Estimated atmospheric emissions from biodiesel and characterization of pollutants in the metropolitan area of Porto Alegre-RS

ELBA C. TEIXEIRA, CAMILA D.P. MATTIUZI, SABRINA FELTES, FLAVIO WIEGAND and EDUARDO R.R. SANTANA

Fundação Estadual de Proteção Ambiental Henrique Luís Roessler, Rua Carlos Chagas, 55/802 90030-020 Porto Alegre, RS, Brasil

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ABSTRACT

The purpose of the present study was to estimate emissions of some pollutants (CO, NOX, HC, SOX, and PM) in diesel fleet due to the addition of biodiesel in different blends, as well as to assess atmospheric pollutant concentrations in the metropolitan area of Porto Alegre (MAPA). The methodology was based on inventories from mobile sources based on US EPA's technical report. Regarding air quality the following parameters were determined: PM_{10}, PM_{2.5}, CO, NOX, O_3, SO_2, HC and PAHs. The results showed a decrease for emissions PM, CO, and HC, and a slight increase for NOX. The characterization of atmospheric pollutants in the metropolitan area of Porto Alegre showed that they are influenced by mobile sources, particularly diesel vehicles. The diagnosis of ratios analysis that was applied to facilitate the identification of sources of PAHs, indicated an influence of diesel oil.

Key words: air quality, biodiesel, diesel, emissions.

INTRODUCTION

The worldwide expansion of the number of vehicles on the roads has been responsible for the increase in pollutant emissions into the atmosphere, thus frequently contributing to worsening air quality, particularly in large urban areas.

Vehicular emissions are one of the major primary sources of fine particles emission, and they might be responsible for approximately 15-50% of the total aerosol mass in urban areas (Sheesley et al. 2007 and references therein). Emissions from diesel engines are chemically complex and contain hundreds of chemical substances divided into gaseous and particulates phases. These emissions contain more particles than those from gasoline engines, with polycyclic aromatic hydrocarbons (PAHs) being their major organic compound (Desantes et al. 2005, Maricq 2007, Phuleria et al. 2007, Ravindra et al. 2008).

Particle size from diesel engines ranges between 20-130 nm, and between 20-60 nm when derived from gasoline engines (Harris and Maricq 2001, Ristovski et al. 2006).

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The use of alternative fuels, such as biodiesel, is being widely discussed in several countries due to an increased environmental awareness and higher petrodiesel prices. Currently, many countries use biodiesel and petrodiesel blends in their cars, e.g. USA, Japan, Brazil, India, and others.

In Brazil, to comply with law 11.097/2005, the use of B2 blend, i.e. 2% biodiesel in petroleum diesel, is mandatory as of January 2008. In January 2010, this mandated level increased to 5% (B5). There is a further possibility of enforcing blends with higher levels or even 100% biodiesel (B100), pending authorization of Brazil's National Oil Agency (ANP).

The present study aims to estimate vehicle emissions of the pollutants CO, NOX, HC, SOX, and PM from the addition of biodiesel to diesel in several blends, as well as to assess atmospheric pollutant concentrations in the metropolitan area of Porto Alegre (MAPA), located in the state of Rio Grande do Sul, Brazil.

**AREA OF STUDY**

The area chosen for this study was the metropolitan area of Porto Alegre (MAPA), at 29°30’S – 30°30’S / 50°25’W – 51°55’W, in the east part of Rio Grande do Sul, located in the south of Brazil. With a total area of 9,800 km², representing 3.76% of the total area of the state, it includes 31 counties, which are the most urbanized area of Rio Grande do Sul. The counties comprising the MAPA are: Alvorada, Araricá, Arroio dos Ratos, Cachoeirinha, Campo Bom, Canoas, Capela de Santana, Charqueadas, Dois Irmãos, Eldorado do Sul, Estância Velha, Esteio, Glorinha, Gravataí, Guaíba, Ivoti, Mon-tenegro, Nova Hartz, Nova Santa Rita, Novo Hamburgo, Parobé, Portão, Porto Alegre, Santo Antônio da Patrulha, São Jerônimo, São Leopoldo, Sapiranga, Sapucaia do Sul, Taquara, Triunfo and Viamão.

