is then created based on the standard deviation in the data, and a root sum square of the equipment error for the steady state operation of the engine.
Appendix B

Statistical Analysis of the Particulate Matter Data Collection

The following section describes the process by which the particulates were measured and evaluated in this research. Extra attention was placed on this portion of the experiment, to provide for quality and repeatability in the data collection. This section contains a description of the procedure, the raw data, and general comments and recommendations about the data.

Particulate samples were collected for each AVL-8 mode, and each fuel blend. Test protocol requires that 5 particulate samples be collected, for statistical quality of the data. The spreadsheet used to analyze the data calculates error bars based on 95% confidence intervals, for a five sample set.

ISO 8178 Part 2 calls for the particulate filters to be pre-conditioned and weighed prior to use, in a controlled environment which is kept to 45% humidity, ± 2%. The scale used to weigh the filters gives a standard deviation across six weighing trials of .005 gm.

It has been the typical procedure in the lab to weigh the filters one time before and one time after particulate collection. For the purpose of trying to minimize the error in the weight of the particulate filters, since the fuel blends were expected to reduce the amount on the filters, the filters were weighed three times before and three times after the collection. The same person weighed the filters, and waited for the scale to stabilize for
10 seconds between each weighing. Also, the scale was tared for each weighing. The average weight was used as the result of the analysis.

The data from this project was collected and put into a spreadsheet for some basic statistical analysis. Preliminarily, the difference between the three samples before and after were compared against the .005 gm standard deviation for the scale weighing range. The spreadsheet gives a True or False indication that the difference is within the standard deviation constraint. The .005 gm was determined by the manufacturer to be the standard deviation from the weighing of the same filter six times. Figure B–1, on the following pages, shows the filter weightings before the particulate collection. BF refers to the before weighing. AF refers to weighing after sample collection. The “Problem” column is used to indicate that the standard deviation being above .005. True in the column would indicate that it is above .005.

Many True indications were observed. This could indicate that there are some other aspects contributing to the error of the data than the scale. The .005 number is much lower than the true error of the entire process.

There are several reasons to explain the results of this data. Since the humidity chamber is to be kept at the designated humidity, the humidifier had to be cycled to keep the humidity constant. The affect of the humidity on the filter is unknown, and not quantified. Another explanation could be malfunction of the humidity sensor. Additionally, particulates could be blown off or somehow removed by static electricity, reducing the weight of the filters.
As recommendations, the following are a few suggestions which may improve the accuracy of this data, even though the source of the inaccuracies has not been quantified. The first effort should be to determine how the standard deviation affects the accuracy of the particulate measurements, to further determine the range in which there is 95% confidence based on the sample size. Some effort should be made to quantify the moisture absorption tendency of the filters, to determine the effect on the filter weight. Also, the humidity chamber environmental controls should be inspected, as well as to confirm the correct operation of the chamber sensors. Further, the device to remove static electricity should be regularly replaced. Also, a ground strip connected from the operator’s wrist to bench top could be put in place as an additional method to remove any possibility of static build up on the filter and in the chamber.
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<th>BF weight 2</th>
<th>BF weight 3</th>
<th>Std dev BF</th>
<th>Problem</th>
<th>AF weight 1</th>
<th>AF weight 2</th>
<th>AF weight 3</th>
<th>Std dev AF</th>
<th>Problem</th>
</tr>
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*Figure B–1: Particulate Filters*
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| dd349m 7  | 4   | 330.078 | 330.088 | 330.085 | 0.0051 | TRUE | 330.809 | 330.814 | 330.81  | 0.0026 | FALSE |
| dd349m 7  | 5   | 329.443 | 329.443 | 329.443 | 0.0000 | FALSE | 330.201 | 330.204 | 330.204 | 0.0017 | FALSE |

(Cont.)
| dd449m 1 | 1 | 321.024 | 321.027 | 321.016 | 0.0057 | TRUE | 321.75 | 321.754 | 321.747 | 0.0035 | FALSE |
| dd449m 1 | 2 | 328.891 | 328.889 | 328.888 | 0.0061 | TRUE | 329.648 | 329.634 | 329.642 | 0.0070 | TRUE |
| dd449m 1 | 3 | 331.501 | 331.517 | 331.506 | 0.0082 | TRUE | 332.195 | 332.19 | 332.187 | 0.0040 | FALSE |
| dd449m 1 | 4 | 324.079 | 324.07 | 324.074 | 0.0045 | FALSE | 324.814 | 324.808 | 324.801 | 0.0065 | TRUE |
| dd449m 1 | 5 | 322.749 | 322.749 | 322.749 | 0.0012 | FALSE | 323.482 | 323.449 | 323.473 | 0.0067 | TRUE |
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| dd449m 2 | 3 | 327.518 | 327.511 | 327.505 | 0.0065 | TRUE | 328.178 | 328.176 | 328.171 | 0.0036 | FALSE |
| dd449m 2 | 4 | 327.5 | 327.497 | 327.485 | 0.0079 | TRUE | 328.123 | 328.129 | 328.123 | 0.0035 | FALSE |
| dd449m 2 | 5 | 324.737 | 324.731 | 324.728 | 0.0046 | FALSE | 325.403 | 325.398 | 325.4 | 0.0025 | FALSE |
| dd449m 3 | 1 | 333.535 | 333.532 | 333.519 | 0.0035 | FALSE | 334.427 | 334.433 | 334.429 | 0.0031 | FALSE |
| dd449m 3 | 2 | 335.459 | 335.461 | 335.454 | 0.0036 | FALSE | 336.141 | 336.143 | 336.143 | 0.0012 | FALSE |
| dd449m 3 | 3 | 335.211 | 335.193 | 335.2 | 0.0091 | TRUE | 335.928 | 335.928 | 335.927 | 0.0066 | FALSE |
| dd449m 3 | 4 | 330.493 | 330.486 | 330.485 | 0.0044 | FALSE | 331.145 | 331.144 | 331.146 | 0.0010 | FALSE |
| dd449m 3 | 5 | 331.24 | 331.252 | 331.252 | 0.0069 | TRUE | 332.016 | 332.022 | 332.004 | 0.0092 | TRUE |
| dd449m 5 | 1 | 323.961 | 323.954 | 323.959 | 0.0036 | FALSE | 324.809 | 324.8 | 324.799 | 0.0055 | TRUE |
| dd449m 5 | 2 | 322.117 | 322.11 | 322.116 | 0.0032 | FALSE | 323 | 322.989 | 323 | 0.0064 | TRUE |
| dd449m 5 | 3 | 330.464 | 330.467 | 330.464 | 0.0017 | FALSE | 331.382 | 331.37 | 331.376 | 0.0060 | TRUE |
| dd449m 5 | 4 | 334.549 | 334.556 | 334.55 | 0.0039 | FALSE | 335.422 | 335.409 | 335.426 | 0.0089 | TRUE |
| dd449m 5 | 5 | 327.562 | 327.558 | 327.546 | 0.0083 | TRUE | 328.387 | 328.398 | 328.381 | 0.0086 | TRUE |
| dd449m 6 | 1 | 333.945 | 333.933 | 333.91 | 0.0076 | TRUE | 334.497 | 334.491 | 334.492 | 0.0032 | FALSE |
| dd449m 6 | 2 | 331.741 | 331.737 | 331.73 | 0.0056 | TRUE | 332.298 | 332.293 | 332.288 | 0.0050 | TRUE |
| dd449m 6 | 3 | 331.211 | 331.119 | 331.122 | 0.0015 | FALSE | 331.746 | 331.74 | 331.738 | 0.0042 | FALSE |
| dd449m 6 | 4 | 330.745 | 330.74 | 330.737 | 0.0040 | FALSE | 331.386 | 331.38 | 331.378 | 0.0042 | FALSE |
| dd449m 7 | 1 | 334.567 | 334.569 | 334.581 | 0.0076 | TRUE | 335.062 | 335.064 | 335.056 | 0.0042 | FALSE |
| dd449m 7 | 2 | 334.411 | 334.412 | 334.4 | 0.0067 | TRUE | 334.875 | 334.866 | 334.87 | 0.0045 | FALSE |
| dd449m 7 | 3 | 331.035 | 331.035 | 331.027 | 0.0046 | FALSE | 331.547 | 331.549 | 331.551 | 0.0020 | FALSE |
| dd449m 7 | 4 | 333.396 | 333.398 | 333.387 | 0.0010 | FALSE | 333.973 | 333.967 | 333.967 | 0.0035 | FALSE |
| dd449m 7 | 5 | 334.197 | 334.192 | 334.183 | 0.0071 | TRUE | 334.702 | 334.699 | 334.697 | 0.0025 | FALSE |
In summary, the particulate data collection distinguished the following:

• there is variability between the weight of each of the same sample, and the causes at this time are unclear

• reducing error in the measurements may not be improved by increasing particulate collection time to increase particulate quantity, or reducing raw filter weight, but may be improved by increasing scale resolution and improving control over the weighing environment.
Appendix C

Mode 4 Repeatability Study

The following section presents all charts and graphs from the data analyzed as part of the repeatability study of Mode 4. This repeatability study was performed in order to quantify the variance of the data, day to day, for the same test conditions. The data was statistically analyzed as described in Appendix A. A portion of the figures were shown in Chapter 4 Results and Discussion, and will be repeated here for a complete picture of the data. Analysis of variance will be used to determine if there is statistical significance to the differences in data points. Some comments and general recommendations from the data will be presented.

*Figure [C–1]* shows the particulate emissions in grams per kilogram fuel consumed.
Even though the error bars are small, this chart indicates that there are some deviations between the samples. The relative error was 1.8 to 2.4 % across the samples in the figure above. Mode 4 was chosen because it is the mode which produces the most particulate emissions for the 5 minute sampling time, and could have the greatest chance of reducing the number of significant figures needed for measurement by the scale. However, the measurements of fuel and air consumption, show some instabilities as well. The particulate measurement is directly impacted by the accuracy of the fuel measurement.

*Figure C–1* shows the particulate emissions comparison, g/kg fuel

*Figure C–2* shows the NO\textsubscript{x} emissions in grams per kilogram fuel consumed.
The NOx data seems to be fairly stable, and within the range of error bars for the other samples. The relative error was 2-5% for the samples in the figure above. The NO and NO2 calibration curves for the FTIR were fit with a 1% or less error. Again, error in the fuel measurement contributes to the error in the NOx results.

*Figure C–2*: NOx Emissions, Mode 4 comparison, g/kg fuel

The NOx data seems to be fairly stable, and within the range of error bars for the other samples. The relative error was 2-5% for the samples in the figure above. The NO and NO2 calibration curves for the FTIR were fit with a 1% or less error. Again, error in the fuel measurement contributes to the error in the NOx results.

*Figure C–3* shows the CO emissions in grams per kilogram fuel consumed.
The CO calibration curve from the FTIR was fit with less than 1% error. Based on the data in the chart, it seems that some variability exists with the CO reading from the FTIR, but taking into account the error bars, the CO value seems to be consistent. The relative error in the above figure is 3.4 to 11.5% across the samples.

