A Review on Green Trend for Oil Extraction Using Subcritical Water Technology and Biodiesel Production

Weal Abdelmoez*, Eman Ashour and Shahenaz M. Naguib

Chemical Engineering Department, Faculty of Engineering, Minia University, Minia, Egypt

Abstract: It became a global agenda to develop clean alternative fuels which were domestically available, environmentally acceptable and technically feasible. Thus, biodiesel was destined to make a substantial contribution to the future energy demands of the domestic and industrial economies. Utilization of the non edible vegetable oils as raw materials for biodiesel production had been handled frequently for the past few years. The oil content of these seeds could be extracted by different oil extraction methods, such as mechanical extraction, solvent extraction and by subcritical water extraction technology SWT. Among them, SWT represents a new promising green extraction method. Therefore this review covered the current used non edible oil seeds for biodiesel production as well as giving a sharp focus on the efficiency of using the SWT as a promising extraction method. In addition the advantages and the disadvantages of the different biodiesel production techniques would be covered.

Key words: biodiesel, non-edible oils, subcritical water, esterification, transestrification

1 Introduction

Energy consumption was inevitable for human existence. There were various reasons for the search of an alternative fuel that was technically feasible, environmentally acceptable, economically competitive, and readily available. The first foremost reason was the increasing demand for fossil fuels in all sectors of human life, be it transportation, power generation, industrial processes, and residential consumption. The second reason was that fossil - fuel resources were non-renewable, and they will be exhausted in the near future. The last reason was the price instability of fuels such as crude oil, which was a serious threat for countries with limited resources. Several alternatives such as wind, solar, hydro, nuclear, biofuel, and biodiesel had been suggested but all of them were still in the research and development stage1. Biodiesel was considered as an important renewable energy source because of its potential to fulfill the energy demand, reduce green house gasses and global warming2.

It was well known that in modern economics, energy had the main role in the advancement of all sectors including agricultural, transportation, telecommunication and industrial sectors. As a result, worldwide energy consumption was expected to grow in a faster rate than the population growth. According to the International Energy Agency (IEA), an anticipated increase of 53% of global energy consumption was foreseen by 2030. Energy had become a key to boost the economic development across the world. Although the fossil fuels were limited and non-renewable, nevertheless the demand of these resources were increasing rapidly3, 4.

Therefore, it became a global agenda to develop clean alternative fuels which were domestically available, environmentally acceptable and technically feasible. According to the Energy Policy Act of 1992 (EPACT, US), natural gas, biodiesel, ethanol, electricity and methanol were the main prospective alternative fuels that could reduce global warming, fossil fuels consumption and exhaust emissions. As an alternative fuel, biodiesel was one of the best choices among others due to its environment friendly behavior and similar functional properties with diesel fuel. Using biodiesel in internal combustion engines could play a great role in reducing fossil fuel demand, environmental impact and the adverse effect on human health. Biodiesel derived from vegetable oil5 had been shown to be a potential alternative replacing petroleum -derived diesel oil for diesel engine6.

In the production of biodiesel more than 95% of feed
stocks came from edible oils. Use of such edible oil to produce biodiesel was not feasible in view of a big gap in demand and supply of such oils as food. Moreover it might cause some problems such as the competition with the edible oil market, which increased both the cost of edible oils and biodiesel. Moreover it would cause deforestation in some countries because more and more forests had been felled for plantation purposes. In order to overcome these disadvantages, many researchers, scientists, technologists as well as Industrialists were interested in non-edible oil sources like palm oil, castor oil, jatropha, jojoba, karanja, neem oil, soap nut, and algae which were not suitable for human consumption because of the presence of some toxic components in the oils. Furthermore, non-edible oil crops could be grown in waste lands that were not suitable for food crops and the cost of cultivation was much lower because these crops could still sustain reasonably high yield without intensive care. However, most non-edible oils contain high free fatty acids. Thus they might require multiple chemical steps or alternate approaches to produce biodiesel, which would increase the production cost, and may lower the ester yield of biodiesel below the standards.

One of the main aims of this review was providing an information on the different available non-edible oil resources for biodiesel production namely castor oil, jatropha, jojoba, karanja, neem oil, soap nut, and algae (Fig. 1).

2 Non-edible oils for biodiesel production

The availability of any feedstock for biodiesel production depends on the regional climate, geographical locations, local soil conditions and agricultural practices of any country. One of the main reasons that make biodiesel production more convenient as an energy substitute was the accessibility of biodiesel feed stocks worldwide.

