Biodiesel: An Overview

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The importance of biodiesel production was analyzed based on scientific articles and patents. A critical analysis was presented on the most used oil sources, the catalysts and the methods to verify the transesterification yields. Also analyzed were the comparative studies on emissions from pure fossil diesel and mixtures with biodiesel in variable proportions. Finally some challenges and considerations focused on technological, agronomic and infrastructure aspects of biodiesel production were indicated.

**Keywords:** biodiesel, emissions, transesterification, vegetable oils

1. Introduction

The predicted shortage of fossil fuel encouraged the search for substitutes for petroleum derivatives. This search resulted in an alternative fuel called “biodiesel”. The concept of biodiesel is still under discussion. Some definitions consider biodiesel as any mixture of vegetable oil and fossil diesel, while others take into account only mixtures of alkyl esters of vegetable oils or animal fats and diesel. The definition adopted by the Brazilian Biodiesel Programme is: “a fuel obtained from mixtures, in different proportions, of fossil diesel and alkyl esters of vegetable oils or animal fats”. Technically speaking, biodiesel is the alkyl ester of fatty acids, made by the transesterification of oils or fats, from plants or animals, with short chain alcohols such as methanol and ethanol. Glycerine is, consequently, a by-product from biodiesel production.

There are several choices for vegetable oil sources. In Brazil, soybean oil is a source that is already scaled up for biodiesel production. Nevertheless, other sources, such as sunflower, peanut, cotton, palm oil, coconut, babassu and, especially, castor oil, may be used in the near future, once their cultivation could achieve an economic up-scaling. The alcohol source in general is methanol. In Brazil, ethanol from sugar cane has a great potential as an alcohol source, since it may ally its non toxicity with a ready availability. But, it is less reactive than methanol and there are some technological problems in its industrial use.

The possibility of using vegetable oils as fuel has been recognized since the beginning of diesel engines. In 1911, Rudolph Diesel presented an engine based on compression-ignition: the diesel engine. At that time there was no specific fuel to feed this engine. Rudolph Diesel used groundnut oil. In the preface of the book *Thermodynamics*, (Chalkey, 1911),\textsuperscript{1} Rudolph Diesel wrote that the diesel engine could be fed by vegetable oils, and that this possibility could help the agricultural

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E-diesel is currently being used in fleet vehicles in the European Union and the United States. Studies carried out with E-diesel indicated significant reductions of particulate matter (PM), sometimes up to 40%, depending on the test methods and operating conditions. The CO and nitrogen oxides (NOx) emissions were significantly lower when a 20% blend of E-diesel was used in a constant-speed stationary diesel engine, as opposed to diesel fuel. The addition of ethanol to diesel may result in a volumetric reduction in sulphur, by as much as 20%, thus significantly reducing SO2 emissions. The major drawback in E-diesel is that ethanol is immiscible in diesel over a wide range of temperatures.

The viscosity of a mixture of sunflower oil and diesel (1:3 v/v) is 4.88 cSt at 40 °C (maximum specified ASTM value is 4.0 cSt at 40 °C). The mixture is not suitable for long term use in diesel engines. A blending of diesel and safflower oil (high level of oleic acid) was tested resulting in good performance, but its long term use is not recommendable because of lubricant thickening. A blending of soybean oil and solvent (1:1 v/v; solvent composition: 48% paraffins and 52% naphthalenes) has a viscosity of 5.12 cSt at 38 °C, and generate heavy carbon deposits and wearing on the engine upper ring.

Transesterification is widely used to reduce vegetable oil viscosity. Most industrial processes employ alkaline catalysis and methyl alcohol. In almost all countries methanol is more available than ethanol.

The transesterification process reduces the molecular weight to ca. 1/3 in relation to the triglycerides, and also, reduces the viscosity and improves the volatility. After the reaction, the products are a mixture of esters of fatty acids, glycerol, alcohol, catalyst and a low percentage of tri-, di- and monoglycerides. Pure biodiesel fuel (100% esters of fatty acids) is called B100. When blended with diesel fuel the designation indicates the amount of B100 in the blend, e.g. B20 is 20% B100 and 80% diesel, and B5 used in Europe is contains 5% B100 in diesel.

## 2. A General View About Research On Biodiesel

A search of scientific articles and patents on biodiesel was undertaken to get a general view about this issue. The search focused on the last 5 years (2000 to September 2004) and it used “biodiesel” as the keyword. The following databases were accessed:

**Scientific articles:** (i) Scielo (database developed by FAPESP, BIREME and CNPq) - www.scielo.br, (ii) American Chemical Society - www.pubs.acs.org, (iii) Elsevier – www.sciencedirect.com

134 scientific articles and 84 published and/or issued patents were found. The information shown in items 2 and 3 is based on these articles and patents.

The growing interest in biodiesel may be testified by the number of scientific articles and patents (Figure 1).

Consulting the Journal of Citation Reports (1997) it was found that 19.4% of articles were published in journals whose Impact Factor (IF) was not defined; 37.3% in journals whose IF was between 0 – 1; 33.6% in journals whose IF was between 1 – 2; and 9.7% in journals whose IF was up to 3.

The search for patents in Industrial Property Offices was focused on Brazilian, American and European Offices (Figure 2). At the Brazilian Office (INPI), 9 registers of patent were found since 2000, but only one issued patent. At the American Office (USPTO), 12 issued patents were found since 2000, and there are 14 published patents. The European Office shows all patents in the European Union and all patent documentation received from other Industrial Property Offices. All these data formed a group of 84 patents that, together with the articles, was used as reference for this topic and the next. It is important to say that among the patents there are INPADOC families.

A patent can be a tool to protect an interesting market. The number of patents at a specific Industrial Property Office reveals the power of this market concerning the technological area protected by the patent (Figure 2). Some patents are submitted on the PCT: Patent Cooperation Treaty. This treaty is signed by about 125 members’ states and allows to invoke the protection for an invention in the main countries of the world, by means of a single application.

Universities are the institutions that devote more attention to write scientific articles about biodiesel. Among the 134 scientific articles analyzed, 86 were from universities; 26 from universities in collaboration with research centres or industries, and 22 from research centres in collaboration with industries. On the other hand, among 84 patents, only three have universities as applicant.