*Figure C–3:* CO Emissions, Mode 4 comparison, g/kg fuel

Figure C–4 shows the hydrocarbon emissions in grams per kilogram fuel consumed.
The hydrocarbon data is based on only the average of two to three samples per mode, which were collected as average numbers from the Heated FID analyzer. The ability to only collect two to three samples was a function of the heavy particulates of Mode 4 clogging the sample lines. Changes in the sampling systems would be recommended so that a greater surface area exists to allow for sampling over a longer time, and therefore giving a more stable reading of hydrocarbons. The relative error in the above figures was 9.4 to 13% across the samples.

*Figure C–4: Hydrocarbon Emissions, Mode 4 comparison, g/kg fuel*
Figure C–5 shows the Brake Specific Fuel Consumption in grams per kilowatt-hour.

It is difficult to determine from Figure C–5 whether the error in the BSFC arises from the fuel consumption measurement or the engine’s operation. The fuel scale being used for this experiment has a resolution of 0.2 kg. During the experiment, the readings were taken so that when the scale changed to the next division, time and fuel were recorded. The error bars in Figure C–5 refer to measurement variability only. The accuracy of the scale has not been included. The relative error is less than 0.1% for the samples in the figure. However, the raw error in the fuel consumption in grams/hour was 8-25%.
Figure C–6 shows the air consumption in grams per minute. The air consumption trend seems as constant as the fuel consumption trend.

The air consumption is based on a voltage signal from a mass air flow (MAF) device. The device was calibrated with a laminar flow element (LFE). Due to errors in the device generating a voltage signal into the main computer logging system, the data was recorded 5 times during each mode. The voltage was recorded from a digital voltmeter, giving a reading up to two decimal places for a voltage value between 6 and 10 volts.
The voltage signal was converted to the flow rate, via the calibration. The mass air flow calibration was fit very well, with a correlation coefficient $R^2$ close to 1. An additional source of error exists between the LFE and the MAF device, in that the LFE which is intended to straighten the air flow may also be blocking the air flow, and thus reducing the signal from the MAF device. The LFE is used for calibration only, and does not reside in the flow passage during engine testing.

*Figure C–7* shows the exhaust temperatures. The exhaust temperature follows the same consistent trend that the fuel consumption does.
Finally, the data indicates that further investigation of the data to confirm the stability of the engine throughout the time the data was collected, which includes speed and load. Additionally, ambient temperature and humidity should be recorded to assess correlation with variations in the engine measurements. Comparisons between the ambient air temperature and charge air cooler temperature could also be helpful in determining factors which could reduce error in the experimental set-up.

*Figure C–7: Exhaust Temperature, Mode 4 comparison, °C*
Appendix D

Controller Area Network (CAN) Data

The following section briefly discusses the principles of the Controller Area Network. In addition, this section will show the types of data are collected from the Navistar engine control (Part Number: 1807349C4).

D.1 Definition of Controller Area Network

CAN, Controller Area Network, is a recommended practice intended for light and heavy duty vehicles for on and off-road applications. The purpose of CAN is to provide an open interconnect system for electronic systems, which then allow Electronic Control Units to communicate with each other using standard architecture. The network management structure is outlined in SAE J1939 and several associated documents. This higher level architecture assigns priority to engine functions, and alert status to changes in engine operation, and engine faults. The documents describe message types, lengths, and structures, in an attempt to provide consistency for programmers. The standard messages can then be used to provide generic service diagnostic tools [122, 123].

For the purpose of these experiments, it was important to understand the basics of CAN operation, so that the message information could be decoded. The CAN system for this experiment was developed by Navistar engineers. It was designed to send out information packages when triggered with a signal message. Also, it was set up for
messages to be sent via the CAN to the microprocessor. These messages contain instructions to update the set point values for engine operation, specifically, injection timing and injection pressure.

### D.2 CAN Data

The data collected by the CAN bus is discussed in Chapter 3, and several charts are shown in Chapter 4. In this section, several other charts will be shown with the balance of the data collected. As discussed in Chapter 3, the CAN data was collected for 10 seconds, with approximately 800-900 points of data for each logged CAN channel being collected in that time frame. (The list of CAN channels is found in Chapter 3.) The charts that were created constituted the average values during that time frame. However, maximum and minimum values were also calculated for each engine setting, as seen in Table [D–1]
Table D–1: CAN data table: Baseline Diesel tests

<table>
<thead>
<tr>
<th>Test Case</th>
<th>Engine Coolant Temp</th>
<th>Engine Oil Temp</th>
<th>Engine Speed</th>
<th>Engine Load</th>
<th>Acceleration</th>
<th>AIR Brake Temp</th>
<th>Single Speed</th>
<th>Single a/g Temp</th>
<th>Dynamic Injection</th>
<th>Engine Injection Control</th>
<th>Engine Manifold Absolute Pressure</th>
<th>Engine Intake Air Temperature</th>
<th>Engine Speed</th>
<th>Engine Oil Temp</th>
</tr>
</thead>
<tbody>
<tr>
<td>1a</td>
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<td>20°C</td>
<td>1200 rpm</td>
<td>80%</td>
<td>40%</td>
<td>19°C</td>
<td></td>
<td></td>
<td></td>
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<td>8°C</td>
<td>20°C</td>
<td>1200 rpm</td>
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<td>40%</td>
<td>19°C</td>
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<td></td>
<td></td>
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<td>40°C</td>
</tr>
<tr>
<td>1c</td>
<td>8°C</td>
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<td>1200 rpm</td>
<td>80%</td>
<td>40%</td>
<td>19°C</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>40°C</td>
</tr>
<tr>
<td>1d</td>
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<td>20°C</td>
<td>1200 rpm</td>
<td>80%</td>
<td>40%</td>
<td>19°C</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td>1e</td>
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<td>80%</td>
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<td>19°C</td>
<td></td>
<td></td>
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<td>40°C</td>
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<tr>
<td>1h</td>
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<td>20°C</td>
<td>1200 rpm</td>
<td>80%</td>
<td>40%</td>
<td>19°C</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>40°C</td>
</tr>
</tbody>
</table>

Most of the data categories are presented in Chapter 4. The rest of the data tables are presented below:
Manifold Absolute Pressure is shown below in Figure D–1.

Figure D–1: Manifold Absolute Pressure, kPa
Barometric Absolute Pressure is shown below in Figure D–2.

Figure D–2: Barometric Absolute Pressure, kPa
Air Intake Temperature is shown below in Figure D–3.

Figure D–3: Air Intake Temperature, °C
Engine Speed is shown below in *Figure D–4*.

*Figure D–4*: Engine Speed, revolutions/minute
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Elana M. Chapman
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ABSTRACT

The use of Dimethyl Ether (DME) blended with diesel fuel was investigated as an alternative fuel for diesel engines. A pressurized fuel delivery system was designed and built. A Navistar T444E, turbocharged direct injection diesel engine was modified to accommodate the pressurized fuel delivery system and enable operation with a blend of diesel fuel and DME. The engine was operated on a diesel fuel-DME blend containing 2% oxygen by mass and cylinder pressure traces were compared with baseline operation with diesel fuel. An eight mode, AVL steady state dynamometer test was adhered to for this comparison.

Miscibility of diesel fuel and DME was examined in various blend ratios. A methodology was developed to utilize a high pressure capillary viscometer to measure the viscosity of pure DME and blends of DME and diesel fuel in varying proportions and at pressures up to 2500 psig.

The comparison of an analysis of the cylinder pressure traces did not indicate any differences in the combustion characteristics of diesel fuel and a blend of diesel fuel and DME with 2% oxygen content. The engine could not be operated at the highest power mode in the AVL eight mode test. Boiling of DME inside the injectors was believed to be the reason. Diesel fuel and DME were found to be completely miscible in all blend ratios. The viscosity-pressure relationship of DME and blends was found to be logarithmic in nature.
Dimethyl Ether (DME) is gaining increasing importance as an alternative fuel for Diesel engines. Properties of DME such as its high cetane number and undetectable smoke emissions have made it an excellent fuel for use in compression ignition (CI) engines. Moreover, due to the presence of an oxygen atom, the absence of a carbon-carbon bond in the DME molecule and its high Cetane number, the oxides of nitrogen (NOx) - particulates tradeoff that plagues the traditional CI engine running on diesel fuel, no longer seems to threaten the CI engine operated on DME. The high Cetane number of DME also helps in reducing the combustion noise, which is another drawback of the diesel fuel fired CI engine [3, 8, 20, 25, 26, 28, 29].

The work done for this thesis was the first part of a project, whose final goal is to modify a shuttle bus that runs on the Pennsylvania State University campus at University Park, PA, to operate on an optimized blend of DIESEL-DME. The first part of the project was to modify the fuel system of the engine to be able to use DIESEL-DME blends in a laboratory environment, under controlled testing conditions. A fuel delivery system was designed and developed to take into account the higher vapor pressure of DME along with its lower lubricity and viscosity. The objective here was to do as few modifications as possible and still be able to use
DME to operate the engine. The lack of lubricity of DME is a major hurdle in using the existing diesel injection equipment, which utilizes the lubricity of diesel fuel to lubricate the parts machined with close tolerances. With this in mind, it was decided to blend DME and diesel fuel to provide lubricity to the blend.

A study was performed to measure the viscosity of DME and various blend ratios of Diesel-DME at pressures up to 2500 psig. A high pressure capillary viscometer was used for this purpose. This viscometer was designed and developed at the Pennsylvania State University [1]. Modifications were made to the viscometer taking into consideration the physical and chemical characteristics of DME and the need to charge the viscometer under pressure.

1.1 What is Dimethyl Ether (DME)?

DME is the simplest ether compound (chemical formula C₂H₆O). Some physical and chemical characteristics of DME are given in Table 1-1. At standard temperature and pressure it is a gas, but can be liquefied under a moderate pressure. This makes DME quite similar to propane and liquefied petroleum gas for handling purposes. DME was first used as an aerosol propellant because of its environmentally benign characteristics. It is not harmful to the ozone layer as the CFCs that it replaced. DME is also easily degraded in the troposphere [4]. A technical information bulletin [2] gives a good overview of the physical and chemical properties of DME.
### Table 1-1: Physical and Chemical Properties of DME [2]

<table>
<thead>
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<tr>
<td>Molecular weight</td>
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</tr>
<tr>
<td>Oxygen content by mass</td>
<td>34.8 %</td>
</tr>
<tr>
<td>CAS Registry number</td>
<td>115-10-6</td>
</tr>
<tr>
<td>Boiling point @ 1 atmosphere</td>
<td>-24.825 °C</td>
</tr>
<tr>
<td>Critical temperature</td>
<td>126.85 °C</td>
</tr>
<tr>
<td>Critical pressure</td>
<td>5370 kPa</td>
</tr>
<tr>
<td>Liquid density @ 25 °C</td>
<td>656.62 kg/ m³</td>
</tr>
<tr>
<td>Vapor pressure @ 20 °C</td>
<td>516.76 kPa</td>
</tr>
<tr>
<td>Flammability limits in air by volume %</td>
<td>3.4 - 18</td>
</tr>
</tbody>
</table>

### 1.2 DME in Diesel Engines

A high Cetane number makes DME an attractive fuel for compression ignition (CI) engines. In CI engines running on a single fuel, the fuel has to be injected at the precise moment just before the firing TDC. This point is called the start of injection (SOI), which varies depending on factors like the load on the engine, speed and ambient conditions. Most modern engines usually have the ability to vary the SOI depending on the above parameters. To be able to meet the above injection requirements and also to be able to end the injection in a certain amount of crank angle degrees, injection of liquid fuel is a necessity. As DME has a high vapor pressure, the fuel system needs to be pressurized to maintain DME in a liquid state. Moreover, the fuel system has to be modified considering the physical and chemical properties of DME.
Another important consideration in the use of DME is that of the physical properties like viscosity and compressibility. Diesel fuel injection systems are designed for diesel fuel, which has a kinematic viscosity an order of magnitude higher than that of pure DME.

