Abstract

Introduction

Experimental setup and methodology

Question 1

Question 2

Average emissions of CO, CO₂, $\overline{NO_x}$

In order to calculate the average CO_2 and NO_{x} emissions, the emission data acquired from the sensors are plotted against time for every test. When possible, the stable region with low variance is selected to average the emissions. The calculated CO_2 and NO_{x} emission means for each test and average of each spark ignition time can be seen in Table 1 and 2.

Gasoline						
Spark time	Tests	CO_2 emission	NO_x emission			
		concentration (%)	concentration (ppm)			
10 CAD	Test 1	14.4167	3162			
BTDC	Test 2	14.4311	3155			
	Test 3	14.4815	3110			
	Average	14.4331	3142			
13 CAD	Test 4	14.4167	3162			
BTDC	Test 5	14.4732	3246			
	Test 6	14.7239	2799			
	Average	14.7197	2781			
16 CAD	Test 7	14.6996	751			
BTDC	Test 8	14.6944	243			
	Test 9	14.6778	146			
	Average	14.6906	380			

Table 1: CO_2 and NO_{x} emissions for all tests for gasoline.

Ethyl Valerate 10%						
Spark time	Tests	CO_2 emission	NO_x emission			
		concentration (%)	concentration (ppm)			
10 CAD	Test 10	13.5300	2916			
BTDC	Test 11	13.5463	2192			
	Test 12	13.5418	1126			
	Average	13.5394	2079			
13 CAD	Test 13	13.5274	98			
BTDC	Test 14	13.5229	48			
	Test 15	13.5183	31			
	Average	13.5229	56			
16 CAD	Test 16	13.4397	20			
BTDC	Test 17	13.4464	14			
	Test 18	13.4266	13			
	Average	13.4556	16			

Table 2: CO_2 and NO_{x} emissions for all tests for Ethyl-Valerate.

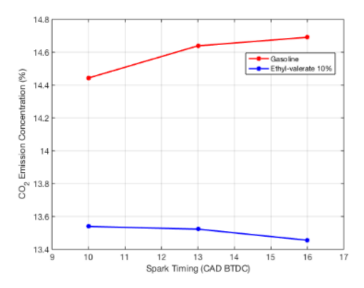


Figure 1: CO_2 emissions plotted for varying spark ignition timings for both fuels.

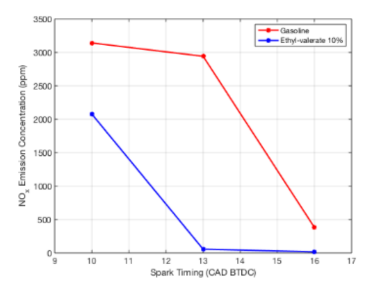


Figure 2: NO_x emissions plotted for varying spark ignition timings for both fuels.

Looking at Figure 1, it is very clear that as the sparking time increased, gasoline showed an increasing $\rm CO_2$ emission trend while the ethyl-valerate had a decreasing trend; the $\rm CO_2$ emission for gasoline peaked at 16 CAD BTDC where the latter peaked at 10 CAD BTDC. In addition, it should be noted that gasoline always had a larger emission of $\rm CO_2$ than ethyl-valerate from 10 to 16 CAD.

As shown in Figure 2, through out the tests gasoline had higher emissions of NO_x compared to ethylvalerate. Unlike CO_2 emissions, both gasoline and ethyl-valerate gave a similar response for NO_x emissions as emissions for both decreased with earlier spark ignition timing. On the other hand, the emission of NO_x for gasoline had a significant drop between 13 and 16 CAD BTDC while the ethyl-valerate experienced a drastic decline between 10 to 13 CAD BTDC. The NO_x emissions for ethyl-valerate between 13 and 16 CAD BTDC is comparatively lower than any other value.