The area is characterized by different industrial typologies, including several stationary sources such as the Alberto Pasqualini oil refinery, two steel mills (Siderúrgica Riograndense and Açôs Finos Piratini), the petrochemical industrial complex, and two coal-fired power plants (Termochar and São Jerônimo). Despite the different industrial sources around Porto Alegre contributing to the total emissions, the major contributions come from an estimated 620,000 vehicles on the roads, representing 20% of the total 3.1 million vehicles of the state (Teixeira et al. 2008).

Due to its location, winter in the MAPA is strongly influenced by cold air masses migrating from polar regions. The seasons are well defined and the rain is evenly distributed all over the year. According to Köppen’s international system of climate classification, the area under study is a type Cfa, i.e. a humid subtropical climate with well-distributed rains all over the year and an average temperature above 22.0°C during the warmest month (Livi 1999). The prevailing wind directions are east (E), east southeast (ESE), and southeast (SE) (Livi 1999). During the day, wind reaches its lowest speed at dawn and early morning, and highest speed in the late afternoon, between 5-7 p.m. This pattern is related to energy availability at the surface (sensible heat) during the day, intensifying local and mesoscale atmospheric circulations. The prevailing wind results from interactions of mesoscale phenomena, especially sea/land breezes (from the Atlantic Ocean and the Patos Lagoon) and valley/mountain breezes (from the nearby Serra Geral mountains to the north of the MAPA).

**METHODS**

**EMISSION ESTIMATION**

The methodology used to estimate the atmospheric emissions in the MAPA was based on the United States Environmental Protection Agency’s methods.
for vehicular emissions inventories (US EPA 2002). Atmospheric emission scenarios with biodiesel added to diesel in different blends - 0% (D100), 2% (B2), 5% (B5), 10% (B10) and 20% (B20) - were applied to the methodology for 2008 and 2010. This allowed forecasts of pollutant emissions, quantified for carbon monoxide (CO), hydrocarbons (HC), sulfur oxides (SO\(_x\)), nitrogen oxides (NO\(_x\)), and particulate matter (PM) emissions.

The working method of the present study consisted of two steps:

1. Data collecting and organization
2. Intermediate calculations to generate the variables for the general equation (1).

\[
E_{\text{pollutant, year}} = \sum_{\text{year-model}} (Fe \times N \times C \times A \times 10^{-6}) \quad (1)
\]

\(E\) = pollutant emission rate (ton/year)

\(Fe\) = pollutant emission factor (g/km)

\(N\) = number of vehicles

\(C\) = fuel consumption (L/year)

\(A\) = vehicle autonomy (km/L)

All steps of calculations were developed by using Microsoft Office Excel software. Initially, the diesel fleet was subdivided into two classes: trucks, vans and pickups, and buses and minibuses, with their corresponding values for kilometers driven and autonomy.

The first step consisted of collecting data about the fleet and fuel consumption of the 31 counties of the MAPA region, which was provided by the Traffic Department of Rio Grande do Sul (DETRAN/RS) and the Brazilian Oil Agency (ANP), respectively.

The second step consisted of calculating the fuel consumption and the emission factor based on data provided by ANP and the Brazilian Institute of Geography and Statistics (IBGE), and by the São Paulo State Environmental Company (CETESB) and US EPA, respectively.

As it is not possible to affirm that a vehicle refueling in a given county will circulate only within that area, it may exist a significant discrepancy between the volume of fuel given by ANP and the fuel volume used in the county. In order to obtain a more realistic fuel consumption profile of that area, a weighting was done with respect to the urban and rural population of each county using statistical data from IBGE.

Emission factors are applied to calculate the estimated atmospheric emissions. In this study emission factors were provided by CETESB (CETESB 2008). These values are obtained in function of average speed, air temperature, and vehicle technology, amongst other variables. According to the Brazilian regulations, CETESB performs annual calculations of emission factors, which are then used as reference for all studies conducted in Brazil in this field. In this study the emissions factors were adjusted with the numbers of diesel vehicles on the roads in 2008 and 2010 and their fuel consumption.

In the present study, the exponential correlation referring to the emission of biodiesel was taken into account (Figure 1). Calculation adjustment to biodiesel was done by applying equation 2 added of the values of coefficient "a" (Table I) and, by considering variables "b" to be close to null, transforming the second term in a unit value. This equation considers the percentage of the biodiesel blend, and it was obtained on US EPA’s technical reports. (US EPA 2002).
\[ E_{\text{pollutant, year}} = \exp[a \times (\text{vol\% biodiesel})] \times \exp[b] \quad (2) \]

**TABLE I**

<table>
<thead>
<tr>
<th>Coefficient “a”</th>
</tr>
</thead>
<tbody>
<tr>
<td>NO\textsubscript{X}</td>
</tr>
<tr>
<td>PM</td>
</tr>
<tr>
<td>HC</td>
</tr>
<tr>
<td>CO</td>
</tr>
</tbody>
</table>

(US EPA 2002).