2.1 Worldwide research on biodiesel in the last five years

USA has prominence in almost all technological and scientific areas. This scenery is the same concerning the production of patents and scientific articles on biodiesel (Figure 3). It is remarkable the effort made by Germany to reach a high scientific and technological level in biodiesel production. The German eminence in this topic is due, among other factors, to the absence of active petroleum deposits in this country and to the governmental support to new technologies based on renewable sources.

3. Source for Biodiesel Production

The source for biodiesel production is chosen according to the availability in each region or country. Any fatty acid source may be used to prepare biodiesel, but most scientific articles take soybean as a biodiesel source (Figure 4). Since the prices of edible vegetable oils, as soybean oil, are higher than that of diesel fuel, waste vegetable oils and non-edible crude vegetable oils are preferred as potential low priced biodiesel sources. Low-quality underused feedstocks have been used to produce biodiesel. This product is comparable in composition, similar in engine performance and emissions, and predicted to be more economical to produce than biodiesel from refined soybean oil.

There are various other biodiesel sources: almond, andiroba (Carapa guianensis), babassu (Orbignia sp.),
barley,\textsuperscript{18} camelina (\textit{CAMELINA SATIVA}),\textsuperscript{19} coconut,\textsuperscript{13,20} copra,\textsuperscript{21} cumaru (\textit{Dipteryx odorata}),\textsuperscript{15} \textit{Cynara cardunculus},\textsuperscript{22} fish oil,\textsuperscript{17,23} groundnut,\textsuperscript{13,16,17,21} \textit{Jatropha curcas},\textsuperscript{24,25} karanja (\textit{Pongamia glabra}),\textsuperscript{26} laurel,\textsuperscript{13} \textit{Lesquerella fendleri},\textsuperscript{27} \textit{Madhuca indica},\textsuperscript{28} microalgae (\textit{Chlorella vulgaris}),\textsuperscript{29} oat,\textsuperscript{18} piqui (\textit{Carvocar sp.}),\textsuperscript{15} poppy seed,\textsuperscript{13,30,32} rice,\textsuperscript{18,21,23} rubber seed,\textsuperscript{31,32} sesame,\textsuperscript{13} sorghum,\textsuperscript{18} tobacco seed,\textsuperscript{33} wheat.\textsuperscript{18}

From a chemical point of view, oils from different sources have different fatty acid compositions. The fatty acids are different in relation to the chain length, degree of unsaturation or presence of other chemical functions (Table 1). Esters prepared with long chain fatty acid or saturated fatty acid show higher cetane number, higher cloud point and cause nozzle clogging. Esters prepared with very unsaturated fatty acid show low cetane number and suffer oxidation easily. Generally, cetane number, heat of combustion, melting point and viscosity of neat fatty compounds increase with increasing chain length and decrease with increasing unsaturation.

Biodiesel properties are strongly influenced by the properties of the individual fatty esters. It therefore appears reasonable to enrich it with certain fatty esters with desirable properties in the fuel in order to improve the properties of the whole fuel. It may be possible in the future to improve the properties of biodiesel by means of genetic engineering of the parent oils, which could eventually lead to a fuel enriched with certain fatty acids, possibly oleic acid, that exhibits a combination of improved fuel properties.\textsuperscript{34}

A suitable source to produce biodiesel should not compete with other applications that reach higher prices, for example pharmaceutical raw material. But the demand for pharmaceutical raw material is lower than for fuel sources. The selection of a source for fuel production must take into consideration the characteristics of oil market (Figure 5).
As much as possible the biodiesel source should fulfil two requirements: low production costs and large production scale. Refined oils have high production costs, but low production scale; on the other side, seeds, algae and sewerage have low production costs and are more available than refined or recycled oils.

To elect a biodiesel source it is necessary also to take into consideration the oil percentage in the plant and the yield of oil per hectare (Figure 6). The palm kernel, for example, has a great potential as a biodiesel source, and has been used by several countries (Malaysia, for example).

In Brazil, biodiesel production has been adjusted to the available crop in each region (Figure 7). In the North, palm kernel and soybean are the most used sources; in the Northeast, castor bean, palm oil, palm kernel, babassu, soybean and cotton seed; in the Central-West, soybean, cotton seed, castor bean and sunflower seed; in the Southeast, soybean, castor bean, cotton seed and sunflower seed; in the South, sunflower seed, soybean, rapeseed and cotton seed.

Soybean occupies a special place among the oil plants that grow in Brazil. This leguminous plant is cultivated in many states of the country and is responsible for ca. 90% of the total of produced vegetable oils. Its participation in the market is mainly related to exportation of seeds in natura and it is an important source of exchange value to Brazil. However, if the exported commodities were refined oil and tart, surely more value could be associated to the exported products.

Soybean and castor seed, because of the large agriculture areas, and palm, especially because of the high level of oil, provide excellent options for expansion of vegetable oil production in Brazil. In the Northeast of Brazil, besides palm, sunflower, cotton and soybean, we could highlight castor seed, which can be cultivated in all states. Nevertheless, a castor seed has presented low productivity, between 600 and 1,000 kg/ha/year. Studies show that this yield could reach 3,000 kg/ha/year with genetic improvement and with possible introduction of irrigated cultivation. Palm oil (Elaeis guineensis) is an important alternative for the sustainable development of some regions of Brazil, including Amazônia.

Ferrari et al. presented a process for biodiesel production by the transesterification of neutral soybean oil and
anhydrous ethanol using NaOH as catalyst. Combinations of biodiesel and diesel in the proportions of 0, 5, 10, 20, 40, 60, 80 and 100% were tested as fuel in an energy generator. The average consumption and mixture performance were analyzed. The tests showed a reduction in diesel oil consumption when mixed with up to 20% of biodiesel.

4. Development of New Catalysts for Biodiesel Production

The preparation of biodiesel is generally carried out using vegetable oils or animal fats as a starting material. The transesterification reaction is preferred to the direct esterification of fatty acid because triglycerides are more available than free fatty acids. Biodiesel is produced by transesterification of the triglycerides with short chain alcohols in the presence of a suitable catalyst. The reaction is shown in Scheme 1. The stoichiometry requires 3 mol of alcohol and 1 mol of triglyceride to give 3 mol of fatty acid esters and 1 mol of glycerine. The overall process is a sequence of three consecutive reversible reactions where diglyceride and monoglyceride are intermediate products.

\[
\begin{align*}
\text{RCOO} - \text{CH}_2 & \quad + \quad 3 \text{ R'OH} \\
\text{RCOO} - \text{CH}_2 & \quad \underbrace{\text{catalyst}} \quad 3 \text{ RCOOR'} + \\
\text{RCOO} - \text{CH}_2 & \quad \text{CH}_2\text{OH} \\
\text{RCOO} - \text{CH}_2 & \quad \text{CH}_3\text{OH} \\
\end{align*}
\]

Scheme 1. Transesterification reaction of triglyceride.