Overall, as highlighted in Figure 1 and 2, gasoline consistently had significantly larger emission of both NO_x and CO_2 between 10 to 16 CAD BTDC, this phenomenon can be explained by the greater number of carbons in gasoline; the molecular formula of ethyl-valerate is $C_7H_{14}O_2$ CITATION and the main component of gasoline is octane, which has a molecular formula of C_8H_{18} CITATION. Greater number of carbons means more CO_2 would be produced and larger amount of oxygen is required, since the formation of NO_x is highly temperature and O_2 concentration dependent, high temperatures and O_2 concentrations will result in high NO_x formation rate. As more oxygen is obtained during the chemical reaction of air with gasoline than the reaction of air with ethyl-valerate, more NO_x is released by gasoline CITATION.

Peak heat release rate

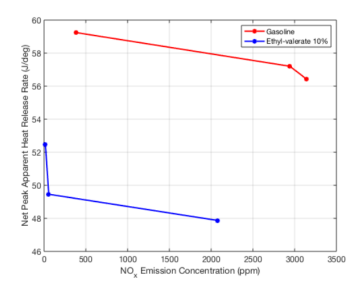


Figure 3: Plot of net peak heat release rate against NO_x emissions for both fuels.

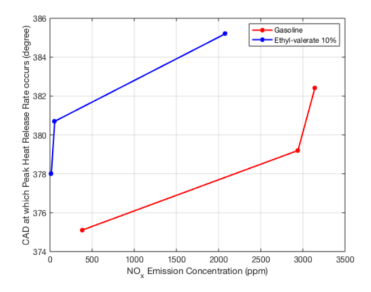


Figure 4: Plot of CAD at which peak heat release rate occurs against NO_x emissions for both fuels.

Theoretically, $\mathrm{NO_x}$ emissions should increase as the peak heat release rate rises. As described previously, high temperature, pressure, and oxygen concentration would result in higher $\mathrm{NO_x}$ formation rate and therefore higher emissions. However, results shown in Figure 3 give a completely opposite response, the emission of $\mathrm{NO_x}$ for both gasoline and ethyl-valerate showed a tendency to decrease as the peak heat release rate increased.

There are a few errors that could have caused errors in calculations.

- 1. In calculations, the assumptions of a perfectly adiabatic system and stoichiometric combustions are made. However, in reality the system is not perfect and there are frictional and environmental heat losses.
- 2. As mentioned previously, as the emissions are recorded with respect to time, emissions are averaged in a stable region when possible. However, emissions didn't reach equilibrium for all tests and were unstable across the cycle. In some of the tests, emissions data displayed decreasing trend over time and didn't become steady. As seen from Table 1, for some spark ignition time

tests, there are significant cycle-to-cycle variations in emissions. For more accurate data, more tests with a steady emission level should be conducted.

- 3. It is possible that the measuring instrument used to detect NO_x and CO_2 emissions may have errors, especially calibration error.
- 4. The heat released may not be uniformly distributed, which makes the measurement inaccurate and leads to deviations from the calculated values.

Question 3

The lower heating value is the amount of heat released in the complete combustion of a specific quantity of fuel at standard conditions (25 °C and 1 bar), without recovering the latent heat of vaporization from the water vapour content in the gaseous products CITATION. The LHV is preferred to HHV in relation to internal combustion engines as it is not practical to recover the energy contained in the gaseous water vapour in typical ICEs, as this would require cooling the combustion products until the water vapour condenses. In most engines the vapour is expelled from the system as part of the hot exhaust gases and is therefore unavailable to do useful work.

Heating values may be determined using a calorimeter or derived empirically by considering an energy balance, as will be done here. The heating value of the renewable fuel, ethyl valerate $(C_7H_{14}O_2)$, was estimated by considering its stoichiometric combustion in dry air. The chemical equation is presented as follows:

$$C_7H_{14}O_2 + 9.5(O_2 + 3.76N_2) \longrightarrow 7CO_2 + 7H_2O(g) + 35.72N_2$$
 (1)

Assuming perfect homogenous mixing of fuel and air, 9.5 mol of dry air (21% oxygen and 79% nitrogen by volume) is required per mole of fuel to achieve complete combustion. The heat released is therefore the difference between the enthalpies of the products and reactants CITATION.

$$Q = \sum_{P} \left(n_i \overline{h_i} \right) - \sum_{R} \left(n_i \overline{h_i} \right) = H_P - H_R \tag{2}$$

The enthalpy $\overline{h_i}$ at any state is a function of temperature and pressure, consists of the enthalpy of formation h_f^{\ominus} at standard reference conditions and sensible enthalpy. For simplicity, only the enthalpy of formation was considered in the estimation as the exhaust temperatures are not known. This causes the heat release to be overestimated and leads to conservative results for thermal efficiency.