After these two steps all the variables of the equation 1 were obtained. Thus, we were able to estimate the emissions of NO\textsubscript{X}, CO, PM and HC in the metropolitan area of Porto Alegre for 2008 and 2010 with different blends of biodiesel.

Figure 2 shows the flow chart based on the calculation methodology of atmospheric emissions and on adjustments of the available variables to equation 2.

**Automatic Samplers**

The equipment used for sampling included a sulfur oxide analyzer (using UV fluorescence method), a PM\textsubscript{10} analyzer (using beta radiation method), a nitrogen oxide analyzer (AC31M using chemiluminescence method), a carbon monoxide analyzer (CO11M using infrared absorption), an ozone analyzer (O341M, absorption of UV light with a wavelength of 254 nm, LCD/UV photometry ozone analyzer), and a total hydrocarbon analyzer (HC 51 M FID Total hydrocarbon analyzer model). All equipments are manufactured by Environnement S.A.

**Stacked Filter Units (SFU) - PAHs**

The experimental method of atmospheric particles (PAHs) consisted of four stages: 1) sampling - sampler (SFU) (Hopke et al. 1997) PM\textsubscript{2.5} – PM\textsubscript{2.5-10}; flow 16.7 L min\textsuperscript{-1}, and Whatman quartz fiber filters; 2) extraction and 3) cleanup were performed according to US EPA TO 13 A norm (US EPA 1999); 4) analysis; the 16 most important PAHs (US EPA 1999) were analyzed by chromatography coupled with a Shimadzu GCMS-QP5050A mass spectrophotometer detector. For further details about the analyses, see Dallarosa et al. (2005, 2008).

**RESULTS AND DISCUSSION**

**EMISSIONS**

Figure 3 shows the distribution of diesel vehicles on the roads in 2008 and 2010 in the MAPA. In this area there where 80,730 and 98,576 diesel vehicles in 2008 and 2010, respectively. An increase of 22% in the total diesel fleet can be seen during these years. This increase was already expected due to the raise in vehicle sales in the state of Rio Grande do Sul during 2009 and 2010. Among the 31 counties of the MAPA, Porto Alegre is the major contributor to atmospheric emissions, since it represents approximately 40% of the total diesel fleet in the area. In 2010, diesel consumption in the MAPA was...
836,838 m³ compared to 736,611 m³ in 2008, and in Porto Alegre it was 305,396 m³ and 268,819 m³ in 2010 and 2008, respectively.

Table II shows the estimated atmospheric emissions for different biodiesel amounts added to diesel: 2, 5, 10, and 20%. The data refer to the MAPA and Porto Alegre in 2008 and 2010. Porto Alegre, as expected, represents about 36-40% of the emissions. It can be observed that PM, CO, and HC emissions decreased, while NOX shows a slight increase. The emissions in the year 2010 are higher in the MAPA and PA when compared to the emissions in 2008; this can be explained by the growth of the diesel fleet circulating in the MAPA during 2009 and 2010, as mentioned previously.

The variation of atmospheric emissions of each type of pollutant for different biodiesel blends added to diesel for 2008 and 2010 are shown in Figure 4 and 5, respectively. As the methodology remains the same, the variation in percentage is the same in both years (2008 and 2010). The changes can be seen in the amount of mass of pollutant that is emitted: with an increase in the vehicular fleet, the quantity mass emitted will raise as well, as seen in Table II.

In both Figures 4 and 5 it is observed an increase in NOX emissions with the addition of biodiesel. This might be explained due to high oxygen levels in biodiesel molecules and improved combustion conditions – temperature and pressure in the combustion chamber are higher, boosting the formation of larger amounts of nitrogen oxides in engines running on biodiesel, by combining oxygen with the nitrogen present in the air (Coronado et al. 2009).