Taking the above properties of DME into consideration a modified fuel delivery system was designed and built. A Turbocharged Direct Injection Diesel engine was operated with this modified fuel delivery system on a DIESEL-DME blend. The engine used was a Navistar T444E diesel engine. Detail specifications of this engine can be found in Table 3-1. This engine was operated at steady state on an eddy current dynamometer. An AVL 8-Mode test schedule was followed to compare the performance of the engine when fired by diesel fuel and DIESEL-DME blend.
Chapter 2

LITERATURE REVIEW

For the past few years, many investigators have studied the use of DME in diesel engines. As with every other alternative fuel, all aspects of using this fuel have been studied. Alternatives to the fuels presently used, are often considered for a couple of important reasons:

- To make use of a previously unutilized energy source (as in natural gas) in transportation systems
- To face the ever-increasing challenge of emissions regulations.

DME appears to satisfy both the above criteria. With the present technology, natural gas in remote locations can be converted to DME, enabling economical transportation to markets and consumers; thus better utilizing the remote natural gas. The clean burning characteristics of DME, i.e., extremely low particulate emissions and low NOx with injection optimization, also help in meeting the emissions regulations.

When studying the viability of any alternative fuel, the mass production process of that fuel is always compared to that of fuels currently being used. The well to wheel efficiency, which is an important aspect in the long term success of any fuel, goes hand in hand with the production process. Combustion characteristics of
the alternative fuel are another aspect subject to investigation. More often that not, in this age, the alternative fuel is cleaner burning than the standard fuels. Finally, the compatibility of the fuel with the materials used in the fuel injection system is under scrutiny. These aspects of DME studies are examined in the following discussion of literature.

2.1 Production Process of DME and its Efficiency

The use of DME as a fuel for the compression ignition (CI) engine started in 1995 with the work done by Sorenson and Mikkelsen [3]. Prior to this, DME was used as a fuel additive to improve the ignition quality of methanol in CI engines. It was also used on a larger scale as an aerosol propellant to replace CFCs as it does not harm the ozone layer.

Hansen et al [4] reported a number of 150,000 metric tons per day as the production capacity of DME. It was solely manufactured from the dehydration of methanol. This paper presents the laboratory and pilot test carried out for large scale manufacture of DME. A single step production process is discussed which will directly convert synthesis gas into DME. This being simpler than the dehydration of methanol, is a cheaper process of DME production. To achieve a production capacity close to what will be required for an alternative for diesel fuel, different types of DME plant setups are discussed. A slow transition from methanol dehydration to stand-alone DME plants is preferred. Direct dehydration of methanol, revamping
existing methanol plants to co-produce DME and stand-alone DME plants for large scale DME production are the methods discussed and compared for their investment and convenience aspects. Direct dehydration is cited as the most convenient route for the introduction of DME as a fuel. For large scale production, however, stand-alone DME plants are most economical.

Verbeek and Van der Weide [5] have carried out an overall study of various aspects of DME as an alternative fuel. An important aspect in the use of any alternative fuel, namely its energy supply and energy security, is discussed in connection with DME. As DME can be produced by synthesis gas conversion, almost any source capable of producing synthesis gas can be used to produce DME. A reason, more important than any other, is discussed for encouraging research and development of DME as an automotive fuel. That is the long term energy security achieved by making use of a variety of energy sources such as remotely located natural gas, coal and biomass. These energy sources combined together are likely to outlast crude oil. Well to wheel efficiencies for various standard and alternative fuels are compared. In a group consisting of diesel, DME, LPG, CNG, LNG, gasoline and methanol, surprisingly enough, DME comes second after diesel in the well to wheel efficiency. This projected efficiency is more than that of gasoline despite the fact that the production efficiency for DME is the second lowest in the group. For the well to wheel CO₂ emissions, the study shows that DME again comes in second after diesel fuel. Interestingly, if the source of DME is renewable, this cuts down the net CO₂ emissions.
Ofner et al. [6] present a good overview of the use of DME in automotive engines. With alternative fuel research being quite common in the past few decades, they emphasize the importance of a global point of view when studying alternative fuels. Besides the well to wheel efficiency, another concept of the “Sustainable Process Index” (SPI) is used in evaluating the environmental impact of DME. SPI compares the mass and energy flow due to a technical process to the natural flows or cycles in the environment. The SPI of various fuels, alternative and traditional, are calculated. The SPI of DME from renewable sources is shown to be the lowest in the group. Simply put, the SPI can be said to be the effect the lifecycle of a particular fuel has on the environment. A low SPI for DME means that DME is the least taxing to the environment amongst the group of fuels discussed.

Phillips et al. [7] give a Canadian perspective of DME as an alternative fuel. The vast natural gas resources in Canada play an important role in the viability of DME as a fuel. The authors have shown that the end user cost for DME per unit volume could be comparable to diesel and lower than gasoline. However, as the energy density of DME is lower that diesel, more DME (in volume and in mass) will be required to complete a specific journey. In the end DME would still end up about 30% more expensive than diesel.
2.2 DME Injection Systems and Injection Characteristics

The earliest work done on pure DME injection were studies in which unmodified diesel injection equipment was used for DME injection with an objective of identifying the areas where modifications would be necessary to successfully use DME and then to optimize that equipment for further improving the injection process.

Kapus and Ofner [8] talk about modifying and optimizing an existing diesel fuel injection system to account for the properties of DME. This is a study of DME injection using standard diesel injection equipment. Physical property data for DME, such as compressibility and velocity of sound were estimated using modeling techniques. A test rig was configured, which measured the pressure oscillations in a tube placed between two injection nozzles. The properties like viscosity and velocity of sound were estimated using these pressure traces. A flow visualization study was also carried out using a glass nozzle. Initiation of cavitation and its effect on the spray development was studied. Different characteristic parameters of the fuel injection equipment (FIE) were varied and the effects of such changes on combustion noise, fuel consumption and NOx emissions were recorded. FIE used on DI diesel engines, was found to be generally more suitable for DME injection than FIE on IDI engines.

McCandless and Li [9] present a “Novel Fuel Injection System” (NFIS) for DME and other alternative fuels. The NFIS was designed to retrofit existing diesel
engines. Low cost was also a major consideration. The design process for a NFIS was initiated by taking into consideration the technical and economic requirements of the customer. Various existing FIE were compared to find the most suitable for modifications. A common rail system was selected as the most suitable. The low viscosity of DME caused a lot of leakage past the solenoid valve and the nozzle. A clever design of equipment, however, enabled the containment of the leakage. The NFIS was also capable of rate shaping, which is a very important control strategy for lower NOx and combustion noise. The pump used to pressurize the DME to injection pressures was an axial piston pump with special shaft seals and wear resistant pumping elements. The fuel injector used, was a modification of a commercially available injector. The nozzle spring and the inclusion of a large volume behind the plunger were the modifications made. Flow bench tests were carried out which showed that the goals such as controllable initial injection rate and system durability were met.

McCandless et al. [10] describe the design and performance of a 275 bar DME pump with variable displacement and variable output pressure. The leakage of DME past the pumping element, which is very common given the extremely low viscosity of DME, was reduced by introducing a “hydraulically long” flow path for the leakage flow. This pump was found to be overall satisfactory in terms of performance and durability. Additionally, it can also be used for other alternative fuels such as methanol, and liquid propane.
Ofner et al. [11] present an injection concept for DME injection. They have included a low pressure “purge” circuit. This purge circuit absorbs all leakages and the DME released from purging the injectors. This system includes a compressor, which then pushes the DME vapors back to the fuel tank. Since this is one of the earlier works on DME injection, a greater emphasis was given on the safety aspect and seals. Contact seals were used. There were two sealing elements in series with the space in between connected to a purge tank to collect any leakage.

Sorenson and Nielsen [12] performed an important study of the wear of plunger type diesel injection pumps pumping DME. A test rig was made out of standard diesel injection equipment, as ASTM standard lubricity test rigs are not suitable for compressed liquids. Several lubricity additives were tested. The tests were run for typically 80 hours. The pumping elements showed wear even after a relatively short time of 80 hours, showing the difficulty in using DME in conventional diesel injection systems. The location and severity of the wear, however, could not be characterized. The drawback of this test was that the areas selected for wear analysis samples were not representative of maximum wear. Also most of the wear observed was the polishing of the plunger due to metal to metal contact. Even though the wear could not be characterized, it was important that DME with all the additives showed some amount of wear. It proved that none of the additives used were effective. Also, no wear whatsoever was observed when the injection equipment was run with standard diesel fuel.
Another important aspect in DME fuel injection is the behavior of DME as it leaves the injection nozzle and enters the high temperature and pressure environment in the combustion chamber towards the end of the compression stroke. The critical point of DME at 127 °C is well below the conditions in the combustion chamber at the instance of injection. This fact makes DME behave differently than diesel fuel once injected into the cylinder.

Glensvig et al. [13] performed a comparative study of the injection properties of diesel fuel and DME. Here, the two fuels were injected into a transparent chamber filled with Nitrogen to a desired pressure and ambient temperature. The penetration rate and the spray angle were the objects of study. As expected, the diesel fuel spray was found to have a faster penetration than DME. The correlations developed for diesel fuel sprays however, gave incorrect results for DME sprays. The spray angle for DME was also found to be almost twice as wide as that for diesel fuel at the same distance from the nozzle tip. The spray angle correlation again was found to give incorrect results in predicting the difference for the two fuels.

Upto this point, all the studies done with Diesel fuel-DME blends have been with premixed solutions of the two fuels. A different approach would be to introduce DME as a liquid or a gas at the very last moment before injection. Recent studies by Sovani et al [14, 15] regarding effervescent Diesel injectors present an idea that may be well suited for injection of Diesel fuel - DME blends. Here, a study of the fuel spray was done by introducing an atomizing gas into the fuel stream before it is discharged from the injectors. It was found that the droplet size of the fuel spray
could be reduced to about 5µm at injection pressures that were about 5 to 7 times smaller than those used in modern diesel injection systems. This concept can be incorporated for use with DME injection. The atomizing gas in this case would be DME. DME could be mixed just before the fuel enters the injectors. This setup would obviate the necessity of a complicated pressurized fuel delivery system used with a mixture of diesel - DME. This method would however limit the maximum percentage of DME in the blend.

