Combustion, Performance and Emissions of a Diesel Power Generator with Direct Injection of B7 and Port Injection of Ethanol

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ABSTRACT

This work investigates a diesel engine operating with diesel oil containing 7% biodiesel (B7) and hydrous ethanol with concentrations varying from 5% to 30%. The experiments were conducted in a 49 kW diesel power generator, equipped with an electronic ethanol injection unit installed in the intake manifold and without any modifications in the diesel oil injection system. The results showed a decrease of in-cylinder pressure and net heat release rate with the use of ethanol at low loads and an increase at high loads, in comparison with B7. Increasing ethanol injection caused increased ignition delay and decreased combustion duration. Fuel conversion efficiency was raised up to 13% with the use of ethanol. The use of 30% ethanol in the fuel caused a reduction of carbon dioxide (CO₂) emissions up to 12% and nitric oxide (NO) up to 53%. Carbon monoxide (CO), total hydrocarbons (THC) and oxides of nitrogen (NOₓ) emissions increased with ethanol addition. The replacement of 20% of diesel fuel by ethanol showed the lowest penalties on NOₓ emissions.

Keywords: ethanol; diesel engine; combustion; emissions; power generation.
1 INTRODUCTION

In recent years, the allowable limits of internal combustion engine pollutant emissions has been reduced as a consequence of increasing concern for the environment and air quality. Ethanol is a renewable fuel that can be obtained from plants such as sugar cane and corn. Several authors highlight ethanol characteristics such as the high latent heat of vaporization, high oxygen content and high burning rate, giving it high potential to reduce oxides of nitrogen (NOx) and particulate matter (PM) emissions from diesel engines [1-3]. Brazil is a major ethanol producer [4] and its main source is sugar cane, which has one of the highest ratio of renewable energy produced to fossil energy used in the production of the fuel.

There are different techniques to use ethanol in diesel engines: blended to diesel oil, fumigated in the intake manifold, dual injection of ethanol and diesel oil, spark ignition conversion and surface ignition [5]. Diesel oil demand can be replaced by up to 50% using ethanol fumigation, up to 90% using dual injection and up to 25% using diesel oil-ethanol blends [6]. Compared with diesel oil, the main limitations of the use of ethanol in diesel engines are the ethanol higher auto ignition temperature, the lower cetane number, which difficult the compression ignition, the lower low heating value (LHV), the lower miscibility of the fuels, mainly with the presence of water, and the lower lubricant properties [7-10].

Bodisco [11] reported that the addition of fuels to the intake air for combustion engines, called indirect injection, has been investigated since the late 1920s, with the commercialization of the first dual-fuel vehicle in 1939. The author points out that methanol and ethanol are suitable for use as secondary fuel. Ethanol fumigation in the intake manifold is a technique of dual fuel engine operation, where ethanol is mixed with the intake air while diesel fuel is injected directly into the combustion chamber as the pilot fuel. This method requires little engine modifications and ethanol is injected into the intake manifold by low-pressure nozzles [12,13]. The high compression ratio of diesel engines is an advantage when exploring the high resistance to detonation of ethanol [14].

Padala et al. [14] conducted experiments in an automotive single engine diesel engine using ethanol fumigation. Diesel fuel was injected directly into the combustion chamber and an ethanol injector was installed in the intake manifold with electronically controlled time and duration of injection. Tests were carried out varying the ethanol content up to 70%, compared to diesel oil, based on the energy required for each demanded engine load. The results showed an increase in the ignition delay with the ethanol addition, reduction of CO emissions and increase in emissions of NOx and THC.
Zhang et al. [15] conducted experiments in a four-cylinder diesel engine with direct injection of diesel oil and rated power of 88 kW. An electronic control unit was developed for controlling the injection of ethanol in the intake manifold. The replacement percentages used, in energy basis, were 10% and 20% ethanol. The results showed an increase in fuel consumption, with ethanol fumigation, and reduction of fuel conversion efficiency at low loads and increase at higher loads, when compared with diesel oil performance. Regarding emissions, the authors reported an increase of total hydrocarbons (THC) and carbon monoxide (CO) emissions, as higher percentages of ethanol were used, mainly at low loads. NOx emissions were reduced, attributed to ethanol cooling effect, but carbon dioxide (CO₂) emission increased.