The transesterification reaction can be catalyzed by both acid and alkaline catalysts, using a homogeneous or heterogeneous catalytic process. Sodium and potassium hydroxide are commonly used as industrial catalysts, since they are relatively cheap and also very active. On the other hand, their utilization in vegetable oil transesterification produces soaps by neutralizing the free fatty acid in the oil and by triglyceride saponification. The soap formation is an undesirable side-reaction, because it partially consumes the catalyst, decreases the biodiesel yield and complicates the separation and purification steps. The removal of these catalysts is technically difficult and brings extra cost to the final product. In addition, the difficulty for recycling and the generation of large waste amounts make the traditional catalysts less favourable.

More recently, there has been an increasing development of new catalysts for biodiesel production. According to Web
of Science Database, 320 articles were found using “biodiesel” as the keyword in the period from 2000 to September 2004. A total of 80 articles focused on biodiesel production. From these, 67 articles reported production by transesterification reaction and 12 articles by esterification. The research showed that enzyme is the most studied catalyst in the last years and few studies have been conducted without a catalyst. The types of catalyst used in the transesterification process is presented in Figure 8.

![Figure 8. Number of articles on transesterification catalysts.](image)

It could be observed that there are more articles about heterogeneous processes than homogeneous processes (Figure 9).

![Figure 9. Number of articles for processes studied.](image)

Finally, the number of articles, according to approaches on different types of heterogeneous catalysts are presented in Figure 10. It clearly shows the increasing use of enzymes as catalyst.

The uses of different available catalysts for the preparation of biodiesel are reviewed. Special attention is given to the recently developed catalysts. Firstly, we will focus on acid and alkaline catalysts for homogeneous processes (section 4.1). Then, new acid and alkaline catalysts will be mentioned for heterogeneous processes (section 4.2).

### 4.1 Catalysts for homogeneous processes

There are a number of reviews in the literature on alcoholyis of triglycerides using conventional homogeneous catalysts.2,16,37,38,41 The alkali catalyzed transesterification of triglycerides is fast and high conversions are easy to achieve at temperatures from 40 to 65 °C. In addition, alkaline catalysts are less corrosive than acidic compounds. The alkaline processes are generally catalyzed by sodium alkoxides, sodium or potassium hydroxides as well as sodium or potassium carbonates. The alkoxide catalysts are more expensive than metal hydroxides. They are more difficult to manipulate since they are very hygroscopic and have only the hydroxide ion as impurity. In this sense, they do not produce soap through triglyceride saponification. The soap formation can also be avoided by using an acid catalyst. Sulphuric, hydrochloric and organic-sulphuric acids are usually preferred. The acids catalyze the free fatty acid esterification to produce fatty acid esters, increasing the biodiesel yield. Nevertheless, the acid catalyzed transesterification is much slower than the alkali catalyzed reaction and also needs more extreme temperature and pressure conditions.

There are few articles about the development of new catalysts for homogeneous processes. In 1995, a study reported the activity and efficiency of guanidine and eight substituted cyclic and acyclic guanidines as catalysts for the transesterification of rapeseed oil.42 The best result showed that only 1 mol% of 1,5,7-triazabicyclo[4,4,0]dec-5-ene (TBD) produces more than 90% of methyl esters after 1 h.

In 2003, metal complexes of the type M(3-hydroxy-2-methyl-4-pyrone),H2O, where M = Sn, Zn, Pb and Hg, were used for soybean oil methanolysis under homogeneous conditions.43 The Sn and Zn complexes showed great activities for this reaction, achieving yields of up to 90 and 40%, respectively, in 3 h using molar ratio 400:100:1 (methanol:oil:catalyst), without emulsion formation. Recently, the same researchers studied the catalyst behaviour of these complexes in the transesterification of different triglyceride sources. The methanolyis of Brazilian oils such as andiroba, babassu, cumaru, palm, piqui and soybean oils were investigated. The catalytic system formed by the tin complex always presents higher conversion than the others.15
4.2 Catalysts for heterogeneous processes

Although transesterification using a conventional alkali catalyzed process gives high conversion levels of triglycerides to their corresponding methyl esters in short times, the reaction has several drawbacks: it is energy intensive; recovery of glycerine is difficult; the catalyst has to be removed from the product; alkaline waste-water requires treatment and free fatty acids and water interfere with the reaction. In order to minimize homogeneous process problems, attempts to use heterogeneous catalyst systems in alcoholysis of triglycerides have been made. These catalysts greatly simplify the post-treatment of the products (separation and purification). They can be easily separated from the system at the end of the reaction and could also be reused. Besides, the use of heterogeneous catalysts does not produce soaps through free fatty acid neutralization or triglyceride saponification. However, the performance is still unfavourable compared to the alkaline homogeneous catalysts.

A large number of heterogeneous catalysts have been reported in the literature, including enzymes, zeolites, clays, guanidines heterogenized on organic polymers, ion-exchange resins, oxides, etc. Some recent results will be summarized below.

The activity of guanidine-containing polymers and their homogeneous analogues was compared as catalysts in the transesterification of soybean oil with methanol. The methanol:oil molar ratio was three times higher in the reactions using the guanidine-containing polymers in order to assure low viscosity for the reaction mixture. The heterogeneous catalysts were used at 5 mol% in contrast to the 3 mol% used with the homogeneous catalyst. They are nearly as active as their homogeneous analogues and reached the same high conversions (>90%) after prolonged reaction time. In addition, the use of polystyrene-bound biguanides as recyclable catalysts was showed for the transesterification of several vegetable oils. Yields above 90% were obtained in less than 15 min. These catalysts were, by far, more reactive and more stable than the polystyrene-supported guanidines previously described.

Insoluble salts of amino acids were used as catalyst for the methanolysis of triglycerides. Some metal salts of amino acids such as those of copper, zinc, cadmium, nickel, lanthanum, cobalt, calcium, magnesium and iron were tested. Zinc-arginate catalyzed the methanolysis of palm lanthanum, cobalt, calcium, magnesium and iron were tested. Zinc-arginate catalyzed the methanolysis of palm lanthanum, cobalt, calcium, magnesium and iron were tested.