The study of literature has shown that the fuel delivery system discussed in the next chapter is the best way to modify the fuel system of the Navistar T444E engine. This attempt to mix diesel and DME will also try to solve the lubricity issues when using pure DME in diesel engines.
Chapter 3

EXPERIMENTAL SETUP

The present study is an investigation of the use of Dimethyl Ether (DME) as a fuel in diesel engines. The study consists of design and installation of an engine testing facility including an engine, dynamometer, control system and instrumentation for exhaust emissions monitoring, as well as configuration of a high-pressure viscometer instrument for measuring the viscosity of blends of diesel-DME. Moreover, the existing fuel system on the engine was modified to enable operation at higher fuel delivery pressures required for the fuel blend.

3.1 Engine-Dynamometer Setup

The engine-dynamometer was installed in a sound reducing test cell. The engine and the dynamometer were mounted on a concrete pad with adequate vibration isolators (Mtech #X pads) between the floor of the test cell and the concrete pad. Heat exchangers were installed for cooling the engine and the dynamometer. An air handling system was also installed to maintain near ambient conditions in the test cell. Care was taken to do a proper alignment of the engine and the dynamometer. The engine flywheel and the dynamometer hub were held parallel to
each other. The driveshaft was kept at an angle of 1.5 degrees. This was in accordance with a driveshaft installation guide. The driveshaft has a rubber damper to dampen driveline vibrations.

3.1.1 Engine

The engine used for the study was the Navistar T444E, 7.3 liter, 4 stroke cycle turbocharged direct injection diesel engine. The selection of this particular engine was based on several important factors. The bus, which was to be modified later, had the same engine model, with the same Engine Control Module (ECM). Secondly, this engine was used in previous studies involving DME. The reason for the popularity of this engine is its fuel injection system. This engine has the common rail Hydraulic Electronic Unit Injector (HEUI) fuel injection system. This system being electronically controlled is highly flexible in operation and hence suitable for developmental work. The relevant specifications for the T444E engine are shown in Table 3-1.
Table 3-1: T444E Engine Specifications [16]

<table>
<thead>
<tr>
<th>Engine Type</th>
<th>Diesel, 4 stroke cycle</th>
</tr>
</thead>
<tbody>
<tr>
<td>Configuration</td>
<td>Over Head Valve V – 8</td>
</tr>
<tr>
<td>Displacement</td>
<td>444 cu.in.</td>
</tr>
<tr>
<td>Bore and Stroke</td>
<td>4.11*4.18 in.</td>
</tr>
<tr>
<td>Compression ratio</td>
<td>17.5:1</td>
</tr>
<tr>
<td>Air Aspiration</td>
<td>Turbocharged and air-to-air intercooled</td>
</tr>
<tr>
<td>Rated power</td>
<td>190 bhp @ 2300 rpm</td>
</tr>
<tr>
<td>Governed speed</td>
<td>2500 rpm</td>
</tr>
<tr>
<td>Peak torque</td>
<td>485 lb-ft @ 1500 rpm</td>
</tr>
<tr>
<td>Combustion system</td>
<td>Direct injection</td>
</tr>
<tr>
<td>Fuel Injection System</td>
<td>HEUI; Common rail</td>
</tr>
</tbody>
</table>

Hower et al. [17] give a detailed description of the engine development process and describe the various important aspects such as the combustion system, fuel injection system and the electronic engine management system for the Navistar T444E.

The engine has two valves per cylinder. The fuel injection system is a state of the art, electronically controlled system that was developed by Caterpillar Inc. Stockner et al. [18] and Glassey et al. [19] describe the HEUI fuel injection system in great detail. Increasingly stringent emissions regulations were the impetus behind the development of this completely new concept in diesel fuel injection systems. This system uses engine oil to activate the injectors. The injection pressure, that can reach as high as 18000 psig (1240 bar), is achieved by using a pressure intensifier. This pressure intensifier is integral to the injector, resulting in the overall compactness of
the system. The electronic control with feedback gives a precise control over the air/fuel ratio, which is very important in controlling exhaust emissions. Another important aspect is the flexible injection timing. Start of injection is critical in minimizing emissions, as well as, fuel consumption. Due to the hydraulic-electronic activation, the injection timing can be tailored for each engine operating condition. Injection timing is decided by the electrical actuation signal that is sent to the injector solenoid. Another concept of “split shot” injection, which is a part of the HEUI fuel injection system, has been shown to reduce combustion noise as well as NOx emissions during idling and low loads.

3.1.2 Dynamometer

The dynamometer that was used was an eddy current absorption dynamometer (Dynamatic Corporation Model #AD-8102). The torque arm is 15.375 inches. The rated capacity is 450 HP between shaft speeds of 3100 to 6000 rpm. The excitation is achieved by a 45V, 14A DC power supply. The dynamometer is water-cooled and requires a cooling water supply of 45 GPM at or below 32 °C. A Lebow tension and compression load cell (model # 3132-500), with a capacity of 500 lbs, measures the load. This load cell was calibrated using free weights as described in the dynamometer operations manual.
3.1.3 Cooling Systems

Three separate cooling systems were used in the test cell. Water to water heat exchangers were utilized for engine and dynamometer cooling. The cooling water source is the same for the engine and dynamometer heat exchangers. The sequence of the heat exchangers in the cooling water line was critical, as the output temperature requirements were very different in both cases. The dynamometer requires coolant at a temperature less than or equal to 32 °C. There is no such requirement for the engine. Thus the dynamometer heat exchanger had to be put upstream of the engine heat exchanger to provide maximum cooling. An air handling system was used to constantly replace the hot air in the test cell with cooler ambient air. A schematic of the cooling systems is shown in Figure 3-1.
Figure 3-1 Cooling Systems
3.1.3.1 Engine Cooling

A water-to-water shell-tube type heat exchanger was used for engine cooling (Young Radiator Company model # F-504-EY-2P). The engine coolant, a 50-50 mixture of ethylene glycol and distilled water, flows through the shell side while treated cooling water flows through the tubes. The engine coolant temperature is maintained at a preset level in the engine block and head by a thermostat valve. There is no external temperature control for engine cooling.

3.1.3.2 Dynamometer Cooling

The dynamometer is water-cooled. A water to water heat exchanger (Affinity Industries Inc., model # 3100) was also used in this case. This heat exchanger has a modulating valve on the cooling water side which is used to control the output temperature. This valve, however, had to be disabled in order to provide the engine heat exchanger with a constant flow. Constant flow was also the requirement of the source of the cooling water.

3.1.3.3 Test Cell Cooling System

An induced draft fan arrangement was used to keep a constant airflow through the test cell. This fan has a Cutler-Hammer adjustable frequency drive to
adjust the fan speed. Constant airflow was important to keep the ambience in the test cell at a steady state. The air is introduced into the test cell from the front end. This also helps maintain an adequate airflow over the charge air cooler, which has a bank of fans to drive air through it. Maintaining the test cell temperature was also important for the operation of exhaust analysis equipment. The Micro-dilution tunnel in particular is very temperature sensitive. The temperature of the diluted exhaust gases should be at or below 52 °C. The test cell air handling system also helps in maintaining this temperature.

3.1.4 Engine flow systems: intake air, fuel

Source of intake air and fuel to the engine are important. The conditions of air and fuel such as the temperatures, humidity and presence of impurities can have a profound effect on the repeatability of engine performance. Close monitoring and control of these variables would be ideal. No control was used on the conditions of air and fuel, but they were monitored and recorded.

3.1.4.1 Intake Air System

Clean filtered air was provided to the engine using standard automotive filters. The temperature of the intake air was close to a temperature of 20 °C, as room air outside the test cell was used for this purpose. Large diameter ducting was used to ensure adequate flow capacity for the engine size. The engine being turbocharged
and intercooled, an air to air intercooler was used to cool the charge air. Cooling air was blown over the charge air cooler using a bank of fans, which in turn was aiding by the air draft from the test cell air handling system. The intake air consumption was measured using an electronic mass air flow meter. This meter is based on the hot wire anemometer principle. The meter is placed between the air filter and a plenum chamber, which dampens out any pulsation in the air flow. This is an important consideration in getting an accurate reading from the mass flow meter. The mass flow meter was calibrated using a laminar flow element. The output from the flow meter is a frequency signal, which was then converted to a voltage signal for ease of measurement. This signal was logged using a computer to give a real time air consumption reading.

3.1.4.2 Fuel Supply System

The fuel tank supplying fuel to the engine is a modified LPG tank that sits outside the test cell. Stainless steel tubes are used to supply fuel from the fuel tank to the engine and return the excess fuel to the tank. High pressure fittings were used with the future application of a pressurized fuel system in mind. The standard fuel filter provided with the engine was used for the baseline runs. For the DME test run, however a modified liquid propane filter was used to handle the high system pressure. In all cases, fuel consumption was measured by using a weighing scale.
with a readability of 2 grams. The time was measured accurately by a stopwatch to a hundredth of a second.

3.1.5 Pressurized Fuel Delivery System for Diesel-DME Blend

DME is a liquefied gas. At standard temperature and pressure, it is a gas, but liquefies under a moderate pressure. The vapor pressure and density changes with temperature are characterized in the Technical Information (ATB-25) bulletin. The fuel delivery system was designed keeping in mind the following important points:

• The vapor pressure of DME and related temperature requirements.
• Material compatibility of the various components in the fuel system with DME
• Lack of lubricity of pure DME.

A schematic of the fuel delivery system is shown in Figure 3-3. The working of the fuel delivery system can be explained as follows:

1. The fuel is delivered from the fuel tank at a pressure of about 90 psig. This overpressure is necessary to keep the DME in a liquid state. Any inert gas is suitable for this purpose. Helium was used as it has a lower solubility in DME than nitrogen.

2. The pressure is then boosted by a gear pump to about 150 psig, depending on the pressure rating of the fuel rail. The rail pressure is maintained at 70 psig in the original fuel system of the engine.
3. The fuel return line pressure is held at about 150 psig by the backpressure regulator. The regulator is a simple spring loaded valve that regulates the flow to keep the backpressure at the desired pressure.