Tutak [16] presented experimental results using E85 (85% ethanol and 15% gasoline) fumigation in a diesel engine, replacing up to 90% diesel oil by E85. For replacement levels above 50%, the engine presented combustion problems. The results showed that E85 fumigation caused lower fuel conversion efficiency at low loads and higher fuel conversion efficiency at high loads. The increase in E85 fumigation increased ignition delay, peak pressure and peak heat release rate. The author evaluated emissions using E85 and found, compared to pure diesel fuel, a reduction in NOx emission at low and medium loads and an increase at full load. The results showed increased CO and THC emissions with increasing percentages of ethanol.

Britto et al. [17] tested a single cylinder diesel engine with compression ratio adjustment with a diesel injector located in the center of the combustion chamber and two ethanol injectors positioned upstream of the cylinder head inlet. The results showed increased CO and THC emissions, attributed to the ethanol cooler effect and non-homogeneous air-ethanol mixture.

This work presents the use of diesel oil containing 7% of biodiesel (B7) directly injected into the engine combustion chamber and hydrous ethanol injection into the engine intake manifold and its effect on combustion, performance and emissions. Ethanol injection was electronically controlled, with concentrations of 5%, 10%, 15%, 20%, 25% and 30%. The experiments were performed in a 44 kW diesel power generator.

2 METHODOLOGY

2.1 Experimental Setup

The main characteristics of the 44 kW diesel engine used in this study shown by Tab. 1. The original engine geometry characteristics and the mechanically controlled diesel oil direct injection system
were not changed for the tests. An electronic injection system was adapted to the engine intake system [18] to control ethanol injection in the dual-fuel operation. The electronic injection system consists mainly of a magnetic sensor and a knock sensor, for engine phase synchronization, and microcontrollers, to control the injection timing, according to engine load demand. A common rail system was developed and adapted to the engine intake manifold for multiport ethanol injection, ensuring that the same amount of fuel is available for each cylinder. The ethanol injectors were positioned 130 mm from the intake valves. The tests were conducted with fixed injection timing of diesel fuel, at 23 crank angle (CA) degrees before top dead center (BTDC), and ethanol injection at top dead center (TDC), in the beginning of the intake stroke.

**Table 1** Diesel engine and generator details

<table>
<thead>
<tr>
<th>EQUIPMENT</th>
<th>PARAMETER</th>
<th>TYPE OR VALUE</th>
</tr>
</thead>
<tbody>
<tr>
<td>ENGINE</td>
<td>Cycle</td>
<td>Four strokes</td>
</tr>
<tr>
<td></td>
<td>Diesel oil injection</td>
<td>Direct</td>
</tr>
<tr>
<td></td>
<td>Bore × stroke</td>
<td>102 mm × 120 mm</td>
</tr>
<tr>
<td>Number of cylinders</td>
<td>4, in line</td>
<td></td>
</tr>
<tr>
<td>Intake system</td>
<td>Naturally aspirated</td>
<td></td>
</tr>
<tr>
<td>Rated power</td>
<td>44 kW</td>
<td></td>
</tr>
<tr>
<td>Number of poles</td>
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<td></td>
</tr>
<tr>
<td>Voltage</td>
<td>220 V</td>
<td></td>
</tr>
<tr>
<td>GENERATOR</td>
<td>Number of phases</td>
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<tr>
<td>Rated power</td>
<td>55 kVA</td>
<td></td>
</tr>
<tr>
<td>Frequency</td>
<td>60 Hz</td>
<td></td>
</tr>
</tbody>
</table>

The intake air mass flow rate was measured through an orifice plate, with maximum uncertainty of ± 2.3 kg/h. The temperatures of ethanol, diesel oil, ambient air, inlet air, orifice plate inlet, exhaust gas and cooling water temperature were monitored using K-type thermocouples. The maximum uncertainty of the measured exhaust gas temperature was ± 7°C; the remaining temperatures were measured with maximum uncertainty of ± 2°C. The inlet air humidity was measured through a thermo-hygrometer with
uncertainty of ± 2.5%, and the ambient pressure was determined through a Torricelli barometer with resolution of ± 1.3 kPa. Diesel oil and ethanol consumption were measured through platform balances, with resolutions of 5 g and 1 g, respectively.