**TABLE II**

<table>
<thead>
<tr>
<th></th>
<th>2008</th>
<th>2010</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MAPA</td>
<td>PA</td>
</tr>
<tr>
<td>NOX</td>
<td>18479</td>
<td>18516</td>
</tr>
<tr>
<td>CO</td>
<td>2550</td>
<td>2518</td>
</tr>
<tr>
<td>HC</td>
<td>870</td>
<td>851</td>
</tr>
<tr>
<td>PM</td>
<td>360</td>
<td>355</td>
</tr>
</tbody>
</table>

**Fig. 3** - Distribution of diesel vehicles on the roads in 2008 and 2010 in the MAPA.

**Fig. 4** - Variation of atmospheric emissions at different biodiesel blends added to diesel oil in 2008 in the MAPA.
Carbon monoxide emissions decreased 40-50% compared to common diesel due to the oxygen present in biodiesel molecules, allowing a more efficient combustion by moving the direction of the combustion reaction toward the production of CO₂ (Maziero et al. 2006).

The higher oxygen levels present in the biodiesel molecule allow the complete burning of the fuel. Consequently, there is a decrease in particulate matter emissions. The absence of sulfur in biodiesel contributes to less sulfate particles formed during combustion, thus contributing even more to a decrease in particulate matter emissions (Dwivedi et al. 2006).

The decrease in emissions of unburned hydrocarbons by the addition of biodiesel to the diesel results from the complete combustion of the fuel, during which CO₂ and water are mainly formed, as opposed to what occurs during diesel combustion (Coronado et al. 2009).

Although the variation of SOₓ emissions with the addition of biodiesel is not provided in the present study, these emissions decreased proportionally to the percent of diesel added to the blend. This occurs because biodiesel is a vegetable-derived fuel and it is free of sulfur (Coronado et al. 2009).

On the National Biodiesel Board (NBB) website there is a tool that calculates the variations in mass of the emissions of some pollutants, according to the addition of biodiesel in different blends (NBB 2010). Table III shows a comparison of the variations, in mass, of the emissions estimated in the present study and the results obtained on the NBB website for the parameters PM, CO, HC and NOₓ in Porto Alegre and the MAPA. It can be seen that data for HC and NOₓ show higher agreement, while there is a significant difference in PM and especially in CO emissions. This might occur because both compounds depend directly on combustion, and the variables used to consider NBB’s calculation were probably at different conditions as those used in the present study.

**TABLE III**
Comparison of the variations on estimated atmospheric emissions of blends of biodiesel in Porto Alegre and the MAPA, with NBB calculation chart.

<table>
<thead>
<tr>
<th>Ton/year</th>
<th>B2</th>
<th>B5</th>
<th>B10</th>
<th>B20</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Estimated</td>
<td>NBB</td>
<td>Estimated</td>
<td>NBB</td>
</tr>
<tr>
<td>NOX</td>
<td>MAPA</td>
<td>63</td>
<td>30</td>
<td>93</td>
</tr>
<tr>
<td></td>
<td>PA</td>
<td>14</td>
<td>11</td>
<td>34</td>
</tr>
<tr>
<td>CO</td>
<td>MAPA</td>
<td>-30</td>
<td>-150</td>
<td>-82</td>
</tr>
<tr>
<td></td>
<td>PA</td>
<td>-12</td>
<td>-55</td>
<td>-30</td>
</tr>
<tr>
<td></td>
<td>PA</td>
<td>-7</td>
<td>-7</td>
<td>-18</td>
</tr>
<tr>
<td>PM</td>
<td>MAPA</td>
<td>-5</td>
<td>-14</td>
<td>-12</td>
</tr>
<tr>
<td></td>
<td>PA</td>
<td>-2</td>
<td>-5</td>
<td>-5</td>
</tr>
</tbody>
</table>

*B2 - blend of 2% biodiesel; B5 – blend of 5% biodiesel; B10 – blend of 10% biodiesel; B20 – blend of 20% biodiesel.*
The results of several authors on pollutant emissions from diesel oil added of different biodiesel blends are show in Table IV. The results are expressed in percentage variation of the emissions of NOX, CO, HC, PM and SOX. Results reported by Morris et al. (2003) and by the National Biodiesel Board (NBB 2010) showed higher agreement with the estimated emission values for B20 in the present study. At lower percents of biodiesel in diesel, results did not show significant variations, with only slight variations in the values found. This might indicate a change of trend. The error margin of the estimated and measured values is sometimes greater than the absolute values found.

