Comparison of genotoxicity of exhaust from a diesel, biodiesel and rapeseed oil powered engine – pilot study

J. Topinka¹, A. Milcova¹, J. Schmuczerova¹, M. Mazac², M. Pechout², M. Vojtisek-Lom²

¹Department of Genetic Ecotoxicology, Institute of Experimental Medicine AS CR, Prague, 142 20, Czech Republic

² Department of Vehicles and Engines, Technical University of Liberec, Liberec, 461 17, Czech Republic

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Presenting author email: jtopinka@biomed.cas.cz

Introduction

Last decades are characterized by massive use of alternative fuels, including biofuels. Since the reports on the toxic effects of exhaust from engines powered by biofuels are often contradictory, it might be of great interest to compare genotoxicity of standard diesel particulate emissions with that of the most frequently used biofuels. For this purpose we performed the pilot study with the aim to identify possible genotoxicity induced by organic extracts from the samples of exhaust of engines running on diesel fuel, biodiesel (neat methylester of rapeseed oil) and neat heated, fuel-grade rapeseed oil. The engines were tested in a laboratory using engine dynamometers.

Methods and Results

In one set of tests, a Zetor tractor engine with an inline mechanical injection pump and no exhaust gas aftertreatment device was tested using the NRSC cycle (also the ISO-8178 test with C-1 weighing, normally used for certification of non-road engines) and the ISO-8178 test with C-2 weighings, representing lowload operation. A sample of undiluted exhaust was drawn through a cartridge with a fluorocarbon-coated filter and two polyurethane foam plugs, with 2.0-3.5 m³ of exhaust sampled. As a marker of the genotoxic potential, DNA adduct levels induced by extractable organic matter (EOMs) in an acellular assay of calf thymus DNA coupled with ³²P-postlabeling in the presence and absence of microsomal S9 fraction (contains enzymes for metabolic activation of genotoxic compounds such as PAHs) were employed. Simultaneously, chemical analysis of 16 priority PAHs in EOMs, including B[a]P was performed. The results suggest that on ISO-8178 non-road engine test cycle, C-2 schedule, representing low engine loads, 100 µg/ml of the organic extract from standard diesel particulate emissions induces highest DNA adduct levels (10.5 adducts/ 10^8 nucleotides), while rapeseed oil and methyl esters of rapeseed oil induce 3.2 and 0.5 adducts/10⁸ nucleotides, respectively. These results correlate with the content of carcinogenic PAHs and B[a]P in the corresponding EOMs.

In a second set of tests, the exhaust was routed to the laboratory main exhaust duct, which has served as an improvised full-flow dilution tunnel, with dilution ratio of approximately 1:100 at idle to 1:15 at full load. From this duct, diluted exhaust was sampled with highvolume samplers (Digitel) on the Teflon coated filters (Pallflex) normally used for ambient air quality measurements, at rates 500-1000 litres per minute, with a target accumulation on the order of 10 mg of particulate mass. Two engines were tested. One was a Cummins ISBe4 engine with a Common Rail fuel injection system and no exhaust gas aftertreatment device, tested using the World Harmonized Stationary Cycle (WHSC) and modified Engine Stationary Cycle (ESC). The ESC cycle was modified by altering the length of each of the 13 modes and including transitions between modes to facilitate continuous sampling. The other engine was the Zetor engine described above, which was tested using the NRSC cycle. Filters were extracted by dichlormethane and genotoxicity of extracts was analyzed by ³²Ppostlabelling of DNA adducts by test described in the previous paragraph. Major results are described in Table 1.

Table1. Genotoxicity of the organic extracts from particulate emissions of selected fuels

particulate emissions of selected fuels					
Engine	Fuel	PM	B[a]P	DNA add.	
fuel	/cycle	mg/kWh	ng/kWh	/kWh*	
injection	-	-	-	+S9	-S9
Cummins	D ¹ /WHSC	6.9	3.5	217	96
ISBe4	R ² /WHSC	7.2	4.9	159	13
Common	D ¹ /ESC	14.1	<2.5	541	140
Rail	R ² /ESC	23.8	11.1	378	145
	B ³ /ESC	20.2	7.3	433	145
Zetor	D ¹ /NRSC	185	< 0.37	2932	828
1505	R ² /NRSC	202	1.4	2351	874
*					

^{*}DNA adducts/10⁸ normal nucleotides/kWh; ¹D-diesel; ²R-rapeseed oil; ³B-biodiesel B-100

Conclusions

1. The emissions of classic diesel contain more of total PAHs, but much less B[a]P and other carcinogenic PAHs.

2. Genotoxicity of particulate emissions of selected biofuels is comparable with a classic diesel.

3. Metabolic activation (+S9) resulted in several fold higher genotoxicity suggesting major contribution of PAHs to the DNA adduct levels. However, directly acting genotoxicants (-S9) are also significant.

4. Genotoxicity is highly dependent on the test cycle (ESC vs. WHSC).

5. Genotoxicity of the emissions is dose/dependent (data not shown).

These results should be taken as preliminary and more detailed study is going on to verify these preliminary findings.

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