The exhaust gas concentration of total unburned hydrocarbons (THC) was measured by a heated flame ionization detector (HFID) analyzer, with resolution of ± 1 ppm. Oxides of nitrogen (NOx) and nitric oxide (NO) concentrations were measured by a heated chemiluminescent detector (HCLD) analyzer, with resolution of ±1 ppm. Carbon monoxide (CO) and carbon dioxide (CO2) concentrations were determined by non-dispersive infrared (NDIR) analyzers, with resolutions of ±1 ppm and ± 0.01%, respectively. In-cylinder pressure was measured by a piezoelectric pressure transducer with resolution of ± 0.5%. A trigger wheel with a magnetic sensor was used to synchronize the pressure data with the engine cycle. The system was set to obtain the pressure data at 0.1 crank angle intervals. Figure 1 shows a schematic draw of the measurement system. The total uncertainty of the results presented is a combination of both the statistical spread and the instrument uncertainty.

Fig 1 Schematics of the experimental apparatus
2.2 Experimental procedure

The fuels used during the tests were diesel oil containing 7% of biodiesel (B7) and hydrous ethanol (92.6% purity). The replacement percentages used were 0% (B7E0), 5% (B7E5), 10% (B7E10), 15% (B7E15), 20% (B7E20), 25% (B7E25) and 30% (B7E30). The tests were conducted at the constant engine speed of 1800 rev/min. The loads were varied from 0 kW to 37.5 kW and the readings at each load were performed after the engine reached the steady state condition, observing the exhaust gas and the cooling water temperatures. The tests were conducted according to ISO 3046-1:2002 standard [20], which shows the correction of the load power and fuel consumption results to standard conditions.

To set the ethanol injection system, the injected ethanol mass was initially estimated considering the same fuel conversion efficiency for standard operation with diesel oil (B7). Thus, the energy contained in a percentage of B7 mass amount consumed at a specific operating condition in the original engine configuration can be provided by a replacement mass amount of ethanol according to [19]:

\[
m_E = P_{\%} \cdot \frac{m_{B7} \cdot Q_{LHV,B7}}{Q_{LHV,E}}
\]

where \(m_E\) is the ethanol mass flow rate (kg/h), \(m_{B7}\) is B7 mass flow rate (kg/h), \(Q_{LHV,B7}\) is the low heating value of B7 (MJ/kg), \(Q_{LHV,E}\) is the low heating value of ethanol (MJ/kg) and \(P_{\%}\) is the percentage of ethanol replacement.

The energy content of both fuels, B7 and ethanol, were considered for calculation of the fuel conversion efficiency in dual fuel operation [15,16]:

\[
\eta_e = \frac{3600 \cdot W}{m_{B7} \cdot Q_{LHV,B7} + m_{E} \cdot Q_{LHV,E}} \cdot 100\%
\]

where \(\eta_e\) is the engine brake fuel conversion efficiency (%) and \(W\) is the engine output power (kW).

The concentration of the exhaust gas component is measured in % or ppm. To convert molar concentration into specific mass concentration per unit of energy produced, in g/kW.h, the following expression is used:
\[ c_{g/kW.h} = c_{ppm \%} \cdot (m_{\text{air}} + m_{\text{gas}}) \cdot \frac{1}{\rho_{\text{ex}}} \frac{W}{\rho} \]  

(3)

where \( c_{g/kW.h} \) is the component specific composition (g/kW.h), \( c_{ppm \%} \) is the component composition (ppm \( \times 10^6 \) or \% \( \times 10^{-2} \)), \( m_{\text{air}} \) is the intake air mass flow rate (kg/h), \( \rho_{\text{C}} \) is the gas component density (kg/m\(^3\)) and \( \rho_{\text{ex}} \) is the exhaust gas density (kg/m\(^3\)).