The methanolation of rapeseed oil was tested in the presence of cesium-exchanged NaX faujasites, mixed magnesium-aluminum oxides, magnesium oxide and barium hydroxide as catalysts for different methanol:oil ratios. Barium hydroxide was particularly effective for a methanol:oil molar ratio of 6:1, at reflux of methanol and after a reaction time of only 1 h. The oil conversion was about 80% with a nearly quantitative ester molar fraction. On the other hand, cesium-exchanged NaX faujasite and mixed magnesium-aluminum oxides required a long reaction time and a high methanol:oil molar ratio to achieve high yields in methyl esters. Recently, the transesterification of soybean oil with methanol in the presence of a series NaX faujasite zeolite, ETS-10 zeolite and metal catalysts were studied. A 6:1 molar ratio of alcohol and a 10% m/v of catalyst were used at 60, 120 and 150 °C. The ETS-10 catalysts provided higher conversions (>80%) than the zeolite-X type catalysts.

Na/NaOH/g-Al2O3 heterogeneous alkaline catalysts were applied for the transesterification of soybean oil with methanol using hexane as co-solvent. The best result (90%) was obtained after 2 h using a methanol:oil molar ratio of 9:1 at 60 °C. Monteiro and Cruz studied the transesterification of soybean oil using mixed oxides as catalysts. The reactions were carried out using a weight ratio of 4.5:6.0:0:3 (methanol:soybean oil:catalyst) at 70 °C for 8 h. The following catalysts were investigated: ZrO2-SiO2, KOH/ZrO2-SiO2, CoO-SiO2, MoO3-SiO2, Na2O-SiO2, La2O3(10%)-MCM-41, MgO(10%)-MCM-41, BaO(10%)-MCM-41, CaO and MgO. Best results were obtained with La2O3(10%)-MCM-41, Na2O-SiO2 and CaO, with conversions of 81%, 76% and 67%, respectively.

The use of commercial sulfonic ion-exchange resin was also reported for production of biodiesel. The methanolation of babassu and soybean oil was compared using Amberlyst-15 and sulphuric acid as a catalyst. The cationic-exchange resin showed better activity than the homogeneous catalyst.

The methanolation of soybean oil was tested using tungstated zirconia-alumina (WZA), sulphated tin oxide (STO) and sulphated zirconia-alumina (SZA) as acid catalysts. The WZA catalyst was the most effective. After 20 h, it achieved conversion higher than 90% at temperatures above 250 °C.

Although the enzymatic process is still not commercially developed, a number of articles have shown that enzyme holds promise as catalyst. These studies consist mainly in optimizing the reaction conditions (temperature,
alcohol:oil molar ratio, type of microorganism which generates the enzyme, enzyme amount, time, etc.) in order to establish the characteristics for industrial applications. Several studies have shown the possibility of recycling the catalyst but the reaction time is still unfavourable if compared to the alkali catalyzed homogeneous processes. Reaction conditions reported in the literature using enzymes are summarized in Table 2.

Other conditions such as pH, use of solvent, use of immobilized or free enzyme and water content are also important in order to obtain higher conversions. An immobilized Candida antarctica lipase (Novozyme 435) was inactivated by shaking it in a mixture containing more than 1.5 molar equivalents of methanol to oil. However, when methanol was added stepwise, the conversion to methyl esters was almost complete (98.4%). The enzyme could be reused 50 times without loss of activity. An even more dramatic increase of the lipase efficiency was shown when it was pre-treated by a consecutive incubation in methyl ester and oil prior to reaction. Candida antarctica (Novozyme 435) was pre-incubated in methyl oleate for 0.5 h and subsequently in soybean oil for 12 h.

5. Methods of Analysis for Characterization of Biodiesel

The vegetable oil or animal fat source reflects basically on the composition and nature of fatty acids present in triglycerides. The type and concentration of fatty acid has an outstanding effect on biodiesel stability, which will influence its storage and also its oxidation. Thus, it is important to know the characteristics of the biodiesel to be produced. A number of methods to analyze biodiesel have been published. These methods are important to identify and quantify the fatty acid composition of biodiesel. Therefore, a search for scientific articles was carried out, focused on these methods, and a discussion of the advantages and disadvantages of the most commonly employed methods is shown below.

The following databases were accessed: Scielo, American Chemical Society and Elsevier Databases. From these databases, 134 articles were found using “biodiesel” as a keyword in the period from 2000 to September 2004.

The search showed that only 50.7% of the articles reported the methods employed for quantifying the fatty acid composition of biodiesel. Among 134 articles only 68 describe the method for biodiesel characterization. Gas Chromatography (GC) and High-Performance Liquid Chromatography (HPLC) are the most commonly employed methods (Figure 11).

5.1 Comparison between GC and HPLC

There is a large variety of analytical techniques for the detection, characterization and quantification of fatty acids (FAs) in biodiesel. The two most common analytical methods for studying fatty acids and triglycerides are GC and HPLC. The comparison of these two methods should focus not only the chromatographic analysis but also the sample preparation.

Although the application of HPLC to fatty acid analysis has increased over the last decade, GC is still the most widely used technique. This well-established procedure coupled with flame ionization detection (GC–FID) is very efficient and rapid when complex mixtures with a broad molecular range are analyzed.

Fatty acids are suitable for separation, quantification and analysis by GC without any derivatization, as demonstrated by James and Martin. However, derivatization of fatty acids

<table>
<thead>
<tr>
<th>Microorganism</th>
<th>Oil</th>
<th>Catalyst</th>
<th>Temp. (°C)</th>
<th>time (h)</th>
<th>Conversion (%)</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Candida antarctica</td>
<td>soybean</td>
<td>4 wt%</td>
<td>30</td>
<td>3.5</td>
<td>97</td>
<td>54</td>
</tr>
<tr>
<td>Cryptococcus spp. S-2</td>
<td>rice bran</td>
<td>2000 U</td>
<td>30</td>
<td>120</td>
<td>80.2</td>
<td>55</td>
</tr>
<tr>
<td>Candida antarctica</td>
<td>soybean*</td>
<td>4 wt%</td>
<td>30</td>
<td>48</td>
<td>93.8</td>
<td>56</td>
</tr>
<tr>
<td>Candida antarctica</td>
<td>cotton seed</td>
<td>30 wt%</td>
<td>50</td>
<td>7</td>
<td>91.5</td>
<td>57</td>
</tr>
<tr>
<td>Rhizopus oryzae</td>
<td>palm b</td>
<td>67 IU g⁻¹</td>
<td>35</td>
<td>96</td>
<td>55</td>
<td>58</td>
</tr>
<tr>
<td>Rhizomucor miehei</td>
<td>soybean</td>
<td>0.9 BAUN</td>
<td>36.5</td>
<td>6.3</td>
<td>92.2</td>
<td>59</td>
</tr>
<tr>
<td>Chromobacterium viscosum</td>
<td>jatropha a</td>
<td>10 wt%</td>
<td>40</td>
<td>8</td>
<td>92</td>
<td>24</td>
</tr>
</tbody>
</table>