According to the National Institute of Standards and Technology, the enthalpy of formation of the test fuel is $\Delta h_f^{\ominus} = -553 \,\mathrm{kJ} \,\mathrm{mol}^{-1}$ CITATION. h_f^{\ominus} of the other compounds were obtained from property tables CITATION. From (1) and (2), the LHB of ethyl valerate ($\mathrm{C_7H_{14}O_2}$) is computed as follows:

$$LHV_{\text{test}} = \left[-553.1 - 9.5 (0+0) \right] - \left[7 (-393.52) + 7 (-241.82) + 35.72 (0) \right] = 3894.4 \,\text{kJ} \,\text{mol}^{-1} \quad (3) \, \text{mol}^{-1} \quad (3) \,$$

Taking the molecular weight of ethyl valerate to be 130.185 kg mol⁻¹, the LHV of the test fuel expressed in mass basis is:

$$LHV_{\rm C_7H_{14}O_2} = 29.91 \,\rm MJ \, kg^{-1}$$
 (4)

The LHV of the reference fossil gasoline is as provided:

$$LHV_{\rm G} = 42.7 \,\rm MJ \, kg^{-1}$$
 (5)

Therefore, the LHV of the fuel blend is the sum of the LHVs of the ethyl valerate and gasoline,

weighted by mass fractions (x_i) , which can be computed from their volume fractions (v_i) CITATION:

$$LHV_{\text{blend}} = x_G \cdot (LHV)_G + x_{\text{C}_7\text{H}_{14}\text{O}_2} \cdot (LHV)_{\text{C}_7\text{H}_{14}\text{O}_2}$$

$$\tag{6}$$

$$LHV_{\text{blend}} = x_G \cdot (LHV)_G + x_{C_7 H_{14} O_2} \cdot (LHV)_{C_7 H_{14} O_2}$$

$$x_G = \frac{v_G \rho_G}{v_G \rho_G + v_{C_7 H_{14} O_2} \rho_{C_7 H_{14} O_2}}$$
(6)

$$\therefore LHV_{\text{blend}} = 0.88267 (42.7) + (1 - 0.88267) (29.913) = 41.20 \,\text{MJ} \,\text{kg}^{-1} \tag{8}$$

The indicated specific fuel consumption ISFC is a measure of the amount of fuel used per unit of power output from the engine. It is given by the following equation CITATION:

$$ISFC = \frac{\dot{m}_f}{\dot{W}} \tag{9}$$

Where \dot{m}_f is the mass flow rate of fuel and \dot{W} is the power output. The power output of the engine may be derived from the indicated mean effective pressure IMEP calculated in Question 1, which is the average pressure required in the cylinder to produce the same amount of work throughout the 4 strokes. As IMEP does not factor friction losses, the power output is overestimated hence the actual fuel consumption is higher. The work done per power stroke is expressed as:

$$W_{\text{power}} = IMEP \times \text{Bore Area} \times \text{Stroke} = IMEP \times \text{Swept volume}$$
 (10)

The Ricardo E6 is a 4-stroke engine running at test conditions of 600 rpm, therefore there is 1 power stroke every 2 crankshaft rotations, i.e. 5 power strokes per second. The power output in kW is:

$$\dot{W} = 5\dot{W}_{\text{power}} \tag{11}$$

The overall indicated thermal efficiency for the engine is by definition the useful work output over total heat input. It is a measure of how much of the fuel's heating value is lost to incomplete combustion, heat loss to surroundings and friction, and other losses. Taking the cylinder to be the system, the heat input is simply the heat released during combustion within the cylinder, equal to mass flow rate of fuel times the fuel heating value.