The in-cylinder pressure data was used to determine the ignition delay, the combustion duration and the net heat release rate. The ignition delay is determined from the time interval between the start of injection and the start of combustion [21]. The start of combustion was determined from the second order derivative of the in-cylinder pressure diagram [22]. Combustion duration was determined as the crank angle interval between the beginning of heat release and the crank angle where there is an accumulation of 95% of the total amount of heat released [6,23]. The heat release analysis was made for provide the effects of the ethanol on combustion characteristics. The apparent net heat release rate was calculated from application of the first law of Thermodynamics [21]:

\[ \frac{dQ_n}{d\theta} = \left( \frac{\gamma}{\gamma - 1} \right) \cdot p \cdot \frac{dV}{d\theta} + \left( \frac{1}{\gamma - 1} \right) \cdot V \cdot \frac{dp}{d\theta} \]  

(4)

where \( dQ_n/d\theta \) is the apparent net heat release rate (J/°CA), \( \gamma \) is the ratio of specific heats, \( c_p/c_v \), \( p \) is the cylinder pressure (Pa), \( \theta \) is the crank angle (°CA) and \( V \) is the cylinder gas volume (m\(^3\)).

3 RESULTS AND DISCUSSION

Figures 2 and 3 show the in-cylinder pressure history and the net heat release rate curves as a function of crank angle and the amount of ethanol injected into the engine intake manifold for the loads of 0 kW and 37.5 kW, respectively. The curves follow the same trend but with a shift of the peak pressure with increasing amount of ethanol due to increased ignition delay. At low loads, the lower temperature of the cylinder gas and the longer ignition delay lead to a peak heat released displaced from the TDC, causing decreased peak pressure.
At the load of 0 kW, the peak cylinder pressure is generally reduced with increased ethanol content in the fuel (Fig. 2). The peak heat release rate peaks does not show a particular trend with ethanol content in the fuel. Despite the larger amount of accumulated fuel during the ignition delay, the concentrations above 5% of ethanol did not show significant increase of peak heat release rate. This can be explained by a poor combustion due to the higher cooling effect caused by ethanol, dominant at low loads, causing also reduced peak pressure.

At high loads, the ethanol-diesel oil-air mixture is richer and more fuel is burned during the premixed combustion phase (Fig. 2). In addition, the cylinder gas temperature is higher and the ignition delay is shorter. This behavior causes an increase in peak pressure and heat released. With increasing engine load, the effect of higher amount of air-fuel mixture formed during the ignition delay and burned in the premixed phase is more evident, being dominant for pressure increase with the increase of ethanol concentration [25].

![In-cylinder pressure and heat release rate at 0 kW](image)

**Fig 2** In-cylinder pressure and heat release rate at 0 kW

Figure 3 shows the results of in-cylinder pressure and heat released for the load of 37.5 kW. There is an increase in peak pressure with increasing ethanol content in the fuel. At this load, the difference in the net heat release rate between different ethanol concentrations is amplified, with an increase in the magnitude of the peak pressure and displacement form TDC. This is caused by the fuel accumulated during the ignition delay and the faster burning of ethanol [13]. Several authors have found similar behaviors of in-cylinder
The increase in heat release can increase local temperatures of the cylinder structure, requiring the evaluation of the engine cooling system efficiency.

Figure 3 In-cylinder pressure and heat release rate at 37.5 kW

Figure 4 shows increased ignition delay with increasing amount of injected ethanol. The high latent heat of vaporization of ethanol decreases in-cylinder gas temperature, thus reducing the cetane number of the ethanol-diesel oil mixture and, as a consequence, increasing the ignition delay [6,13,14,16,24]. Furthermore, when B7 is injected in the cylinder it is not taken by air only, but by an ethanol-air mixture with lower oxygen concentration when compared with pure air, thus increasing the ignition delay. The in-cylinder gas dilution is increased with increasing ethanol injection [14,16]. The results also shows that engine load influences ignition delay, since it is dependent on the cylinder gas temperature and pressure conditions [21]. For B7E0 operation the ignition delay decreased at high loads, due to high in-cylinder temperature. For ethanol operation, at low loads, the in-cylinder temperature is low and the cooling effect of ethanol fumigation is dominant. At high loads and high ethanol concentration, the gas dilution effect and the low cetane number of ethanol increase the ignition delay. The shortest ignition delay, 21.7°CA, was verified for B7E0 fuel at 37.5 kW, and the longest ignition delay, 26.2°CA, occurred to B7E30 at the same load.
**Fig 4** Variation of ignition delay with ethanol concentration and load power

Figure 5 shows the combustion duration as a function of engine load and fuel. Increasing ethanol concentration decreased the combustion duration, justified by the higher heat release rate in the premixed combustion phase caused by the addition of ethanol in the air and increased ignition delay. The addition of ethanol increases the amount of oxygen in the mixture, which reduces the pyrolysis process and increases oxidation during combustion, reducing the combustion duration [23].