<table>
<thead>
<tr>
<th>TABLE IV</th>
<th>Compared results (%) of various studies on pollutant emissions from diesel oil added of different biodiesel rates.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>B2</td>
</tr>
<tr>
<td>Pollutant</td>
<td>CO</td>
</tr>
<tr>
<td>MAPA (2010)</td>
<td>-1.3</td>
</tr>
<tr>
<td>Corrêa (2005)</td>
<td>-0.1</td>
</tr>
<tr>
<td>Oliveira and Costa (2002)</td>
<td>-</td>
</tr>
<tr>
<td>B10</td>
<td>B20</td>
</tr>
<tr>
<td>Pollutant</td>
<td>CO</td>
</tr>
<tr>
<td>MAPA (2010)</td>
<td>-6.2</td>
</tr>
<tr>
<td>Corrêa (2005)</td>
<td>-0.3</td>
</tr>
<tr>
<td>Oliveira and Costa (2002)</td>
<td>-</td>
</tr>
</tbody>
</table>

B2 - blend of 2% biodiesel; B5 – blend of 5% biodiesel; B10 – blend of 10% biodiesel; B20 – blend of 20% biodiesel.

AIR QUALITY

It is important to note that data will be here presented from 2006, 2007 and 2008. Table V shows the total average concentrations of PM$_{10}$, PM$_{2.5}$, CO, NO$_X$, O$_3$, SO$_2$, HC and PAHs measured in the MAPA during 2006, 2007 and 2008. There was a decrease in CO in the atmosphere in 2008. This compound is originated mostly from mobile sources that operate on the Otto cycle. This gasoline fleet is responsible for the emission of 160,000,000 ton/year of CO in the MAPA (Teixeira et al. 2010). Despite the greater number of vehicles on the roads in recent years, the addition of catalytic converters to automobiles has substantially reduced CO emissions from the combustion of common gasoline. However, CO emissions during the “cold start” of engines, when the catalyst is cold and has not reached operating temperature, can still be substantial, even for those vehicles that are equipped with catalytic converters (Gaffney and Marley 2009). Higher biodiesel blends foreseen to be added to diesel will allow a decrease in emissions and concomitantly a decrease of this compound in the atmosphere (Figure 5). Studies about macro-regions of Rio Grande do Sul (Teixeira et al. 2010) showed that the MAPA had higher CO concentration in the air due to gasoline-fueled vehicles. However, even if the annual mean concentration of CO is decreasing, there are concerns about an increase in the number of vehicles on the roads and, consequently, in fuel consumption (Teixeira et al. 2010).

SO$_2$ has shown an increase in 2006 and 2007, with a slight decrease in 2008. Despite vehicles have been showing to be major contributors to SO$_2$ emissions in the MAPA, some stationary sources located in the area, e.g. an oil refinery and coal-fired power plants, are probably also related to SO$_2$ emissions and contribute to higher emission levels measured in 2007. Some authors (Geng et al. 2009) have reported SO$_2$ in urban areas from vehicles and coal-fired power plants. These activities might speed up SO$_2$ concentration in the atmosphere (Nguyen and Kim 2006).
Table V shows NOx levels in 2006, 2007 and 2008. A slight decrease in NOx concentration from vehicular emissions from 2006 to 2008 can be observed. In recent years, despite the higher number of vehicles on the roads, there has been an improvement in fuel and lubricant oil quality, vehicle maintenance and operating conditions. Not all diesel vehicles had electronic injection until 2008, therefore they were probably contributing to a higher concentration of NOx in the atmosphere. Diesel vehicles on the roads are a major source of NOx. Studies (Gaffney and Marley 2009) have estimated that diesel engines produce five times the amount of NOx per mass of fuel burned when compared to gasoline vehicles.

Exhaust gases from motor vehicles can contribute significantly to atmospheric particles emissions, mainly those of an aerodynamic diameter of less than 10 µm (Künzli et al. 2000). In urban areas, traffic is one of the major contributors to atmospheric particles such as PM2.5 (Heinrich et al. 2005), but mainly the PM1.0 fractions (Gaffney and Marley 2009). Table V shows PM10 and PM2.5 average concentrations in the MAPA from 2006 to 2008. The area studied herein (MAPA) has a heavy traffic of heavy diesel trucks, and just in Porto Alegre diesel consumption was 259,181 m³ and 268,819 m³ in 2007 and 2008, respectively. The heavy traffic of diesel engines significantly contributes to produce ultrafine particles (Vallius et al. 2000) and more carbonaceous particulate matter (Gaffney and Marley 2009). These ultrafine particles have a very large surface area onto which the organic contaminants present in the diesel exhaust can adsorb. The emission depends on a number of factors such as vehicle age, engine design and operating conditions, lubricant oil and fuel quality, as well as environmental conditions (Maricq 2007, Yanowitz et al. 2000). It is believed that, with the higher percent of biodiesel added to diesel, there will be a significant decrease in ultrafine particles emissions and, consequently, the quality of air will improve (Zhang et al. 2011).