4. This fuel then passes through a heat exchanger, where it is cooled down.

5. After cooling, the fuel is then fed to the inlet of the pump.

![Figure 3-2: Vapor pressure curve for DME (Dymel A) [2]](image)

**3.1.5.1 Fuel Pressure and Flow Requirements**

At 20 °C the vapor pressure of DME is about 62 psig. Keeping DME in a liquefied state calls for pressurizing the entire fuel system from the fuel tank up to
the fuel injectors. The vapor pressure also changes rapidly with temperature. The vapor pressure curve for DME is shown in Figure 32. The pressure of the fuel system is hence dictated by the fuel temperature. The upper limit of pressure, however, is decided by the pressure rating of the fuel rail. The engine used in the study has a common rail injection system. Each cylinder head has a fuel rail running along its length, which is the source of fuel for the pressure intensifier in the fuel injectors. In the original fuel system of the engine, the pressure in the rail is maintained at 70 psig. This facilitates proper filling of the pressure intensifier. The fuel rails in the cylinder head form a dead head system. This means that there is no fuel return once the fuel enters the fuel rail. It is because of this that the fuel temperature in the rail approaches the engine coolant temperature in the head. This layout of the fuel system was modified to accommodate a fuel return from the cylinder heads. A study was performed in which the temperature of the fuel in the fuel rail was recorded in conjunction with the fuel consumption of the engine, for the 8 modes of the AVL test. The temperature of the fuel in the fuel rail, which was about 80 °C, would be too high for the DME to remain in the liquid state at the operational pressure of 150 psig. A minimum flow rate value was calculated so as to keep the temperature of the fuel in the rail below 50 °C. The vapor pressure of DME at this temperature is about 150psi. This pressure, more or less dictated the final fuel temperature as it exited the fuel rail. The fuel delivery pump was sized based on the above calculations. In addition to excess flow rate, cooling of the returned fuel was resorted to, to maintain the required fuel temperature.
Figure 3-3: Pressurized Fuel Delivery System
3.1.5.2 Design Considerations for Fuel System

DME is known to be incompatible with the common gasket materials such as Viton and buna-N, used in diesel service. Data provided by DuPont Inc. indicated Kalrez to be the best material for DME in terms of compatibility. For economic considerations however, this material was used sparingly. Other materials such as butyl rubber, Teflon and neoprene have also been found to be compatible, though not to the same degree as Kalrez. Stainless steel was used for the fuel lines as a safeguard against corrosion. All the other components such as valves and regulators were also made of stainless steel.

Selecting a pump for pumping DME was challenging due to the properties of DME such as its low lubricity and low viscosity. Due to the vapor pressure of DME, the pump housing is required to handle pressures up to 250 psig. Positive displacement pumps such as vane pumps, diaphragm pumps and gear pumps were considered. Gear pumps were found to be economical as well as convenient to operate. With these considerations, a gear pump (Tuthill Pump CO, California model #TXS2.6PPT3WN00000) was selected. This pump has a magnetically coupled AC motor. This configuration does not have the driveshaft going through the pump housing, which in turn, obviates the need for seals. The gear material is Ryton (Polyphenylene sulphide), which was found to be compatible with diesel and
DME as per the data by provided DuPont Inc. The pump body seals are made of Teflon.

The fuel filters on the engine could not be used because of the high pressure of the fuel. The minimum pressure in the fuel lines was 90 psig. This required the use of special filters, which could withstand higher pressure. A diesel water separator was used as a primary filter. This is rated at 100 psig. The final filter was a LPG filter rated at 500psig. The mesh size of the filter was matched to the engine specifications of 3 micron.

The fuel tank was made out of a modified 100 lb capacity LPG cylinder which was pressure tested prior to use. This tank was fitted with a 1/2” NPT fitting at the bottom for liquid exchange.

3.1.5.3 Engine Operation Procedure With the Pressurized Fuel Delivery System.

The pressurized fuel delivery system made it essential for a particular operating procedure to be followed. The pressurized fuel delivery system was controlled using separate controls from the engine. A particular procedure was followed while charging the fuel tank with diesel and DME to guarantee a reasonable accuracy in mixture preparation. Studies showed that diesel and DME are miscible in any proportion. Taking this fact into consideration, no mixing apparatus was used. The tank filling procedure is as follows:

- The fuel tank is purged with Helium to get rid of any air present in it.
• The tank is placed on a weighing scale and the initial reading is noted.

• A predetermined quantity of diesel is introduced in the tank under gravity.

• Care is taken to note the reading before and after filling, with the hoses on or off, but always the same way.

• The fuel tank is then pressurized with helium to about 50 psig or to about 10 psig less that the DME cylinder pressure. This pressure differential is sufficient to force DME with a sufficient high velocity into the fuel tank to aid the mixing of diesel and DME.

• The quantity of DME is again predetermined by the ratio of diesel to DME required.

• Additional helium is now introduced in the fuel tank to raise the pressure to about 90 psig.

• In addition to the entering velocity of DME, the blend is allowed to stand for about 6 hours, by which time the blend is assumed to be completely homogeneous.

• The fuel pump in this case is electric driven. For this reason it is turned ON and the backpressure regulator is set to 150 psig before the engine is fired.

• During engine operation, the temperature of the fuel exiting the rail needs to be monitored to prevent it from going above a certain limit.
3.1.6 Data Logging and Test Cell Control

One of the main purposes of an engine test cell is to produce data for various experimental settings. Real time data acquisition is very important to ascertain the steadiness of the engine operating parameters during a steady state test. This also helps in making a statistical interpretation of the data to get an estimate of the variation of the parameters during testing. Control systems on the other hand, help in maintaining desired conditions in the test and also act as safety checks for various aspects in test cell operation. Together these two systems help in the day to day activity of a test cell. A Programmable Logic Controller made by Modicon was used for data acquisition as well as test cell control. In addition to this a data acquisition card by Kiethley Instruments was also used.

3.1.6.1 Data Acquisition System

A PC based system was used for acquisition and logging of the various data output from the engine and the dynamometer. Type K thermocouples were used to record various temperatures. Table 3-2 shows the measurement equipment used for each type of measurement done.
Table 3-2: Data Acquisition

<table>
<thead>
<tr>
<th>Data type</th>
<th>Sensing instrument</th>
<th>Recording instrument</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature</td>
<td>Type K thermocouples</td>
<td>PC based Modicon PLC</td>
</tr>
<tr>
<td>Cylinder pressure</td>
<td>Kistler pressure transducer (6125A non-cooled)</td>
<td>PC based DAS 1800 data acquisition card</td>
</tr>
<tr>
<td>Engine control module data</td>
<td>Engine sensors</td>
<td>PC based National Instruments CAN card</td>
</tr>
<tr>
<td>Engine speed</td>
<td>Hall effect sensor.</td>
<td>PC based Modicon PLC</td>
</tr>
<tr>
<td>Engine load</td>
<td>Lebow load cell (model # 3132-500)</td>
<td>PC based Modicon PLC</td>
</tr>
<tr>
<td>Air Flow</td>
<td>Mass Air Flow</td>
<td>PC based DAS 1800 data acquisition card</td>
</tr>
<tr>
<td>Fuel Flow</td>
<td>Electronic Mass Scale</td>
<td>Manual observation</td>
</tr>
</tbody>
</table>

3.1.6.2 Test Cell Control

The engine and dynamometer operation was controlled using a Programmable Logic System (Modicon). The PLC had some shut off limits for various temperatures and engine speed. The OEM Accelerator Position Sensor (APS) was used to control engine fueling. No feedback control was deemed necessary for
engine speed as the dynamometer and OEM APS were able to keep the engine speed to within a few rpm of the desired value. The air handling and cooling systems and the pressurized fuel delivery system had separate controls of their own.

3.1.7 Engine Testing and Exhaust Emissions sampling strategy

One of the main objectives of this project was to demonstrate the operation of the T444E engine on a blend of diesel and DME. Due to the nature of combustion in a diesel engine, particulates and NOx are the main pollutants formed in the exhaust. CO, CO\textsubscript{2} and hydrocarbon emissions are usually less than its counterpart, the spark ignited engine due to leaner air fuel mixtures and high thermal efficiency.

3.1.7.1 Engine Testing Procedure.

A steady state dynamometer testing strategy was adopted to examine the engine performance and emissions with different fuels. The AVL 8 mode steady state test was followed as it closely approximates the US-FTP heavy duty engine testing procedure. This test was also used in previous studies involving the T444E engine and DME [20]. The AVL 8 mode test is shown in the Table 3-3
The AVL 8 mode test dictates the speed of each mode in relation to the idle and rated speed. The load is fraction of the maximum brake torque at that speed.

The speed and load for the 8 modes can be calculated as follows:

*Speed (rpm) = Low idle + %speed * [ (rated - low idle) / 100 ]

where,

low idle = 700 rpm

rated speed = 2300 rpm

**Load % = is the percentage at each speed as per the torque curve.

Table 3-3: AVL 8 Mode Test

<table>
<thead>
<tr>
<th>Mode</th>
<th>RPM (%)*</th>
<th>Actual rpm</th>
<th>Load (%)**</th>
<th>Actual load (ft-lb)</th>
<th>Wt. Factor %</th>
<th>Engine HP</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0</td>
<td>700</td>
<td>0</td>
<td>0</td>
<td>35</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>11</td>
<td>876</td>
<td>25</td>
<td>84</td>
<td>6.34</td>
<td>13.97</td>
</tr>
<tr>
<td>3</td>
<td>21</td>
<td>1036</td>
<td>63</td>
<td>224</td>
<td>2.91</td>
<td>44.12</td>
</tr>
<tr>
<td>4</td>
<td>32</td>
<td>1212</td>
<td>84</td>
<td>357</td>
<td>3.34</td>
<td>82.38</td>
</tr>
<tr>
<td>5</td>
<td>100</td>
<td>2300</td>
<td>18</td>
<td>77</td>
<td>8.4</td>
<td>33.90</td>
</tr>
<tr>
<td>6</td>
<td>95</td>
<td>2220</td>
<td>40</td>
<td>178</td>
<td>10.45</td>
<td>75.24</td>
</tr>
<tr>
<td>7</td>
<td>95</td>
<td>2220</td>
<td>69</td>
<td>307</td>
<td>10.21</td>
<td>129.79</td>
</tr>
<tr>
<td>8</td>
<td>89</td>
<td>2124</td>
<td>95</td>
<td>409</td>
<td>7.34</td>
<td>165.20</td>
</tr>
</tbody>
</table>
The torque curve provided by the engine manufacturer was used to calculate the load, though it was also verified during testing. Exhaust emissions were measured and quantified at each of the 8 modes after allowing sufficient time to attain a steady state. The exhaust temperature, as well as the oil and coolant temperatures, were closely monitored to decide whether the engine has reached a steady state or not. The speed of the engine was precisely controlled with an accelerator position sensor (APS/IVS) switch. This, along with a precise load control, helped in maintaining the steady states repeatable and close to that required by the modes.

3.1.7.2 Exhaust Sampling Strategy.

As discussed earlier, the exhaust was collected and characterized for particulates, NOx, CO, CO₂ and hydrocarbons. The same sampling and characterization procedure was followed to ensure consistency for all the fuels used.