$$\eta_{th} = \frac{\dot{W}}{\dot{Q}_{in}} = \frac{\dot{W}}{\dot{m}_f \times LHV_{\text{blend}}} \tag{12}$$

Therefore, the efficiency and ISFC are related by the following:

$$\eta_{th} = \frac{1}{ISFC \times LHV_{\text{blend}}} \tag{13}$$

Using the above relations, the calculated values for ISFC and indicated thermal efficiency for each spark timing are presented in Table 3, with the corresponding IMEP for reference. The thermal efficiencies were plotted against spark timing in Figure 5.

Fuel	Spark time (deg BTDC)	IMEP (bar)	Indicated specific fuel consumption $(kg J^{-1})$	Indicated thermal efficiency
Reference	10	8.0096	8.0218×10^{-8}	0.292
fossil	13	8.1568	7.7742×10^{-8}	0.302
gasoline	16	8.1614	7.8758×10^{-8}	0.297
10% Ethyl	10	7.6880	7.8958×10^{-8}	0.307
valerate	13	7.8622	7.8158×10^{-8}	0.311
	16	7.9461	7.6111×10^{-8}	0.319

Table 3: ISFC consumptions and indicated thermal efficiencies.

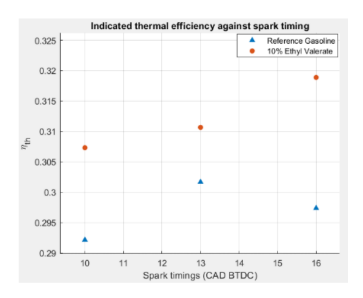


Figure 5: Plot of indicated thermal efficiency against spark timing

It is observed that thermal efficiency of the engine running on the fuel blend is higher than that of gasoline at all spark timings. Although the engine produces less work on the blend, the heat input requirements are also lower.

The trend shows that in general, the indicated thermal efficiency increases with retarded spark timings. This is to be expected as earlier spark timings resulted in higher IMEPs, which translates to a higher indicated power output. Spark timings that are too advanced result in the gas expanding when the piston is already moving down from TDC, hence work done by the gas is not maximised. However, if the spark timing is too early, the gas expands against the piston when it is moving up towards TDC and it is not doing useful work. It has been determined experimentally that optimal power and efficiency for a spark ignition gasoline engine is achieved at 31 CAD BTDC (Zareei and Kakaee, 2013) CITATION. Therefore, for the range investigated in this experiment, an increase in indicated thermal efficiency with spark timing was expected to be observed.

For this reason, the efficiency for the gasoline fuel at 13 or 16 CAD BTDC was most likely to be anomalous as it does not fit with expectations of the trend. Although the IMEP showed a linear increase with later spark timings, the ISFC of the 13 CAD BTDC trial was significantly lower than 16 CAD BTDC. The reason for this anomaly can possibly be attributed to the measured fuel flow rate during test 4, which ranged from $0.18\,\mathrm{g\,s^{-1}}$ at the beginning of the test to $0.1\,\mathrm{g\,s^{-1}}$ by the end, giving a lower average fuel flow rate compared to the other two tests ran at the same spark timing. Since air mass flow rate remained relatively constant, and the methodology called for the air fuel ratio to remain constant throughout, this must be the result of an error, human or otherwise. This thus increased the average ISFC and thermal efficiency of the gasoline at 13 CAD BTDC. Removing the results of test 4 gives us a revised thermal efficiency closer to 29.6%, which is much more reasonable given the expected trend. The revised thermal efficiencies with test 4 removed were plotted against spark timing in Figure 6.

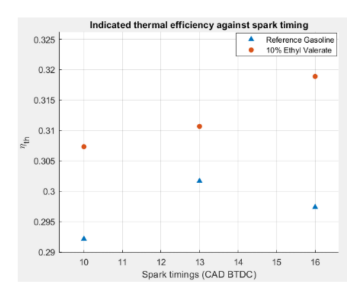


Figure 6: Plot of indicated thermal efficiency with test 4 removed.