* Degummed oil. b From waste bleaching earth sample. c Ethanol was used instead of methanol.
for GC analysis is performed to increase the volatility of the substances, to improve separation, and to reduce tailing. Recent developments in the technology of columns with bonded phases offer spectacular separation and little phase bleeding. Nevertheless, if a higher sensitivity is demanded, then derivatization is used. In recent times, GC has been used for the separation and analysis of geometric and positional isomers. Meanwhile, MS detection is a powerful technique for diagnostic fragmentation of saturated and unsaturated FAs, and analysis of branching positions in FAs.

The major advantages of HPLC over GC are lower temperatures during analysis, which reduces the risk of isomerization of double bonds, and the possibility of collecting fractions for further investigation. The speed of analysis, selectivity and sensitivity are important parameters in HPLC that may be improved with derivatization.

Regarding the elution order of FAs, both methods are based on chain length and degree of unsaturation. FA retention times increase with carbon number. Regarding analysis time, sample preparation is much more time consuming in HPLC than in the GC method. Nevertheless this disadvantage is largely compensated by run time.

### Table 3. Summary of some parameters for analyzing biodiesel by GC

<table>
<thead>
<tr>
<th>Column: l x id x ft (phase)</th>
<th>Injector Temp. (°C)</th>
<th>Oven Temperature</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>30 m x 0.25 mm x 0.25 μm (polyethylene glycol)</td>
<td>250</td>
<td>165-180 °C (4 °C min⁻¹)</td>
<td>73⁺</td>
</tr>
<tr>
<td>25 m x 0.53 mm x 1 μm (acified polyethylene glycol)</td>
<td>-</td>
<td>180-200 °C (4 °C min⁻¹)</td>
<td>74</td>
</tr>
<tr>
<td>25 m x 0.32 mm x 0.52 μm (% diphenyl and 95% dimethyl polysiloxane)</td>
<td>250</td>
<td>150-225 °C (5 °C min⁻¹)</td>
<td>57⁺</td>
</tr>
<tr>
<td>30 m x 0.32 mm x 1 μm</td>
<td>300⁺</td>
<td>190-215 °C (6 °C min⁻¹)</td>
<td>59⁺</td>
</tr>
</tbody>
</table>

⁺ Splitless. ⁻ FID 250 °C. ⁺⁺ FID 280 °C. ⁺⁺⁺ FID 300 °C.

### Table 4. Summary of some parameters for analyzing biodiesel by HPLC

<table>
<thead>
<tr>
<th>Column: l x id (TM)</th>
<th>Mobile phase</th>
<th>Temp (°C)</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>250 mm x 4.6 mm (STRODS-II)</td>
<td>methanol</td>
<td>40</td>
<td>75⁺</td>
</tr>
<tr>
<td>250 mm x 4 mm (LiChro CART RP-C18)</td>
<td>hexane:isopropanol:methanol</td>
<td>-</td>
<td>76⁺⁺⁺⁺</td>
</tr>
<tr>
<td>300 mm x 7.8 mm (GPC-Styragel)</td>
<td>toluene</td>
<td>-</td>
<td>77⁺⁺⁺⁺</td>
</tr>
<tr>
<td>300 mm x 7.5 mm (GPC-Styragel 0.5)</td>
<td>tetrahydrofuran</td>
<td>35-40</td>
<td>78</td>
</tr>
</tbody>
</table>

⁺ 1.00 mL min⁻¹. ⁻ Injection loop of 10 μL. ⁺⁺ Injection loop of 500 μL.

### Table 5. Solvent systems and detecting reagents parameters by TLC

<table>
<thead>
<tr>
<th>Solvent system (v/v/v)</th>
<th>Detecting reagents</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hexane:ethyl acetate:acetic acid (90:10:1)</td>
<td>iodine vapour</td>
<td>24</td>
</tr>
<tr>
<td>Isohexane:diethyl ether:acetic acid (80:20:1)</td>
<td>ultraviolet light</td>
<td>80</td>
</tr>
<tr>
<td>Chloroform:petroleum ether (1:3)</td>
<td>-</td>
<td>81</td>
</tr>
<tr>
<td>Petroleum ether:diethyl ether:acetic acid (85:15:1)</td>
<td>iodine vapour</td>
<td>82</td>
</tr>
<tr>
<td>Hexane:ethyl acetate:acetic acid (90:10:1)</td>
<td>sulphuric acid: methanol 1:1</td>
<td>64</td>
</tr>
</tbody>
</table>

The HPLC method is sensitive and precise and may be considered a good optional tool for the routine determination of 18:1 (Δ⁹), 18:2 (Δ⁹,12) and 18:3 (Δ⁹,12,15) in any oil. However, when a more complete study of FA profile is required, GC is the method of choice, since HPLC may present co-eluting peaks.

### 5.2 Parameters of GC and HPLC analyses for quantifying biodiesel conversion

Following the searches on the types of analysis employed for biodiesel characterization, some parameters were detached which are usually used in GC and HPLC. These parameters are presented in Table 3 for GC and Table 4 for HPLC.

### 5.3 Parameters of ¹H NMR and TLC analysis

The final product of the transesterification can possess some traces of triglycerides which can damage some components of the GC equipment, for example the column. To avoid this problem, it is important to be sure about the conversion achieved by the transesterification of the vegetable oil.
$^1$H NMR spectroscopy is a method used to quantify the conversion of vegetable oils in methyl esters by transesterification reaction. The relevant signals chosen for integration are those of methoxy groups in the methyl esters at 3.7 ppm (singlet) and of the $\alpha$-carbonyl methylene groups present in all fatty ester derivatives at 2.3 ppm. The latter appears as a triplet, so accurate measurements require good separation of this triplet from the multiplet at 2.1 ppm, which is related to allylic protons.