3.1.7.2.1 Sampling of Particulate Emissions.

The BG1 microdilution test stand was used to measure the particulate emissions. A detailed description of the instrument can be found in the operations manual [21]. The raw exhaust passes through a micro dilution tunnel, where it is mixed with shop air in a predetermined dilution ratio. This diluted exhaust then passes through a Pallflex membrane filter (Emfab™). The filters are 90mm in diameter. The particulates are collected on the filter as the gas passes through it.
These filters are preconditioned in a humidity controlled environment for 24 hours before and after collection of particulates. The temperature of the gas stream and the filter is kept at or below 52°C. The selection of operational parameters such as the dilution ratio, the total flowrate and the sampling time was done based on a trial and error method. These conditions were adjusted so as to collect sufficient sample during a sampling duration of 5 minutes. With the sampling time fixed, the dilution ratio and total flow were adjusted. It was found that as the total flow increased above 150 SLPM the mass of particulates collected in a particular amount of time actually decreased. This was attributed to a very high face velocity. The dilution ratio is an important parameter. Lapuerta et al [22] have shown the effect of dilution ratio and other sampling conditions such as filtering temperature and pre and post processing of the filters on the measured specific particulate emissions. The effect of filter temperature and the dilution ratio on the specific particulate emissions is discussed. It is shown from data gathered, that the dilution ratio has a more profound effect on the mass of particulates collected, than the filter collection temperature (12% reduction for a dilution ratio change from 5 to 25 as opposed to a 3.7% reduction for the filter temperature change from 25 °C to 45 °C). Keeping this in mind it was decided to keep the dilution ratio for collection constant and allow the collection temperature to vary. The range of dilution ratios in our case was bound by the amount of particulates collected within 5 minutes, on the higher side and the filter temperature on the lower side. By trial and error, 8:1 ratio was found to be suitable. A higher dilution gave an insufficient sample mass, while a lower
dilution ratio resulted in higher filter temperatures. The 8:1 dilution ratio was used throughout all the testing done i.e. for all modes of all tests with different fuels. This removed the variation due to sampling conditions from the results. As cooling of the gas sample was unavailable, the collection temperature varied from mode to mode. To account for the sample to sample variation, 5 samples were taken for every mode in every test. The filters were weighed before and after sampling, every time after equilibration. The humidity controlled chamber is Electro-tech Systems, Inc. (Model # 506A). The humidity controller was by the same manufacturer (model # 514). The humidity in the chamber was maintained at a relative humidity of 45%. The scale used for weighing the samples was a Sartorius microbalance with a resolution of 1µg.

Another instrument that was used for particulate analysis is the Series 5100 Diesel Particulate Measurement System (Rupprecht and Patashnick Co., Inc.) A detailed description of this can be found in the Operating manual. Okrent [23] also gives a description of the instrument as well as a sample application. This instrument samples the raw exhaust gas. The sampling line was heated to keep any condensation from taking place. The ceramic filter can be maintained at a specified temperature during collection. After the collection phase, comes the analysis phase. During this phase, the ceramic filter is heated from 250 °C to 750 °C. The temperature is ramped at a particular rate.
3.1.7.2 Oxides of Nitrogen, CO, CO₂ and Hydrocarbon Emissions.

A Fourier Transform Infrared analyzer was used to quantify the NOx, CO and CO₂ emissions. The one used was made by The Nicolet Instrument Corp. (model # 552), with a He-Ne laser. Moisture free exhaust gas was passed through the cell. The exhaust gas was stripped of moisture in a chiller. This was then directed to the FT-IR and an oxygen analyzer. The FT-IR measures the absorption spectrum of the exhaust sample gas. Prior to sampling, a background spectrum is taken with nitrogen as the purge gas. The FT-IR was calibrated using span NO, NO₂, CO and CO₂ gases. Various concentrations were made using mass flow controllers. The solvent gas used was nitrogen.

3.1.8 In Cylinder Pressure Measurement

A Kistler pressure transducer was used to measure the pressure in the cylinder through the engine cycle. The transducer used in this case was a noncooled type (6125 A), along with a suitable charge amplifier. A minimum of 40 consecutive traces and a maximum of about 140 traces were collected depending upon the speed of a particular mode of engine operation. These raw pressure traces were then processed using a software program, PTRAN by Optimum Power Technologies to get a pressure vs. crank angle trace for the recorded cycles. The same software was also used to get an average pressure trace of the collected cycles for each mode.
The heat release analysis was done using the generic equation as described in Heywood [24].

\[
\frac{dQ_n}{dt} = \frac{\gamma}{\gamma - 1} p \frac{dV}{dt} + \frac{1}{\gamma - 1} V \frac{dP}{dt}
\]  

(3.1)

Here;

- $Q_n$ is the net heat release
- $\gamma$ is the ratio of specific heats.
- $V$ is the cylinder volume
- $P$ is the cylinder pressure.

In actual calculations time was substituted by the crank angle.

The specific heat ratio was assumed to be 1.33 for all the cases for the 8 mode test. This analysis was done on the ‘Average Pressure Trace’ output of PTRAN. The resulting plots are compared in Chapter 4. Various combustion events like the start and end of combustion were estimated using the Heat release curves obtained as shown above.
3.2 High Pressure Viscometer Setup

To optimize the performance of a fuel injection system for a particular fuel or fuel blend, it is very important to have a good estimate of the physical properties of those fuels. It is equally important to know the change in properties with change in pressure and temperature. An experiment was set up to measure the viscosity of diesel, DME and their blends at various pressures. The high pressure viscometer apparatus used for this work was designed and built at The Pennsylvania State University in 1962-63. This equipment was modified to allow for charging of pressurized liquid samples, as is necessary when dealing with compressed liquids. Johnson [1] in his Master’s thesis, gives a detailed description of the design and use of the apparatus. The equipment, very simple in design, nevertheless is extremely accurate in viscosity measurement up to a pressure of 10000 psig.

3.2.1 Description of High Pressure Viscometer

A schematic diagram of the apparatus setup is shown in Figure 3-4. The setup consists of a pressure intensifying system, a pressure measurement system, a constant temperature bath and the viscometer pressure vessel.
Figure 3-4: High Pressure Viscometer Setup

- Pressure Measurement System
- Sample Cylinder
- Pressure Gauge
- Viscometer Pressure Vessel
- Constant Temperature Bath
- AMINCO Hydrogenation Bomb
- Compressed Helium
- Oil Reservoir
- Regulated Air Supply
- Air Driven Pressure Intensifier
- 10000 psig
- 5000 psig
3.2.1.1 Pressure Supply and Measurement Equipment

Pressurized helium produces the high pressure required for this equipment. Initial studies were carried out with nitrogen as the pressurizing gas. It was found, however, that nitrogen is highly soluble in DME. Higher the solubility of a gas in DME, higher is the error in measurement of viscosity. For this reason it was decided to use helium, which is less soluble in DME as compared to nitrogen. Pressures up to 2000 psig can be attained by simply connecting a commercial helium gas cylinder to the system. For higher pressures, the air operated pressure intensifying pump is used. This system consists of an AMINCO dehydrogenation bomb with a volumetric capacity of 4.5 liter. This bomb is filled with helium from the gas bottle at the available pressure. This is further compressed by pumping oil into the bomb. A SAE30 oil was used in this case. For higher pressures, however, a SAE50 oil is recommended. The oil is pumped by an air driven pump (Teledyne Sprague Engineering Model # S-216-J-150). The 150 in the model number, indicates the factor by which the pressure of oil can be raised as compared to the pressure of the supplied driver air.

The pressurization procedure can be explained as follows with reference to Figure 3.4:

- Valve B, the return valve is closed and valve A is opened. The outlet oil pressure gauge will read the pressure of the helium in the bomb.
• Driver air is supplied to the pump at a pressure of about 25 psig. An air lubricator is installed upstream of the pump. If the air inlet valve to the pump is open, it will cause the pumping plunger to pump oil into the hydrogenation bomb. The addition of oil increases the pressure of helium already present in the bomb. There is a one way valve in the oil supply line, which prevents the oil from flowing back into the pump. The pressure of the helium in the bomb is read by an accurate pressure gauge.

• When the desired pressure is reached, the air inlet valve to the pump can be closed to stop the pump. The valve A is then closed. High pressure after the pump is relieved slowly by gradually opening the valve B.

• The bomb is now ready to supply pressurized helium to the viscometer apparatus. The pressure of helium is always kept a bit higher than the required pressure in the viscometer.

• The amount of oil in the bomb can be estimated by observing the change in the oil level in the oil reservoir.

• Care should be taken to prime the pump with oil. This can be achieved by gravity feeding the oil.

High pressure gauges are used to measure the pressure in the system. These gauges have a maximum readability of 5000 and 10000 psig and have a rated accuracy of 0.5% of full scale. For the present study, the gauge with a capacity of 5000 psig was used for its higher resolution.
3.2.1.2 Constant Temperature Bath

For the present study, the viscometer housing was kept at a constant temperature of 100 deg F. Water was used as the bath fluid. It was maintained at 100 deg F, using a temperature controller by Omega Engineering (Model CN9000A). This controller reads the temperature by means of a K-type thermocouple. There is a feedback loop that controls the electric immersion heater. Using this system, the temperature in the bath can be maintained to within 0.1 deg F. The bath is made out of a glass jar 12 inches in diameter. The water level in the bath is maintained at a certain height so that the viscometer pressure vessel is fully immersed in water. A submersible pump is used for water circulation, which helps maintain a constant temperature throughout the water bath. The bath temperature control is typically kept ON continuously to assure a stable operating temperature.

3.2.1.3 Viscometer Operation Procedure

The operating procedure for the viscometer is described with respect to Figure 3-4. Prior to beginning the viscosity measurement, the sample for viscosity measurement is prepared in the sample cylinder and maintained at a pressure of about 500 psig. The sample cylinder is then kept in the water bath overnight to ensure thermal equilibrium between the sample and the water bath. The AMINCO hydrogenation bomb is charged to the required pressure by operating the pressurizing equipment. The viscometer capillary is assembled in the pressure vessel and is brought up to the
operating pressures. The bath is brought up to the operating temperature of 100 deg F. The following steps are then taken to charge the viscometer capillary with the fluid and measure its viscosity:

1. The 5000 psig pressure gauge is used to measure the pressure in the viscometer housing. At this point valves 1, 3, 4, 6, 9 and 12 are open and valves 2, 5, 7, 8, 10, 11, 13 and 14 are shut off.

2. The sample cylinder is connected to the viscometer housing at valves 2 and 5. Valve 2 is opened to equalize the pressure between the sample cylinder and the viscometer housing. Valve 5 is used to introduce the desired amount of sample into the viscometer housing. The level of the sample is observed through the glass windows on the viscometer housing. As soon as the desired level is reached, the valve 5 is closed. The sample cylinder can then be disconnected from the viscometer housing.

3. As the test fluid is already at the operating temperature, the viscometer capillary can be immediately filled with the sample. To accomplish this, valves 3 and 4 are shut off. Helium is then vented very slowly to the atmosphere through valves 1 and 2. Valve 2 can be left fully open and the rate of capillary filling can be controlled by modulating valve 1. It is very important to fill the capillary very slowly. Rapid filling may cause boiling and separation of the sample. A rule of thumb is to allow the same time for filling as it takes for drainage. The liquid level in the capillary should be a little above the uppermost etched line above the degasification bulb as shown
in Figure 3-5. At this point valves 1 and 2 are closed. Valves 1 and 4 are then opened slightly to equalize the pressure between the top of the capillary and the reservoir. Valve 3 is then opened so as to read the correct pressure in the viscometer housing. Valve 1 is then closed immediately to maintain the level of the liquid in the capillary. The filling process can reduce the pressure to a great extent. It is important to begin with a pressure higher than desired. With sufficient practice, the desired level in the capillary can be reached at the desired pressure.