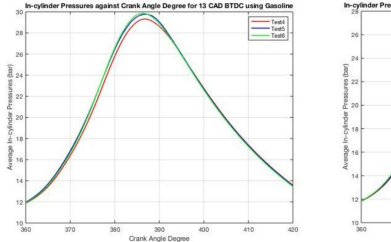
In general, the LHV is an inaccurate estimate for heat input to the system. As mentioned previously the estimated LHV of the fuel blend overestimates the heat released in its combustion, hence the thermal efficiency is likely underestimated. Furthermore, using the product of fuel mass flow and LHV as Q_{in} assumes that perfect combustion occurs, and the fuel is burnt completely. In reality, there will not be a homogenous mixture of air and fuel, hence there will be pockets of unburnt fuel that leave in the exhaust gases. In this manner, Q_{in} is again overestimated.

Additionally, as mentioned previously since the power output is an overestimate, since it was derived from the IMEP which this does not consider frictional losses which may occur in the moving parts of an ICE. Hence, the useful power will be lower in practical purposes and thus the thermal efficiencies calculated are also an overestimate.

Lastly, it is important to consider whether it is appropriate to evaluate the system using thermodynamic analysis. Thermodynamic efficiency of the spark ignition engine is derived from the closed system analysis of the Otto cycle, whilst our experiment deals with a system that undergoes a mechanical cycle instead of a thermodynamic one. Though thermodynamic analysis is not strictly accurate with regards to our system, it is still useful for observing trends within our data. Our analysis allows us to conclude that efficiency increases with increased spark timings, and for all spark timings, the ethyl valerate fuel blend yields a higher thermal efficiency, meaning more energy is converted to useful work.

Question 4

Question 5



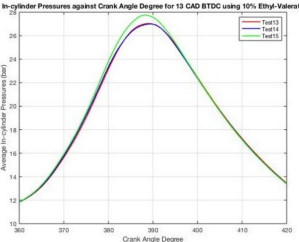


Figure 7:

The three cycles show a variation in the instantaneous in-cylinder pressure cycles. The reason for the cycle-to-cycle variation in in-cylinder pressure is due to the variations in mixture motion, cyclic variations in the local equivalence ratio and the mixture composition near the spark plug. The air-fuel mixture might not be identical for every cycle that takes place which could create the variance in the pressure.

There could be defects created during the manufacturing in the throttle valve of the engine or the defects could be due to wear, which would cause the intake of air-fuel mixture to be inconsistent, affecting the in-cylinder pressure. There could be a larger amount of intake, than what is required, into the combustion chamber which will cause the pressure to be irregular from cycle to cycle as we see the curves in figure 1 have different peaks.

The vary in the speed of the engine could also have an effect on the in-cylinder pressure as in reality, the engine will not run at a constant speed due to discontinuities in flow, resistance and the change in momentum. Therefore, the time duration of the combustion cycle will change as the speed changes which will have an impact on the in-cylinder pressure during the cycle.

Incomplete combustion of the fuel could be a reason for the change in in-cylinder pressure as unburnt fuel left over could affect the cycles after, causing the pressure in the cylinder to change as we get a varied amount of heat release in each cycle due to the inconsistency in the amount of fuel present for combustion.

This cycle-to-cycle variations in in-cylinder pressure are not desirable as they lead to lower efficiency, limitations in power output and higher emissions. The varying pressure leads to varying amount of power output and this could affect the whole system as it may cause it to wear over time. This is due to the change in the pressure causing the components to experience a changing force which could cause damages. This varied power output limits performance and reliability of the engine.