**Fig 5** Variation of combustion duration with ethanol concentration and load power
Figure 6 shows the fuel conversion efficiency results for the different ethanol concentrations and engine load. Ethanol fumigation in diesel engine affects its performance by different factors. The gases inside the cylinder are cooled when ethanol is admitted during the intake stroke, which causes poor air-fuel mixture formation and tends to reduce the fuel conversion efficiency. The addition of ethanol tends to increase the ignition delay, as shown in the above results, causing a larger fuel amount to burn during the premixed phase of combustion. Furthermore, the air-ethanol mixture burns faster, compared with B7, which may increase the fuel conversion efficiency by reducing the time for heat exchange between the gases and the combustion chamber walls [13,26]. These factors act simultaneously and are dependent on ethanol percentage in the fuel and engine load.

![Graph of Fuel Conversion Efficiency](image)

**Fig 6** Variation of fuel conversion efficiency with ethanol concentration and load power

In general, there was a reduction of fuel conversion efficiency at low loads and increase at high loads (Fig. 6). At low loads the cooling effects of ethanol vaporization are dominant, decreasing the in-cylinder gas temperature. Besides, the low amount of ethanol injected form an air/fuel mixture that can be too lean to allow combustion to proceed, resulting in reduced fuel conversion efficiency [16]. At high loads, the larger quantity of fuel burned increases the cylinder gas temperature. These effects, together with the faster burning of ethanol and increased amount of fuel burned in the premixed phase of combustion, in
comparison with B7, are determinant to decrease or increase the fuel conversion efficiency [1,13,26,27]. In addition, at low loads, the reduction of the peak pressure and heat release rate could have contributed to reduce the fuel conversion efficiency. At high loads, there was an increase of these two parameters, having, therefore, increasing fuel conversion efficiency. The largest reduction occurred for B7E20 at the load of 10 kW, when the fuel conversion efficiency was 6.6% lower than operation with B7E0, and the largest increase was 13.0%, found for operation with B7E30 at the load 37.5 kW.

Figure 7 shows the behavior of the specific CO2 emission for the different ethanol concentrations and loads. As the load was increased, CO2 emission was reduced for all ethanol concentrations. At a given load, in general, increasing ethanol concentration in the fuel also caused a reduction of CO2 emission. The reductions obtained for CO2 emissions were up to 12.2%, for B7E25. The decrease of cylinder gas temperature due to the high ethanol latent heat of evaporation inhibits CO oxidation, which leads to a reduction of CO2 emission [14,15]. The presence of oxygen and a single carbon bond in ethanol molecule increases fuel conversion efficiency (Fig. 6), helping to reduce CO2 production (Fig. 7).

![Fig 7 Variation of specific carbon dioxide emissions with ethanol concentration and load power](image)

Figure 8 shows that CO specific emissions was increased with ethanol concentration in the fuel for all engine loads tested. Zhang et al. [15] explain that CO emission tend to increase in situations where there is incomplete combustion, under low temperatures and very lean mixtures, and is controlled by the
local equivalence ratio, being an indicative of combustion quality. CO oxidation occurs at high combustion temperatures. At low loads, the addition of ethanol leads to a reduction of cylinder gas temperature, resulting in quality loss and lower combustion oxidation of CO to CO₂ in the expansion stroke, increasing the CO emission [12,15]. With the addition of ethanol, there is a reduction of excess air in the cylinder, which has a strong influence to increase CO emission [3]. At high loads, the effect of ethanol vaporization to decrease cylinder gas temperature is reduced, due to the higher combustion temperatures attained, thus reducing the differences of CO emissions between the ethanol blends and B7.