TLC is commonly used for qualitative analysis. It is fast and effective, and also can be used to verify the conversion of the oil during the reaction. After this previous analysis, the product can be quantified by GC. Some solvent systems and detecting reagents commonly used for evaluation of starting material conversion during the reaction are shown in Table 5.

6. Biodiesel Production Technology

Nowadays, the production of biodiesel in Brazil occurs mainly on an experimental scale with plants of small capacity (up to 1,000 t year$^{-1}$). However, units of medium scale of ca. 5,000 t year$^{-1}$ and of high capacity, up to 30,000 t year$^{-1}$, are being built. The transesterification process in an industrial unit with 1,800 t year$^{-1}$ is described here: the vegetable oil, animal fat or recycled oil is stored and, after acid correction, is transferred to the principal reactor. In a second reactor, the catalyst (KOH or NaOH) and the alcohol (MeOH or EtOH) are homogenized and transferred to the principal reactor. The reaction is completed in 40 min, in temperatures between 30 and 40 °C. The biodiesel (superior layer) is then separated from the raw glycerine (inferior layer) by decantation (on batch process) or by centrifugation (on continuous process). The inferior layer, beside raw glycerine, contains part of the alcohol that could be recovered and reused in the process. The raw glycerine could be purified and used, for example, in the cosmetic industry (Figure 12).

The separation step is one of the most important in the biodiesel production, since it is essential for the specification of the product according to legislation. The fuel should have similar characteristics to those of fossil diesel and, in case it does not reach the claimed specification, should be reprocessed. In order to assure the product purity and avoid engine and also environmental damage, levels of free fatty acid, alcohol, water, and glycerine should be at a minimum. The yield of biodiesel from the process can vary from 70 to 90% depending on the type of starting material, reagents and operational conditions. The transesterification is affected by alcohol:oil molar ratio, catalyst, temperature, reaction time and the content of water and free acids.

One of the most important variables affecting the yield of ester is the alcohol:oil molar ratio. The molar ratio is associated with the type of catalyst used. An acid catalyzed reaction needs a 30:1 ratio of BuOH:soybean oil, while an alkali catalyzed reaction requires only a 6:1 ratio to achieve the same ester yield for a given reaction time. Bradshaw and Meuly described that amounts of methanol above 1.75 equivalents tend to prevent the separation of the glycerine by gravity thus adding unnecessary expense to the process. Amounts of alcohol from 1.10 to 1.75 equivalents represent a practical range.

A continuous process of transesterification of triglycerides comprises bringing a stream of triglycerides into contact with a stream of alcohol in the presence of an alcoholysis catalyst. The continuous introduction of a stream of the liquid reaction product into a flash chamber, at low pressure, allows unreacted alcohol to flash into the vapour, and thus the remaining liquid contains only esters and glycerine. Transesterification can occur in different temperatures depending on the type of oil employed. The reaction of castor oil with methanol occurs most satisfactorily at temperatures in the range from 20 to 35 °C. At temperatures lower than 20 °C, the reaction proceeds so slowly as to be uneconomical. Higher temperatures than 35 °C cause loss of catalyst due to saponification, and deleterious amounts of soap are formed. The reaction can be done using a alcohol:oil molar ratio from 6:1 to 12:1 and from 0.05 to 0.35% (in weight of oil) of NaOH as catalyst. Many patents have been published, based on the transesterification route of vegetable oils or residual fats.

In the 90’s, some countries developed schemes to produce biofuels, using the methanol route and vegetable oil or residual fats as a starting material. In particular, Germany implemented a program for biodiesel production.
The vegetable oil is extracted from rapeseed, which additionally fixes nutrients in the soil, in particular, nitrogen. In the extraction process, the seed is pressed and, beside the oil, tart is obtained as a by-product, which may be added to animal feed. The industrial units of biodiesel production, in Germany, have a high level of automation. However, they operate only during part of the year due to climatic conditions. There are over 1,000 gas stations for distribution of biodiesel. At the beginning, biodiesel was used mainly in vehicular fleet in big cities. The biodiesel price to the consumer is lower than fossil diesel. The difference can reach 12%, and this can be explained by the reduction of governmental taxes.89

In France, for example, biodiesel also has been used in urban buses, in proportions up to 30% in fossil diesel (B30), in order to decrease vehicle emissions in big cities. The biodiesel chain is encouraged with the adoption of governmental decisions like reduction of taxation.

In Brazil, there is already a government law establishing the use of 2% mixtures of biodiesel in petroleum diesel (B2) by 2008.88 After that, this proportion will rise to 5% (B5). This strategy has the advantage of decreasing vehicle emissions, without changes in diesel engines.

7. Emissions

Increasing concern about combustion related pollutants, such as particulate matter (PM), sulphur and nitrogen oxides, CO, metals, total organic carbon and volatile organic compounds (VOCs), amongst others, is driving governments to put more stringent requirements on fuel regulations.89

Exhaust emissions from diesel burning in motor vehicles contain hundreds of compounds, either in the gas or particulate phases. Several of those compounds are proved, or suspected to be, carcinogenics, such as formaldehyde, acrolein, PAHs, nitro-PAHs, benzene, sulphate, etc.

Amongst vehicular fuels, diesel also produces larger quantities of fine particulate matter, which consists basically of carbonaceous material, metals, soluble organic fraction (SOF) and sulphate.90 Epidemiological studies have associated the daily exposure to fine particulate matter (d$_{\text{p}}$ ≤ 2.5 μm) to mortality and morbidity. Also, statistical analysis has suggested that sulphate from fine particulate matter is very correlated to total mortality,91 while mutagenicity associated with the organic fraction of diesel emissions is largely dependent on the sulphur and aromatic content of the fuel.90

As a consequence, the use of alternative fuels in order to reduce the environmental impacts of diesel emissions has been extensively investigated. Trends in the regional use of biomass-derived fuels, such as alcohols, biodiesel and agricultural residues - as a proposed control initiative against elevated carbon monoxide levels in urban areas - have expanded to a global scale. The general interest in biofuels obviously results from economic considerations but, more recently, increasing attention has also been due to the role that their combustion products may have on the question of global warming and on smog formation, as well as an alternative to substitute methyl tert-butyl ether as a gasoline additive.