Question 6

Thermal engines present in modern cars rely on the combustion of fuel in oxygen to release energy CITATION. Ideally, this process should involve the complete combustion of all fuel with no excess oxygen; an air-fuel mixture containing the precise ratio of air to fuel required to ensure this is called a stoichiometric mixture. For a gasoline powered vehicle, this ratio is approximately 14.7:1 CITATION. It is important to maintain the air-fuel mixture in the combustion chamber of vehicles as close as possible to the ideal stoichiometric mixture as this ensures that the combustion process is most efficient, with the highest fuel economy and lowest emissions CITATION.

In fact, burning an air-fuel mixture that contains too much fuel and too little air (a rich mixture) will result in some of the fuel not being combusted, but released with the exhaust gases instead. This results in poor fuel economy as a portion of the fuel is wasted, as well as damage to the environment as the release of the unburned hydrocarbon fuels in the atmosphere will cause pollution. On the other hand, the combustion of a mixture that has too much oxygen/air and not enough fuel (a lean mixture) can have even more adverse consequences due to the increase in combustion chamber temperature that this process results in. The high temperatures can cause damage to the vehicle itself, e.g., damaging the pistons and spark plug, as well as promoting the production of Nitrous Oxides (NO_x) which are remarkably damaging to the environment, as they are directly linked to smog and acid rain, and have indirect effects on human health, like causing breathing problems CITATION CITATION.

However, it is of relevance to mention that the stoichiometric ratio is not ideal for all operating conditions; in fact, when the vehicle is operating under a greater load, for instance, when it is accelerating, the engine is required to produce more power, thus a richer air-fuel mixture (a ratio between 13:1 and 12:1 for gasoline powered vehicles) which would produce more power should be burned instead CITATION. Thus, controlling and optimizing the air-fuel ratio is of vital importance for high engine performance and for maintaining high fuel economy as well as reducing emissions. It is the function of O_2 sensors to monitor this ratio in modern cars by gauging the concentration of Oxygen in the exhaust gas (rather than the air-fuel mixture itself) and return this information to the Engine Control Unit (ECU) of the vehicle to control the ratio.

O₂ sensors are electronic transducers located in the exhaust pipe of vehicles. They are used to measure the proportion of Oxygen in the exhaust gas released by the vehicle and compare it to that measured in the atmospheric air in real time CITATION. These sensors are usually "solid-state electrochemical fuel cells [...] that provide a certain output voltage depending on the amount of oxygen in the exhaust system relative to that in the atmosphere" CITATION. The sensor usually operates between two voltages (e.g., between 0.2 V and 0.8 V for narrowband sensors), the lower one associated with a lean mixture, the higher one with a rich mixture, and a middle voltage (0.45 V for narrowband sensors) associated with a nearly stoichiometric, optimal mixture CITATION. The output voltage, alongside with other sensors' readings, is fed back to and processed by the ECU which automatically adjusts the air-fuel mixture as required by injecting more or less fuel into the mixture. The adjusted air-fuel mixture then undergoes combustion and releases new exhaust gases which will then be analysed by the O₂ sensor and the process described above will continually repeat itself to maintain the ideal ratio. The use of this sensor permits closed loop operation of the air-fuel mixture – the information from the sensor is gathered and processed in real time, enabling the ECU to continuously change the air-fuel ratio, if needed, to match the desired one – ensuring optimal combustion conditions and engine function at all times.

In modern vehicles, O_2 sensors are also used to measure the efficiency of catalytic converters. A catalytic converter is a device typically placed in the exhaust system of a vehicle that chemically changes some of the harmful gases (namely NO_x and Carbon Monoxide) present in the exhaust gas of vehicle into more innocuous gases (Nitrogen, Carbon Dioxide and Water) in order to reduce exhaust emissions. By placing one oxygen sensor before the convertor and one after it, the difference in oxygen

concentration between the gas that is yet to pass though the converter and the gas that has already passed through it can be measured, and this difference is used to assess the efficiency of the converter.

Question 7

Part A: On Production

A study published by the Royal Society of Chemistry shows that the esterification of valeric acid (VA) with ethanol catalysed by amino acid ionic liquids is a promising route of production for ethyl valerate (EV)CITATION. VA is obtained from cellulosic materials, a cheap and abundant renewable feedstock for biofuels which, importantly, does not compete with food supply like biofuels derived from starch-based plants.