These organic contaminants emitted by vehicles, especially by diesel engines, can be polycyclic aromatic hydrocarbons. Some of these compounds have shown carcinogenic and mutagenic properties (Chang et al. 2006, Kawanaka et al. 2004, Villalobos-Pietrini et al. 2007). Studies on PAHs in fine particles conducted by several authors (Dallarosa et al. 2008, Bourotte et al. 2005, Hays et al. 2003, Sheu and Lee 1997) have shown that the highest concentration of these compounds is found in fine particles. Along with their small size, these particles have a very large surface area onto which other organic contaminants present in the diesel exhaust can adsorb. Polycyclic organic matter compounds with more than three aromatic rings are usually associated with diesel particulates.

Table V shows the total average concentration of PAHs associated with atmospheric particles in the MAPA. Between 2006 and 2008 a small decrease was observed in concentration of PAHs. As mentioned before, during this period not all diesel vehicles had electronic injection, therefore a contribution in order to reduce the concentration of PAHs in the atmosphere did not exist. In the MAPA the alcohol fleet corresponds to only 4% (Teixeira et al. 2010),
and besides that, according to some authors (Allen et al. 2008), the emission of PAHs, from the use of hydrated alcohol, is almost negligible compared to gasoline. Other authors (Abrantes et al. 2009) reported that the emission of PAHs ethanol from vehicles were on average 92% lower than PAHs emission from the gasohol vehicle.

Figure 6 shows the seasonal average concentration of PAHs in the MAPA. A higher concentration of these compounds can be observed in fine particles on the winter. Many factors influence the concentration of PAHs in the atmosphere. In this case the concentration of PAHs ranged in relation to particles size because the smaller particles have higher specific surface area and sorption capacity. Another factor observed was the seasonal variability of PAHs concentration that can be explained by chemical and photochemical degradation due to differences in sunlight intensity, temperature, and washouts effects. In the MAPA, which has a strong influence of traffic from heavy-duty gasoline and diesel vehicles, there is an increase in the emissions from primary sources of combustion during the winter (e.g., motor vehicles) and a decrease in photochemical oxidation of PAHs by solar radiation. Consequently, those conditions favor PAHs accumulation in the atmospheric particles (Di Filippo et al. 2010). These data are similar to those already reported in previous studies, e.g. Lee (2001), Bae et al. (2002), and Fang et al. (2006), who reported higher PAHs concentrations in the coldest periods.

Among the 31 counties of the MAPA, the three sites (Canoas, Sapucaia do Sul and FIERGS/Porto Alegre) are the major contributors to atmospheric emissions (approx. 48.0%) due to the great number of diesel vehicles with NOX emissions of approximately 7,767 tons each year. In 2007 and 2008, diesel consumption in these areas was 343,334 m³ and 356,102 m³, respectively (Teixeira et al. 2010). As described above, in this region the alcohol fleet corresponds to only 4%, and besides that, according to some authors (Allen et al. 2008, Abrantes et al. 2009), the emission of PAHs from the use of hydrated alcohol is very low and almost negligible when compared to gasoline emissions.

In order to characterize the major sources of PAH emission, the study of ratios was applied (Tsapakis et al. 2002, Vasconcellos et al. 2003). The ratios were calculated in order to facilitate the identification of sources of anthropogenic emissions, especially vehicular emissions by means of the BaA/BaP; Bgp/ID; Pyr/BaP; BaP/(BaP+Chry; Fluo/(Fluo + Pi; ID/(ID +BP) ratios.

The characteristic ratios of diesel oil and gasoline BaA/BaP; Bgp/ID; Pyr/BaP; BaA/(BaA+Chry) used for characterizing emissions from these sources allowed assessing the influence of diesel oil in the MAPA (Table VI). The BaA/(BaA+Chry) ratio, whose value of 0.4 was obtained in the present study, lies within the emission range for heavy diesel engines (0.38-0.65) (Kavouras et al. 2001). It was observed that even when the value nears its lowest limit, it still shows an influence of diesel in the MAPA.