The vehicular fleet in Brazil is quite unique in the world, as its light-duty vehicles, running in the largest cities, are fuelled with either a blend 22-24% (v/v) of anhydrous ethanol in gasoline – namely gasohol - or pure hydrated ethanol (95%). At the beginning of the 90’s, about 41 % of the light-vehicles running, for example, in Salvador - the capital of Bahia, a big state in the north-eastern region of Brazil - were fuelled with pure hydrated ethanol, and that proportion was probably very similar to those in many other big Brazilian cities.92 Meanwhile, in the middle of the 90’s, the atmospheric levels of ethanol measured in that city were in the range of 12.5 to 154.9 ppb.93

There is evidence that the use of alternative fuels such as ethanol, leads to an increase in the atmospheric levels of acetaldehyde. The acetaldehyde-to-formaldehyde ratios are well established in many works.94-96 The weighted aldehyde emission factors, for the ethanol vehicles operated under the urban driving cycle without catalytic converter, were in the range of 19 to 46 mg km$^{-1}$ for formaldehyde and 143 to 246 mg km$^{-1}$ for acetaldehyde. Immediately after the adoption of the use of converters, substantial reductions were observed, although the acetaldehyde emissions remained higher than formaldehyde.97

Volatile organic compounds (VOCs) play a significative role in the formation process of photochemical smog. Photooxidants are formed in the atmosphere when VOCs react with nitrogen oxides (NO$_x$), in the presence of sunlight and OH radicals. In urban areas, vehicle fuels constitute an important source of VOCs and NO$_x$ emissions to the atmosphere, either by burning or evaporative losses. A comparative study of the two types of light-duty vehicular fuels presently used in Brazil - gasohol and hydrated ethanol - as potential precursors for ozone formation in the atmosphere of urban cities, was conducted outdoors, using two Teflon chambers which were filled, respectively, with each fuel, at high initial VOC to NO$_x$ ratios and exposed to the sunlight. Ozone formation, as well as NO$_x$ concentration profiles were measured directly using ozone and NO$_x$ continuous analyzers. The results showed that, for the same initial volumes of the two fuels into the bags, the ozone peak concentrations are on average 28% higher for alcohol
compared with gasohol. In addition, the ozone formation process showed a tendency to start earlier for gasohol but, once started, its concentration increased up to two times faster than for alcohol fuel.98

From a perspective of the air quality in large urban centres, especially those to which public transportation is largely dependent on heavy-duty diesel vehicles, as is the case of the majority of Brazilian cities, several studies have pointed to the fact that biodiesel burning, either pure or in mixtures with diesel, can reduce the emissions of carbon monoxide (CO), particulate matter (PM), total hydrocarbons (THC) and sulphur compounds, although quite different conclusions can also be found related to other substances, since an increase in NOx emissions with mean quantities of 610 ± 29 and 614 ± 30 ng kW−1 h−1 in particulate phase of diesel and B20 respectively, of 3818 ± 296 and 2920 ± 1105 ng kW−1 h−1 in vapour phase. Since the lighter PAHs were not determined, it was not possible to establish a comparison between their emissions from both fuels. Benzo(a)pyrene (BaP), a carcinogenic and mutagenic PAH, was present in both fuels, much less are available with approaches related to unregulated substances.

Regarding regulated substances, no significant differences were observed between both fuels. The determined emissions (g kW−1 h−1) for diesel and B20 were, respectively: THC (0.6 ± 0.1 and 0.5 ± 0.1); CO (2.1 ± 0.2 and 1.9 ± 0.2); NOx (5.3 ± 0.3 and 5.3 ± 0.4); PM (0.26 ± 0.02 and 0.27 ± 0.03). Also, soluble organic fractions (SOF) were 49 ± 1% for diesel and 52 ± 12% for B20.

Regarding the unregulated substances, on the other hand, the sum of PAHs, both in vapour and particulate phases, were higher for diesel, as expected since aromatics are virtually absent in biodiesel. The average emissions, respectively, 9661 and 8416 ng kW−1 h−1, for diesel and B20 in particulate phase and 985 and 418 ng kW−1 h−1 in vapour phase. These values show a clear predominance of PAHs in particulate over vapour phase. Amongst the 10 PAHs determined in this work, beginning in the benz(a)anthracene, the most abundant was chrysene, with average emissions, in diesel and B20 respectively, of 3818 ± 296 and 2920 ± 1105 ng kW−1 h−1 in particulate and 354 ± 97 and 131 ± 31 ng kW−1 h−1 in vapour phase. Since the lighter PAHs were not determined, it was not possible to establish a comparison between their emissions from both fuels. Benzo(a)pyrene (BaP), a carcinogenic and mutagenic PAH, was present in emissions with mean quantities of 610 ± 29 and 614 ± 30 ng kW−1 h−1 in particulate phase of diesel and B20 respectively. The mean values were considerably lower in the vapour phase (73 ± 29 and 24 ± 12 ng kW−1 h−1).

Similar conclusions were also obtained by Bünger et al.,117 regarding the PAH emissions by both diesel and methyl ester of rapeseed oil, running in passenger cars. They also found higher mutagenic potency in exhausts
from diesel, when compared to that from rapeseed oil ester, and have concluded that this was probably due to the greater amount of PAHs, in the emissions from the fossil fuel. Costa Neto et al.\textsuperscript{118} presented the utilization of used frying oil for the production of biodiesel, and discussed the performance of biodiesel in diesel engines. The emissions derived from engines running on unused vegetable oils were compared to those derived from engines running on conventional diesel. Two nitro-PAHs were also determined, namely 2-nitrofluorene (2-NFLU) and 1-nitropyrene (1-NP). The mean values for their sum, in emissions from diesel and B20 were, respectively, 156±22 and 248±169 ng kW\textsuperscript{-1} h\textsuperscript{-1}, with predominance of NFLU. Regarding the carbonyl compounds, the sum of the twelve, found in this work, gave values of 79.3 and 94.5 mg kW\textsuperscript{-1} h\textsuperscript{-1} for diesel and B20. Formaldehyde, followed by acetaldehyde, were the two most abundant in both fuels, with average values of 42.7 and 15.8 mg kW\textsuperscript{-1} h\textsuperscript{-1} in diesel and 50.6 and 16.8 mg kW\textsuperscript{-1} h\textsuperscript{-1} in B20. Results of this study point to an increase in carbonylic compound emissions, when biodiesel is blended with diesel, although different and conflicting results can be found in the literature.\textsuperscript{110,119} Biodiesel blends with diesel are expected to emit less light aromatic compounds than pure diesel, since in the vegetal oil there is virtually no aromatic content. Turriobaldassari et al.\textsuperscript{90} had observed this in their work, with mean values of 29.9 and 13.9 mg kW\textsuperscript{-1} h\textsuperscript{-1} for diesel and B20, respectively, benzene and toluene comprising more than 60% of the total. Despite this, it is important to highlight that in other works distinct results were also reported.\textsuperscript{110,119}