Several amino acid ionic liquids can be used to catalyse the esterification process, including proline bisulfate (ProHSO₄), glycine bisulfate (GlyHSO₄), and alanine bisulfate (AlaHSO₄). The synthesis of these catalysts involves processes in water without by-products and post-processing, thus these preparations are highly efficient and low cost.

The esterification process catalysed by $ProHSO_4$ shows the most promise, with conversion rate above 99.9% and 100% selectivity for a 7-hour reaction at 80 °C. As a comparison, the esterification process produced no EV in the absence of an acidic catalyst at room temperature and only 67% of EV found using concentrated sulfuric acid at 80 °C. The lipase process took up to 9 days to reach a conversion rate of 97.8% and is thus much less efficient.

The difference is due to the reaction mechanism. The reactants and ionic liquid catalyst are miscible in a single phase before reaction, but the produced EV has low solubility in the ionic liquid, resulting in a biphasic system, in which the unreacted VA and ethanol remained in the ionic liquid layer. Additionally, $ProHSO_4$ acts as water absorbent since water as a by-product is also miscible in the ionic liquid layer, thus shifting the equilibrium of the reaction to the products without water removal. As a result, $ProHSO_4$ facilitates the auto phase separation of the product from the reactants and by-products, favouring the conversion of VA and achieving a high EV yield, whilst restraining the reversible reaction of the ester.

 $ProHSO_4$ is also easily recyclable from the products by a simple removal of water under vacuum taking place at $100\,^{\circ}$ C for two hours. The reused catalyst showed similar catalytic performance to the original, with EV yield remaining above 95% for five additional cycles of the reused catalyst.

The sources of these lignocellulosic biomass from which VA is derived must also be considered when discussing EV as a potential biofuel. The revised Renewable Energy Directive (EU) 2018/2001 emphasizes the direct negative impact the procurement of biomass may have due to indirect land use change (ILUC). The production of biofuel may lead to the extension of agricultural land into areas with high carbon stock such as forests, wetlands and peatlands, the clearing of which may negate the positive effects of greenhouse gas savings from the increased use of biofuels CITATION.

One way to mitigate the impact of ILUC is to produce lignocellulosic biomass from crop residues of agriculture, which does not require the extension of agricultural land. A study published in GCB Bioenergy estimated that in the reference period from 2011 to 2015, the total production of lignocellulosic material from crop residues in the EU28 is 419MT CITATION. From this estimate a sustained supply of crop residue for the production of biofuels can be deduced, though there are other important applications for crop residues, such as those related to soil quality and resilience.

Additionally, the design of the supply chain is an important factor when considering the economic and environmental sustainability of our biofuel, of which the transportation of biomass plays a critical role. As such, the location of residue production within the EU28, the distance between production, processing and consumption hotspots, the timing of biomass supply, and the variability of crop yields year on year due to weather fluctuations are all critical factors to examine. Moreover, the GHG emissions during the transportation of the biomass must also be accounted for in the life cycle assessment of our biofuel.

In conclusion, ethyl valerate shows great promise of displacing gasoline if its production is catalysed by amino acid ionic liquids, namely proline bisulfate ($ProHSO_4$). The time and temperature of the production processes are relatively low whilst achieving conversion and selectivity of nearly 100%. Ethanol is readily available, and the catalyst can be easily mass-produced due to the lack of post processing needed. Additionally, $ProHSO_4$ is easily recyclable and can be reused multiple times with minimal impact on catalytic activity, improving its economic and environmental sustainability.

The main feedstock of lignocellulosic biomass, from which valeric acid is derived, is crop residues from agriculture and they are expected to play a major role in the production of advanced biofuels. Of course, valeric acid is not the only use of biomass both in the context of biofuel production and agricultural applications, though an optimised supply chain will increase the availability and decrease the cost of procurement. The design of the supply chain is a complicated process due to the factors mentioned previously, but also presents exciting opportunities, such as the densification of biomass with techniques such as baling, pelleting, and pyrolysis to reduce logistic costs CITATION. Ongoing research in this field and government incentives such as the EU's Horizon 2020 program shows there is great potential for crop residues as a feedstock for lignocellulosic biomass.