![Graph: Variation of specific carbon monoxide emissions with ethanol concentration and load power](image)

**Fig 8** Variation of specific carbon monoxide emissions with ethanol concentration and load power

Figure 9 shows the specific THC emission for the different ethanol concentrations. As it occurred for CO emissions, THC emission increased with the addition of ethanol. Unburned hydrocarbon formation is strongly dependent on combustion quality; thus, the results indicate the use of ethanol provokes poor combustion. Several factors may have caused the increase of THC emission, such as temperature reduction due to ethanol injection, which causes lower rates of vaporization and mixture formation. For Chauhan et al. [9], at low loads, the indirectly injected ethanol is unable to impinge on surfaces and be distributed, causing poor fuel distribution, large amounts of excess air, low exhaust temperature and lean air-fuel mixtures tend to escape, resulting in higher unburned fuel. The air-ethanol mixture trapped in cylinder crevices during the compression and combustion processes, under increasing cylinder pressure, returns to
the cylinder during the expansion and exhaust strokes, under decreasing cylinder pressure, and partially undergoes a post-flame oxidation process. The unburned amount escape though the exhaust valve and contribute to the increase of hydrocarbon emissions, a characteristic phenomenon of port fuel spark ignition engines [13,27]. At high loads, the temperature increase allows more burning of the air-fuel mixture, improves the quality of combustion and reduces the increase of hydrocarbon emission [13]. Several researchers, such as Surawski et al. [3], Chauhan et al. [9], Tutak [16] and Yao et al. [27], also report the increase of CO and THC emissions.

![Variation of specific total hydrocarbon emissions with ethanol concentration and load power](image)

**Fig 9** Variation of specific total hydrocarbon emissions with ethanol concentration and load power

Figure 10 shows that NO emission was reduced for all loads when ethanol was used as fuel. The reductions attained were up to 53.2%, for B7E30. Surawski et al. [3] also found reduction of NO emission for all engine power range with ethanol concentration up to 40%. Several factors affect NOx formation and emissions in a diesel engine. The oxygen content and low cetane number of ethanol lead to high temperature peaks and cause increased NOx, but, on the other hand, the high enthalpy of vaporization and low flame temperature of ethanol lead to decreased NOx emissions. The effects of ethanol in NO emissions are dependent on engine operating conditions and fuel characteristics [28]. Several authors associated reduction of NO emissions to decreased gas temperature caused by ethanol fumigation [15]. This effect is more significant at low loads, since at high loads richer air-fuel mixtures tend to increase combustion temperature.
Despite the reduction of NO emission with ethanol fumigation, it was observed an increase of specific NO\textsubscript{x} emissions for all loads (Fig. 11). The increase in NO\textsubscript{x} emission is due to the increase of NO\textsubscript{2} formation, as Padala et al. [14] and Tutak [16] showed it. This behavior can be caused by the cooling effect of ethanol, as the decrease in temperature favors NO\textsubscript{2} formation. Heywood [21] explains that one of nitrogen dioxide formation mechanism is by the combination of NO formed in the flame front with hydroperoxyl radical (HO\textsubscript{2}), also producing hydroxyl (OH). NO\textsubscript{2} would subsequently combine with oxygen element (O) to form NO and molecular oxygen (O\textsubscript{2}), unless the NO\textsubscript{2} formed in the flame is quenched by mixing with cooler fluid.

Zhu et al. [23] explain that the NO oxidation efficiency depends on the formation of hydroperoxyl radical. The authors cite studies that shows that ethanol could produce HO\textsubscript{2} through the thermal degradation behavior of alcohols with NO, and that HO\textsubscript{2} is promptly formed during oxidation of oxygenated fuels. Thus, ethanol can be a source of HO\textsubscript{2} and increase the oxidation of NO into NO\textsubscript{2}, especially at low and medium loads. Furthermore, their results showed significant increases of unburned fuel emission, which favors the formation of NO\textsubscript{2} due to the presence of ethanol, a good oxidizing [15], and due to the cold fuel condition, which inhibits the conversion of NO\textsubscript{2} to NO [21].
From an analysis of Figs. 10 and 11 for each fuel curve when the engine operated with ethanol, there is a tendency of increasing NO emissions for loads below 10 kW and no single trend is observed for higher loads. From the difference of NOx and NO concentrations, it can be inferred that NO2 concentration always increase with decreasing load. At low loads the relatively low temperatures attained in the cylinder do not favor the decomposition of NO2 into NO, while, as the load and combustion product gas is increased, a larger amount of NO2 is decomposed into NO.