4. At this time the viscometer is left to equilibrate for a couple of hours. This is to ensure that the entire system has reached a constant desired temperature.

5. After about two hours, the viscometer is ready for viscosity measurement. Care should be taken that the liquid level hasn’t changed significantly in the capillary. Starting drainage from different liquid levels may introduce errors as will be discussed later. Valve 1 is now opened to allow the liquid to start flowing down. The stop watch is started as soon as the lower liquid meniscus comes in line with the top etched line on the efflux bulb. The time is stopped when the lower meniscus reaches the lower etched line at the bottom of the efflux bulb. The time is measured with a resolution of 0.01 seconds. The measured efflux time along with the characteristic distance are the only two observations required of the viscosity test run. The characteristic distance is the distance between the liquid level at the bottom of the viscometer housing
Figure 3-5: Viscometer Capillary

- Upper fill line
- Degasification bulb
- Time start etch line
- Efflux bulb
- Time stop line
- Capillary
and the bottommost etched line on the capillary. This distance is measured by a cathetometer with a resolution of 0.005 cm.

3.2.1.4 Sample Preparation

The samples tested for viscosity were blends of Diesel and DME in various proportions. A schematic of the apparatus used for sample preparation is shown in Figure 3-6. It consists of a transparent pressure vessel used to hold the mixture under a pressure of about 90 psig. The pressurizing gas is Helium, which is chosen for its lower dissolution in DME. The following steps describe the preparation of a homogeneous sample and its transfer into the sample cylinder.

1. About 100g of DME are introduced into the sample cylinder. The sample cylinder is then pressurized to about 90 psig. The difference in mass of the sample cylinder with and without DME gives the exact amount of DME present in the cylinder.

2. The observation vessel is pressurized with Helium to about 90 psig. The sample cylinder filled with DME is attached in the circuit 1. The valves connecting the sample cylinder to the observation vessel are opened. Under the influence of gravity, DME flows down into the observation vessel. The circuit 1 helps in equalizing the pressures in the two connected vessels. Sufficient time is allowed for all the DME in the sample cylinder to drop down into the observation vessel. The sample cylinder is disconnected and
Figure 3-6 Diesel-DME Sample Preparation
weighed. The difference in mass is the mass of DME introduced into the observation vessel.

3. This known mass of DME dictates the mass of Diesel fuel to be put in to get the desired Diesel-DME blend ratio. The required amount of Diesel is then introduced into the observation vessel in a similar manner. The velocity of the Diesel fuel entering the observation vessel helps in forming a homogeneous mixture instantly.

4. This mixture is then introduced into the sample cylinder through circuit 2. The mixture blend ratio is calculated by taking the actual mass numbers as observed.

3.2.2 Viscometer Calibration and Data Analysis

The viscometer is a capillary type viscometer, in which a constant volume of liquid is allowed to flow under gravity into a reservoir of the same liquid. Reference lines are etched on the glass viscometer for measurement of head. The parameters measured are;

- time required for a predetermined volume of liquid to drain through the capillary with a resolution of 0.01 seconds.
- a characteristic distance, the distance between the bottommost etched line on the viscometer bulb to the liquid level in the reservoir in the viscometer housing.
The procedure followed for calibration of the viscometer is the same as that followed for viscosity measurement. The calibration is performed by allowing a liquid with a known viscosity to drain from the capillary. A series of runs are performed by varying the characteristic height. For every run, the characteristic height and drainage time are recorded. A viscometer constant is calculated as follows:

\[
\text{viscometer constant (s/ cSt)} = \frac{\text{drainage time (s)}}{\text{viscosity of calibration liquid (cSt)}}.
\]

For viscosity measurement, runs are performed within the range of characteristic height used during the calibration runs. The viscometer constant is obtained using the calibration curve and the characteristic height during viscosity measurement run. The constant together with the measured drainage time gives the viscosity of the liquid. Detailed discussion of the data analysis can be found in [1].
Chapter 4

RESULTS, DISCUSSIONS AND CONCLUSIONS

The ability to operate the Navistar T444 E engine on a fuel blend of Diesel-DME with a 2% O content was demonstrated. The pressurized fuel supply system that was designed to provide a liquid fuel blend to the fuel injectors/intensifier unit was also tested for its suitability and practicality. The viscosity of Diesel-DME blends with varying blend ratios was measured. Observations were made regarding the miscibility of diesel and DME for the blend ratios considered.

4.1 Miscibility Studies of Diesel - DME Blends

To ensure stable operation of a direct injection diesel engine, it is imperative that the fuel blend is always maintained in a homogeneous liquid state. The miscibility of the fuels is therefore an important aspect of any fuel blend.

The miscibility tests of Diesel-DME blends were carried out in a pressurized vessel with a transparent window for observation. Blend ratios were varied in steps of 25% by mass. Thus 0, 25, 50, 75 and 100 were the percentages of DME by mass in a blend comprising of DME and Diesel. Diesel has a density that is higher than that of DME. To separate the effect of gravitation on the mixing process, diesel was introduced as the lower layer in the observation vessel. DME, which was introduced
from the top as a liquid under pressure, formed the upper layer. The pressure was maintained at about 90 psig. This same pressure was later to be used in the pressurized fuel tanks for the engine. This non-homogeneous mixture was left undisturbed and the quality of the mixture was observed after various time intervals. This process ensured that the mixing would be only because of molecular diffusion and not because of any physical mixing via turbulence or density difference. It was observed that irrespective of the blend ratio, the blend became a homogeneous mixture in about 6 hours.

Under these testing conditions, minimal fluid mixing took place to assist the formation of a homogeneous mixture, relying primarily on molecular diffusion. Introduction of one fuel into the other at some velocity greatly helps the mixing process.

Separation of the two fuels is also an important issue. Once a homogeneous mixture is formed, the fuels do not separate until DME starts boiling. Boiling may occur due to a reduction in pressure or an increase in temperature. Boiling is very likely to occur in the fuel rail of a diesel engine due to elevated temperatures. During the test run with a 2% O₂ content Diesel-DME mixture, unsteady operation of the engine and a loss of power were observed. This could have been due to the boiling of the DME present in the blend. The resulting fuel blend entering the fuel injectors was a mixture of liquid and DME vapors. This is thought to be the cause of power loss and unsteady operation. The temperature of the fuel in the rail was measured at 53 °C, which is higher than the boiling point of DME at the rail pressure of 150 psig.
4.2 Viscosity of Diesel - DME Blends

The high pressure viscometer capillary was calibrated using a Cannon Certified Viscosity Standard N 1.0. Figure 4.1 shows the calibration curve obtained for the capillary used. The procedure followed to obtain the calibration was similar to the procedure for data taking. The calibration curve is typically a straight line. Mass was used as the controlled variable in determining the composition of the samples. Table 4-1 shows the samples used for viscosity measurement.

Table 4-1: Samples for Viscosity Measurement

<table>
<thead>
<tr>
<th>Sample number</th>
<th>Percent of DME by Mass</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>2</td>
<td>74</td>
</tr>
<tr>
<td>3</td>
<td>50</td>
</tr>
<tr>
<td>4</td>
<td>26</td>
</tr>
<tr>
<td>5</td>
<td>0</td>
</tr>
</tbody>
</table>

The remaining portion of the sample was made up by an Emissions Certification Diesel Fuel (ECD-LS) from Specified Fuels, a federal emissions certification fuel. Figure 4.2 shows the effect of pressure on the viscosity of the various liquid samples. The kinematic viscosity is plotted on a logarithmic scale. Johnson [1] notes that a plot
of the logarithm of the kinematic viscosity versus the pressure results in a straight line. DME and diesel fuel-DME blends also follow a logarithmic relation with pressure.

The slope of this line can be used to extrapolate to higher pressures with a fair amount of accuracy. The line representing the ECD-LS fuel starts from ambient pressure. For the remaining samples with DME as a constituent, the starting point was 500 psig to ensure that the samples remain in a liquid state. The pressure range in which the viscosity measurements are made are typical in the low pressure circuit of a diesel engine. Previous studies, done with pure DME as the fuel, state that lower injection pressures can be used for DME as the condition in the cylinder just before the firing TDC allows a very rapid vaporization of DME.

Another use of the viscosity versus pressure relations for the various blends is to choose or design a fuel injection system for the optimized blend ratio. There are two ways in which the fuel system can be designed for an engine running on a Diesel-DME blend. The first one is to examine the capability of existing fuel systems. The deciding factor in this case will be the minimum viscosity that an existing fuel injection system can handle. The other way would be to optimize a blend ratio for a particular engine considering the exhaust emissions benefits and the energy density tradeoffs and to use the viscosity data of this particular blend to design a fuel injection system for that blend ratio.

Figure 4.3 shows the response of the kinematic viscosity to the ratio of DME in the blend. This graph shows the effect of pressure on the viscosity of a particular blend.
Sivebaek et al. [30] have measured the viscosity of pure and blended DME. The viscosity was measured at the vapor pressure at that temperature. The viscosity of pure DME as measured by Sivebaek et al. is 0.185 cSt at a temperature of 25 °C. In the present study, the viscosity of DME was not measured at the vapor pressure. This value can however, be estimated by extrapolating the pure DME viscosity line to lower pressures. The viscosity of DME in Figure 4.2 shows a value of about 0.199 cSt, however, at a different temperature.
Figure 4.1: Viscometer Capillary Calibration Curve

\[ y = -23.486x + 423.3 \]

\[ R^2 = 1 \]

Distance between the bottommost etched line on the capillary and the liquid level (cm)

Viscometer constant (s/cS)
Figure 4.2: Pressure-Viscosity Relationship for Diesel-DME Samples @ 100 F
Figure 4.3: Relation of Blend ratio and Viscosity
4.3 Engine Operation With Diesel-DME Blend

The first Diesel-DME blend to be run in the engine was one with a blend ratio so as to contain 2% O by mass. This particular number was chosen for the blend to be equivalent to blends of diesel and other oxygenates in terms of the Oxygen content for comparison. The performance of the existing fuel system was studied and modifications were performed as described in the Chapter 3. Back to back AVL 8-mode tests were run with the ECD-LS fuel and the 2% O content Diesel-DME blend.