Another ratio, ID/(BghiP+ID), also showed to be a good indicator for diesel engine emissions (0.35-0.70) (Kalaitzoglou et al. 2004). The average ratios obtained for the studied years were 0.415 and 0.429, leading to the conclusion that vehicles with catalytic converters are not enough to reduce the impact of some pollutants on the environment. Emissions of fuel pollutants still represent a significant burden. Other studies reached the same conclusion (Fang et al. 2004, Manoli et al. 2004, Guo et al. 2003).
Table VII shows the correlation matrix of the studied pollutants. PM$_{10}$ correlates with CO and CO correlates with NO$_X$. These results evidence the influence of vehicular emissions in the MAPA. This table shows that NO$_X$ and O$_3$ have a significant negative correlation. The negative correlation of NO$_X$ with O$_3$ confirms the so-called ‘titration effect’ of NO on O$_3$; NO reacts with O$_3$ to produce NO$_2$ (NO + O$_3$ → NO$_2$ + O$_2$). The correlation of NO$_X$ with PAHs and the negative correlation with O$_3$ evidence the influence of mobile sources, particularly emissions from diesel engines in the area of study.

**CONCLUSION**

The major benefit of biodiesel to the environment is the significant decrease in atmospheric pollutant emissions compared to petroleum diesel oil. Estimated data on biodiesel addition to diesel oil indicate a decrease in pollutant levels, particularly for B20. Pollutant emissions showed a decrease in PM, CO, and HC, while NO$_X$ experienced a slight increase.

Vehicle emissions were usually the most significant source of air pollution in the MAPA. This was intensified by emissions from diesel engines.

Table VI

<table>
<thead>
<tr>
<th>Year</th>
<th>Fluo/ (Fluo + Py)</th>
<th>ID/(ID + BP)</th>
<th>BaA/ BaP</th>
<th>BaP/ (BaP + CRI)</th>
<th>BGP/ ID</th>
<th>Pyr/ BaP</th>
</tr>
</thead>
<tbody>
<tr>
<td>2006</td>
<td>0.629</td>
<td>0.415</td>
<td>1.28</td>
<td>0.480</td>
<td>2.37</td>
<td>1.11</td>
</tr>
<tr>
<td>2008</td>
<td>0.514</td>
<td>0.429</td>
<td>0.95</td>
<td>0.495</td>
<td>1.88</td>
<td>2.02</td>
</tr>
</tbody>
</table>

The characterization of atmospheric pollutants in the MAPA showed that they are influenced by mobile sources, particularly diesel engines.

Even with more vehicles on the roads in recent years, the average concentration of the atmospheric pollutants (PM$_{10}$, PM$_{2.5}$, CO, NO$_X$, O$_3$, SO$_2$, HC and PAHs) from vehicles, particularly those with diesel engines, decreased in 2006, 2007 and 2008 in the MAPA. This is confirmed by the correlation of PM$_{10}$ with CO, and of NO$_X$ with CO and PAHs.

There is evidence that biodiesel would increase the NO$_X$ emissions, but the measures of the concentrations of air quality indicated little difference and a decrease for NO$_X$. There is the need to introduce more studies in the future, when a higher content of biodiesel can confirm this trend or not.

**ACKNOWLEDGMENTS**

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**RESUMO**

O objetivo do presente estudo foi estimar as emissões de alguns poluentes (CO, NO$_X$, HC, SO$_2$ e MP) na frota diesel devido à adição de biodiesel em diferentes misturas, bem como avaliar as concentrações de poluentes atmosféricos na região metropolitana de Porto Alegre (MAPA). A metodologia foi fundamentada em inventários de fontes móveis baseados em relatórios técnicos da US EPA. Em relação à qualidade do ar foram determinados os seguintes parâmetros: PM$_{10}$, MP$_{2.5}$, CO, NO$_X$, O$_3$, SO$_2$, HC e HPAs. Os resultados mostraram uma diminuição das emissões de PM, CO e HC e um ligeiro aumento de NO$_X$. A caracterização dos poluentes atmosféricos na região metropolitana de Porto Alegre mostrou que eles são influenciados por fontes móveis, principalmente, por veículos a diesel.
A análise de diagnóstico das razões, que foi aplicada para facilitar a identificação das fontes de HPAs, indicou influência de óleo diesel.

**Palavras-chave:** qualidade do ar, biodiesel, diesel, emissões.

**REFERENCES**


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