![Graph](image)

**Fig 11** Variation of specific oxides of nitrogen emissions with ethanol concentration and load power

The studied diesel engine does not have any exhaust gas aftertreatment technology. Thus, when operating with diesel oil only, the engine does not meet the current Brazilian emissions standards for diesel engines, PROCONVE (Air Pollution Control Program for Motor Vehicles) P7, based on Euro V European standards for heavy-duty engines, which requires systems as exhaust gas recirculation (EGR) or selective catalytic reduction (SCR). Ethanol addition to B7 increased CO2, THC and NOx emissions, however aftretreatments systems could lead to a reduction in these emissions. SCR system is a well-stablised technique to achieve very low levels of NOx [29-31]. Tadano et al. [29] showed reductions up to 93% in NOx using SCR systems in a diesel engine. EGR systems are also very effective in reducing NOx emissions [32-33], but increases smoke emissions, which can be reduced by ethanol addition to the fuel [33-35].

Another extensive used aftertreatment system is diesel oxidation catalyst (DOC), used to oxidize THC and CO emissions to H2O and CO2, respectively, for diesel oil with low sulfur content [36]. This
system is able to reduces CO and THC emissions to the standards limits [36-42]. Zhang et al. [41] and Wei et al. [42] also showed that alcohol fumigation in diesel engines increased CO, NO\(_2\) and THC emissions, but, after catalytic conversion, the concentrations of these components were significantly reduced, up to 38% for THC and 16% for CO [41], and for levels near diesel fuel operation with DOC system [42]. For diesel fuel operation, Wei et al. [42] showed reductions up to 90% in CO and THC emissions for high loads.

4 CONCLUSIONS

Ethanol was added to B7 in a diesel engine by fumigating in the intake manifold and without any change in the original engine geometry characteristics and in the mechanically controlled diesel oil direct injection system. Ethanol addition increased ignition delay and reduced combustion duration. Peak cylinder pressure and heat release rate were reduced at low loads and increased at high loads. The fuel conversion efficiency was reduced at low loads up to 6.6%, using 20% ethanol in the fuel, and increased up to 13.0% at high loads, for 30% ethanol. CO\(_2\) and NO emissions were reduced in all engine load range tested, reaching up to 12% CO\(_2\) reduction and 53% NO reduction with 30% ethanol concentration in the fuel. CO, THC and NO\(_x\) emissions always increased with the use of ethanol. The partial substitution of 20% of diesel fuel by ethanol fumigation at moderate to high loads in an engine with unmodified diesel injection settings is recommended for the lowest penalties on NO\(_x\) emissions. In general, ethanol fumigation in a stationary diesel engine may require optimization of diesel injection and/or exhaust gas aftertreatment to meet emissions standards.
ACKNOWLEDGMENTS

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Prof. Francisco R. Cunha  
Editor-in-Chief  
Journal of the Brazilian Society of Mechanical Sciences and Engineering

Dear Editor,

Thank you for the Reviewers’ useful comments and suggestions on the structure of our manuscript. We have modified the manuscript accordingly, and detailed corrections are listed below point by point:

Reviewer #1: The previous comments and indications are well reflected on the revised manuscript. Therefore, I would like to recommend this revised version as an accepted paper for the journal of BSME after the correction according to following editing comment.

Comments
In the manuscript, please remove the section numbers of 5. Acknowledgement and 6. References in pages 19 as Acknowledgement and References.

Authors’ response: The number 5 was removed from the Acknowledgments section title, and the number 6 was removed from the References section title, on page 19.

Reviewer #2: I agree with revisions done by the authors and suggest the paper be accepted for publishing.

The manuscript has been resubmitted to your journal. We look forward to your positive response.

Sincerely yours,

Prof. J. R. Sodré, Ph.D.
Corresponding author