4.3.1 Modified Fuel System Performance

To maintain the DME in a liquid state, a fuel rail pressure of 150 psig was used. A necessary condition was to keep the temperature of the fuel in the rail below 50 °C. To help maintain the desired temperature in the fuel rail a chiller bath was used to cool the fuel as it exited the fuel rail. Figure 4.4 shows the measured temperature of the fuel in the fuel rail of the engine for different configurations of the fuel system. As can be seen in the unmodified fuel system, the temperature of the fuel in the rail approaches the coolant temperature of around 80 °C. This is because the fuel system on the T444E engine is a dead head system where the fuel entering the rail must be injected into the engine over the next few engine cycles. The fuel return takes place just before the fuel enters the fuel rail.
Figure 4.4: Temperature of Fuel in Rail

Mode

Temperature of Fuel (deg C)

Base_unmodified
Base_pressurized
DME
With this “dead head” setup, it is impossible to maintain the temperature of the fuel in the rail below the required 50 °C during steady state engine operation. In the modified fuel system, the fuel passing through the rail is returned allowing for cooling of the fuel in the return path. The drop in temperature seen in Figure 4.4 was due to the cooling of the fuel as it returns from the rail. It can be seen however, that the fuel temperature for modes 4 and onwards was still higher than 50 °C. This resulted in an unsteady operation of the engine for mode 8. The engine was not able to maintain the required speed of 2124 rpm at a brake torque of 409 lb-ft. As seen in Figure 4.5 the engine speed kept on falling. The unstable running condition seen at Mode 8 (the data represents approximately 2.5 minutes of running time), was caused by boiling of the DME in the transition passage from the fuel rail to the injector, where the fuel pressure is still equal to the rail pressure and the temperature higher than the fuel rail. In the stock fuel system, the fuel enters the injectors at a temperature in excess of 80 deg C. The T444E engine management system does not change injection timing to compensate for fuel temperature, which could lead to a poorer fuel spray in the cylinder if the fuel temperature is much less (in our case it was maintained below 50 deg C with the exception of Mode8).
Figure 4.5: Engine Instantaneous Speed Comparison for Mode 8

Desired Speed = 2124 RPM

Sample points
4.3.2 Incylinder Pressure Trace Analysis

Cylinder pressure for cylinder number 1 was recorded for all the modes of the AVL test for both the fuels. Effect of the different fuels on combustion events such as start of burn and combustion duration was observed. A rate of heat release (ROHR) calculation was carried out. Figure 4-6 compares the “effective” ignition delay between the two fuels for various modes. The ignition delay numbers shown are a sum of the actual ignition delay and a "hydraulic+electric" delay in the fuel injectors. The hydraulic delay is caused by a finite amount of time that passes between the electronic command signal from the ECM to the injector and the occurrence of actual fuel injection. As the injector is a HEUI, the oil temperature and oil pressure influence this hydraulic delay. The electrical delay is the time delay between the initiation of a signal by the ECM, being converted to a higher voltage by the Injector Driver Unit (essentially a capacitive discharge ignition coil), then proceeding to open the oil supply solenoid to the injectors. In the absence of accurate knowledge of the hydraulic and electric delays, it is safe to assume that the delays will be constant at a particular engine operation mode regardless of the fuel used. Monitoring of the pressure and temperature of the oil supply to the injector was performed during the two runs with diesel and 2%O blend to show similar conditions. Based on this, the ignition delay numbers were reduced as a ratio between a particular mode and the maximum delay recorded.
Figure 4-6: Ignition Delay Comparison
Figures 4-7 to 4-14 present a pressure trace comparison for all the modes of the AVL test between the two fuels. By looking at the heat release curves a comparison can be made between the diesel and blend pressure traces. The ROHR curves calculated here compare very well with the curves generated by commercial software such as PTRAN. Mode 2 seems to be the only mode where the "split-shot" injection is in evidence. The "split shot" is due to a spill port in the injector. The amount of pilot injection is the same for any operation mode of the engine. Mode 2 seems to be the only mode in which the pilot injection can really be differentiated from the rest of the injection. In general, slightly higher ROHR and peak cylinder pressures are observed for the operation with 2% O fuel blend. Mitsumasa Ikeda, et al [31], have studied the combustion of 8% DME blend (by mass) in diesel (2% O corresponds to 5.747% DME by mass). They have observed a "delayed heat release" for 8% DME blend as compared to Diesel fuel. The have also noted an increase in max ROHR with an increase in DME concentration. The present work with 5.747% DME and the above reference with 8% DME seem to indicate results opposite to expectations considering the higher Cetane number of DME. In both studies the injection timing was maintained the same for both fuels. This may be the cause of the anomaly. With a higher Cetane value, the diesel-DME blend will need an optimization of injection timing to strike a balance between too high ROHR and too high THC emissions. Thesis work by Matt Stoner [32] also suggests that there is not a significant difference in the ROHR (rather burn rate) of the different oxygenate blends with 2% O content. All oxygenates considered there, were within the boiling range of diesel
fuel. Boiling point of DME being drastically less than that of diesel fuel, an expectation was to see some effect of 2% O diesel-DME blend on the ROHR. However, this physical advantage of lower boiling point may not be realized at an Oxygen level of 2% (DME 5.747% by mass).
Combustion Event | Diesel | Diesel-DME Blend |
<table>
<thead>
<tr>
<th></th>
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<th></th>
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</thead>
<tbody>
<tr>
<td>Start of Injection (Degrees ATDC)</td>
<td>-6</td>
<td>-6</td>
</tr>
<tr>
<td>Start of Ignition (Degrees ATDC)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>End of Burn (Degrees ATDC)</td>
<td>22</td>
<td>22</td>
</tr>
<tr>
<td>Max ROHR (J/crank angle degree)</td>
<td>28.78</td>
<td>23.74</td>
</tr>
</tbody>
</table>

Figure 4-7: Pressure Trace and ROHR-Mode 1
Combustion Event | Diesel | Diesel-DME Blend
---|---|---
Start of Injection (Degrees ATDC) | -8 | -8
Start of Ignition (Degrees ATDC) | 0 | 0
End of Burn (Degrees ATDC) | 30 | 30
Max ROHR (J/ crank angle degree) | 37.70 | 40.61

Figure 4–8: Pressure Trace and ROHR-Mode 2
**Combustion Event** | **Diesel** | **Diesel-DME Blend**
--- | --- | ---
Start of Injection (Degrees ATDC) | -8 | -8
Start of Ignition (Degrees ATDC) | 1 | 1
End of Burn (Degrees ATDC) | 42 | 42
Max ROHR (J/crank angle degree) | 83.87 | 88.69

*Figure 4-9: Pressure Trace and ROHR - Mode 3*
Combustion Event | Diesel | Diesel-DME Blend |
<table>
<thead>
<tr>
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<th></th>
<th></th>
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</thead>
<tbody>
<tr>
<td>Start of Injection (Degrees ATDC)</td>
<td>-8</td>
<td>-9</td>
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<tr>
<td>Start of Ignition (Degrees ATDC)</td>
<td>1.5</td>
<td>0.5</td>
</tr>
<tr>
<td>End of Burn (Degrees ATDC)</td>
<td>45</td>
<td>45</td>
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<tr>
<td>Max ROHR (J/crank angle degree)</td>
<td>115.55</td>
<td>125.19</td>
</tr>
</tbody>
</table>

Figure 4-10: Pressure Trace and ROHR-Mode 4
Combustion Event | Diesel | Diesel-DME Blend
--- | --- | ---
Start of Injection (Degrees ATDC) | -18 | -17
Start of Ignition (Degrees ATDC) | 1.5 | 2.5
End of Burn (Degrees ATDC) | 52 | 52
Max ROHR (J/crank angle degree) | 44.53 | 46.74

Figure 4-11: Pressure Trace and ROHR-Mode 5
Combustion Event | Diesel | Diesel-DME Blend |
-----------------|--------|-----------------|
Start of Injection (Degrees ATDC) | -14    | -14             |
Start of Ignition (Degrees ATDC)    | 4      | 4               |
End of Burn (Degrees ATDC)          | 58     | 56              |
Max ROHR (J/ crank angle degree)    | 63.83  | 66.27           |

Figure 4-12: Pressure Trace and ROHR - Mode 6
Combustion Event | Diesel | Diesel-DME Blend |
<table>
<thead>
<tr>
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<tbody>
<tr>
<td>Start of Injection (Degrees ATDC)</td>
<td>-10</td>
<td>-10</td>
</tr>
<tr>
<td>Start of Ignition (Degrees ATDC)</td>
<td>6</td>
<td>10</td>
</tr>
<tr>
<td>End of Burn (Degrees ATDC)</td>
<td>75</td>
<td>75</td>
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<tr>
<td>Max ROHR (J/crank angle degree)</td>
<td>69.07</td>
<td>76.36</td>
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</tbody>
</table>

Figure 4-13: Pressure Trace and ROHR-Mode 7
Combustion Event | Diesel | Diesel-DME Blend
--- | --- | ---
Start of Injection (Degrees ATDC) | -13 | -12
Start of Ignition (Degrees ATDC) | 2 | 4
End of Burn (Degrees ATDC) | 70 | 70
Max ROHR (J/crank angle degree) | 89.54 | 87.20

Figure 4-14: Pressure Trace and ROHR-Mode 8
Figure 4–15: Peak Cylinder Pressure Comparison
4.4 Conclusions

The present study was the first attempt to use a blend of Diesel-DME in the Navistar T444E engine at The Pennsylvania State University. The pressurized fuel system was proven as a possible concept in the conversion of diesel engines to operate on diesel-DME blends with minimal modifications. This fuel system will be improved with successive iterations. Drawbacks of this system, which have come to light, are discussed below.

- It was necessary to keep the fuel temperature below 50 °C for DME to remain in the liquid state. It was seen on Mode 8 of the AVL test with the 2% O content fuel blend, that possible vaporization of the fuel caused reduced power output and unsteady operation of the engine. To prevent vaporization of DME, fuel cooling was implemented. In the original design by the manufacturer, the temperature of the fuel is close to 83 °C as it enters the HEUI fuel injector. By the time it is injected into the cylinder, the fuel temperature will have raised to a value much higher than that. This leads to a very good spray breakup, better atomization and mixing. In the modified fuel system for the blend, the fuel temperature is much lower than that designed by the manufacturer. This will perhaps lead to a poorer spray breakup and hence poorer mixing in the cylinder. This problem however, will only arise for very low concentrations of DME in the blend. With increasing amounts of
DME, its boiling point being much lower than diesel, the spray breakup and mixing will be much better even at lower fuel temperatures and injection pressures.

- The temperature of the fuel coming out of the fuel rail was monitored and kept below 50°C. However, the fuel is heated all along its way from the entry into the HEUI injectors to the tip of the injectors exposed to the combustion chamber. This rise in temperature was not recorded. Vaporization of the fuel inside the injectors will also lead to unsteady engine operation as seen in mode 8 of the AVL 8-mode test. This rise in fuel temperature through the injectors should be either modeled or measured to estimate its impact on the injection process.

The pressure trace analysis and comparison of peak pressures does not indicate any significant difference between the burning of diesel and diesel-DME blend with 2% oxygen content. Figure 4.15 compares the peak pressures for engine operation on baseline diesel and diesel-DME blend. The peak pressures appear similar for operation with both fuels.

The viscosity studies show a logarithmic relation of viscosity and pressure as expected. These values can be important in design of the injector nozzle and the injector opening pressure to get an optimized spray pattern. The high miscibility of DME with diesel fuel in any proportion is an encouraging sign and will allow any mixture to be prepared.
Another important issue in use of DME as a fuel is its higher compressibility as compared to Diesel fuel. This will impact the injection process and should be measured for various blends of Diesel fuel and DME. The high pressure viscometer apparatus used for the viscosity work in this thesis can be converted to measure the bulk modulus.

Diesel-DME blends with higher DME concentrations are currently being studied. An optimum blend ratio will be identified with reference to exhaust emissions and fuel system compatibility.
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