Part B: On Efficiency and Emissions

The efficiency and emissions of the ethyl valerate test fuel blend must also be examined to better judge its suitability in replacing gasoline. For all spark timings investigated, the thermal efficiency of the ethyl valerate test fuel blend was higher when compared to gasoline (see Figure 6). This is caused by the lower indicated specific fuel consumption using the test fuel blend at all spark timing when compared to gasoline (not including test 4). This result is slightly unexpected given the higher energy density of gasoline, meaning the ISFC of gasoline should be lower.

The reason for this is likely to be the narrow range of spark timings investigated. As previously mentioned, experimental data has shown 31 CAD BTDC to be the optimal spark timing for the combustion of gasoline in a spark ignition engine. If the range of spark timings investigated were to be increased, we would likely see a range of spark timings across which the ISFC of gasoline was higher than the test fuel blend, leading to higher thermal efficiencies.

For all spark timings investigated, the emissions of both CO_2 and NO_x were lower for the test fuel blend when compared to gasoline. Additionally, the emissions of both gases were lower for earlier spark timings when using the test fuel blend; whilst the opposite was true for gasoline, emissions were higher for earlier spark timings.

The difference in the amount of CO_2 produced can be explained by the chemical composition of the fuels. Since gasoline, whose main component is octane ($\mathrm{C}_8\mathrm{H}_{18}$), is more carbon heavy than ethyl valerate ($\mathrm{C}_7\mathrm{H}_{14}\mathrm{O}_2$), thus blending ethyl valerate with gasoline will reduce the carbon dioxide produced during combustion. Similarly, since more air is required for the combustion of air with gasoline than the test fuel blend, more NO_x is produced since the formation of it is highly oxygen dependent.

Interestingly, temperature and fuel viscosity dependence seem to have little effect on the NO_x emissions. Theoretically, NO_x emissions increase with both increasing combustion temperature and fuel

viscosity. Table 4 shows that for all spark timings, the exhaust gasses are hotter for the test blend when compared to gasoline, which suggest combustion temperatures are higher. Furthermore, the test blend had a much higher viscosity of 1.08 mPas compared to 0.61 mPas for gasoline.

Spark Timing	Average Exhaust Ga	s Temperature (Celsius)
	10% Ethyl Valerate	Reference Gasoline
10 CAD BTDC	477.7	461.1
13 CAD BTDC	492.7	481.0
16 CAD BTDC	493.3	484.3

Table 4: Average exhaust gas temperature for all spark timings.

Both factors suggest that $\mathrm{NO_x}$ emissions should be higher for the ethyl valerate test blend, though we do not observe this in our data. Again, it is important to note the narrow range of spark timings across which our results were collected. Plotting $\mathrm{NO_x}$ emissions against spark timing (see FIGURE X) suggest that $\mathrm{NO_x}$ emissions for ethyl valerate will increase beyond the $\mathrm{NO_x}$ emissions for gasoline, which looks to have reached a maximum near 10 CAD BTDC. However, these trends are by no means conclusive and must be tested experimentally.

In conclusion, ethyl valerate performed better than gasoline at all spark timings both in terms of thermal efficiency and pollutant emissions. However, due to the narrow range of spark timings investigated, the test fuel blend cannot conclusively be recommended as a potential replacement for gasoline. At different spark timings, gasoline may outperform the test fuel blend. The analysis would also benefit from data about additional pollutants such as unburnt fuel, carbon monoxide, and particulates. Furthermore, efficiency and emissions are not the only parameters to be examined. For example, for power sensitive applications such as high-performance vehicles, IMEP is a much more important parameter to consider. Overall, the performance the ethyl valerate test fuel blend shows promise, though much more data would be needed to make a definitive judgement.

Results and discussion

Conclusions

References