8. Conclusions

The number of articles and patents on biodiesel testifies to the growth of interest in this subject. Especially USA, Japan and European countries have dedicated efforts to reach a good technological level for biodiesel production. Soybean is the most commonly used oil source, but the chosen source changes from place to place according to the availability. Regarding the seeds, good sources should have low production costs, high percentages of oil and high yields per hectare. The increasing use of recycled oil as a biodiesel source is due to its lower cost in relation to raw or refined oils. From the chemical point of view, good oil for biodiesel production must be rich in long chain and low unsaturation level fatty acids.

The transesterification process in an industrial unit can be summarized as following: the oil, after acid correction, is transferred to the principal reactor. In a second reactor, the catalyst (KOH or NaOH) and the alcohol (MeOH or EtOH) are homogenized and transferred to the principal reactor. The reaction is completed in 40 min, at a temperature between 30 and 40 °C. The biodiesel (upper layer) is then separated from the raw glycerine (lower layer) by decantation (on batch process) or by centrifugation (on continuous process). The lower layer, besides raw glycerine, contains part of the alcohol that could be recovered and reused in the process. The raw glycerine could be purified and used, for example, in the cosmetic industry.

There has been a continuous development of new catalysts for biodiesel production, especially for heterogeneous processes. Regarding the development of new catalysts for homogeneous processes, there are few articles in the literature. Among the heterogeneous catalysts, it could be observed that enzymes are the most commonly studied in the last few years. However, the performance is still unfavourable compared to the alkaline homogeneous catalysts. The use of different available acid and alkaline catalysts has been reviewed and particular attention has been given to the more recently developed ones.

GC and HPLC are the most commonly employed methods for characterization of biodiesel composition. HPLC is a technique that gives good results but the cost of the analysis is a little higher than GC. Thus, GC is preferred to quantify biodiesel.

Results available till now, regarding comparative studies between emissions from pure fossil diesel and mixtures, in variable proportions, with biodiesel, are still conflicting in several aspects. While some studies report, for biodiesel, overall reductions in almost all regulated substances, others point to comparable values or even increases in emissions. Also, while regulated emissions are by now already reasonably well documented in several works, non-regulated emissions, on the contrary, lack more substantial research. Differences between emissions, produced by fuel burning, depend on factors that are not only related to the type of fuel, but also to engine characteristics and test conditions. The combination of all of these variables may be leading to the different conclusions, among the studies conducted till now, but is also an encouragement for further researches.

9. Brazilian Perspectives

Despite the potential in Brazil for oleaginous cultivation, and consequently for biodiesel production, there are challenges for setting up and consolidating a wide program of biodiesel use as fuel. The principal challenges and considerations focused technological, agronomic and infrastructure areas.
9.1 Technological perspectives

(i) Develop new process of transesterification (heterogeneous catalysts, ethylic route, etc.), with the possibility of reduction of by-products and costs with separation and purification of biodiesel; (ii) improve the product stability by use of additives; (iii) optimize the industrial plants to obtain control of the continuous process; (iv) improve the quality standard of biodiesel to avoid motor and environmental damage; (v) investigate new uses for glycerine; (vi) evaluate and control the quality of vehicle emissions in motors using biodiesel as fuel.

9.2 Agro-economic perspectives

(i) Plan and execute an ecologically sustainable agricultural zoning of oleaginous cultivation; (ii) assure supply of raw material with minimal costs to the producer; (iii) establish financial support to amplify the cultivation of oil plants; (iv) develop research to select new varieties and systems of tilling with low environmental impact; (v) intensify search for genetic improvement of oil plants, with the purpose of increased productivity and yield of oil for biodiesel; (vi) develop technology to use the tart castor bean and tart soy as animal feed.

9.3 Infrastructure perspectives

(i) Improve the infrastructure of transport and distribution; (ii) improve the connections of the productive chain.

9.4 Environmental perspectives

(i) Reduce the emissions of harmful species (e.g. CO, particulate matter, sulphur compounds and THC); (ii) get a clear diagnosis on the environmental impacts of biofuel uses, along with its advantages and benefits.

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Abbreviation List

\( \Delta \) position of double bond in fatty acids
ANEEL Agência Nacional de Energia Elétrica
ANP Agência Nacional de Petróleo
ASTM American Society for Testing and Materials
B100 pure biodiesel
B20 20% B100 and 80% diesel
B30 30% B100 and 70% diesel
B35 35% B100 and 65% diesel
B5 5% B100 and 95% diesel
B2 2% B100 and 98% diesel
BaP benzo(a)pyrene
BIREME Latin American and Caribbean Center on Health Sciences Information
CNPq Conselho Nacional de Desenvolvimento Científico e Tecnológico
ETS-10 titanosilicate structure-10 zeolite
FAPERJ Fundação de Amparo à Pesquisa do Estado do Rio de Janeiro
FAPESB Fundação de Amparo à Pesquisa do Estado da Bahia
FAPESP Fundação de Amparo à Pesquisa do Estado de São Paulo
FAs fatty acids
FID flame ionization detection
GC gas chromatography
HC hydrocarbons
‘H NMR hydrogen nuclear magnetic resonance
HPLC high performance liquid chromatography
IF impact factor
INPADOC International Patent Documentation Center
INPI Instituto Nacional da Propriedade Industrial
MS mass spectrometry
2-NFLU 2-nitrofluorene
NO\(_x\) nitrogen oxides
1-NP 1-nitropyrene
PAHs polycyclic aromatic hydrocarbons
PCT Patent Cooperation Treaty
PM particulate matter
SOF soluble organic fraction
STO sulphated tin oxide
SZA sulphated zirconia-alumina
TBD 1,5,7-triazabicyclo[4,4,0]dec-5-ene
TGA thermo-gravimetric analysis
THC total hydrocarbons
TLC thin layer chromatography
USPTO United States Patent and Trademark Office
\( v/v \) volume/volume proportion
VOCs volatile organic compounds
WZA tungstated zirconia-alumina

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