It was very important that any potential feedstock for biodiesel production should be attainable at the lowest price and in an abundant compared to diesel in the competitive market.

2.1 Algae oil (cyanobacteria)

Algae (micro and macro) oils are one of the best sources for synthesis of biodiesel. Micro algae are classified as diatoms (bacillar-iophyceae), green algae (chlorophyceae), golden brown (chryso-phyceae) and blue-green algae (cyano phyceae). Microalgae may be the potential and economical source of biodiesel because of high yielding feedstock. In microalgae the oil content can exceed 80% by weight of dry biomass and oil levels of 20–50% are common in any micro algae.

Micro algae have long been recognized as potentially good sources for bio-fuel production because of their high oil content, rapid biomass production (double their biomass production within 24 h) and it is much faster and easier to


2.2 Castor oil (Ricinus communis)

Castor plant has the botanical name R. communis of the family Euphorbiaceae. It contains about 46–55% oil by weight. Castor is originally a tree or shrub that can grow above 10 m high, reaching an age 4 years. Castor grows in the humid tropics to the sub-tropical dry zones (optimal precipitation 750–1000 mm, temperature15–38°C). The oil of castor plant is viscous, pale yellow, non-volatile, non-drying oil with a bland taste and sometimes used as a purgative and the oil has a slight odor. Relative to other vegetable oils castor oil has a good shelf life and it does not turn rancid unless subjected to excessive heat and is a good source of raw material for varieties of other field like adhesives, coatings, soaps, lubricants, paints and dyes.

2.3 Jatropha oil (Jatropha curcas L.)

Jatropha plant has the botanical name J. curcas, and it is a multipurpose bush small tree belonging to the family of Euphorbiaceae; it can grow under a wide variety of climatic conditions like severe heat, low rain and sometimes used as a purgative and the oil has a slight odor. Relative to other vegetable oils castor oil has a good shelf life and it does not turn rancid unless subjected to excessive heat and is a good source of raw material for varieties of other field like adhesives, coatings, soaps, lubricants, paints and dyes.

2.4 Jojoba oil (Simmondsia chinensis)

The seed of jojoba plant, which has botanical name S. chinensis, appears to be a promising scope for cultivation in arid and semi-arid areas. The Jojoba seed is nut shaped and is around 1–2 cm long, with red-brown to dark-brown color. The seed contains 45 and 55 wt% of Jojoba oil-wax. The chemical structure of jojoba oil allows its use in many lubricating oil formulation and biodiesel production. Jojoba oil and its derivatives find applications in the fields of cosmetics, pharmaceuticals and lubricants.

2.5 Karanja seed (Pongamia pinnata)

Pongamia oil is a non-edible oil extracted from seeds of P. pinnata (L.) Pierre, family Fabaceae commonly known as ‘Karach’, ‘Karanja’ in Assam. Karanja is medium sized fast-growing ever green tree, 12–15 m height, branches spread into hemispherical crown of dense green leaves. It has a disagreeable odor and bitter taste. Seeds are 10–20 mm long and light brown in color. Pongamia can survive in adverse conditions like drought, heat, frost, salinity, etc. The karanja seed kernel contains 27–39 wt% oil. The oil has been used mainly for leather tanning, lighting and to a smaller extent in soap making, medicine, and lubricants.

2.6 Neem (Azadirachta indica)

Neem (A. indica) is a fast-growing tree that can reach a height of 15–20 m. Neem can be grown on very marginal soils that may be very rocky, shallow, dry or pan forming. The neem seed contains 35–45% of oil. Neem products have been observed advantageous to be anthelmintic, anti-fungal, anti-diabetic, antibacterial, antiviral, contraceptive and sedative. Neem gum is used as a bulking agent and for the preparation of special purpose food (for diabetics) and neem leaf paste is applied to the skin to treat acne.

A neem tree can produce many thousands of flowers, and in one flowering cycle, a mature tree may produce a large number of seeds. Neem trees start bearing harvestable seeds within 3–5 years, and full production may be reached in 10 years, and this will continue up to 150–200 years of age.

2.7 Soap nut (Sapindus mukorossi)

Soap nut (S. mukorossi) is a fruit of the soap nut tree and the plant grows very well in deep loamy soil sand leached soils, so cultivation of soap nut in such soil avoids potential soil erosion. (30% oil content). Also it was used in a commercial cleanser and medical applications based on its usages including antidermatophytic.

The oil from soap nut had been considered as promising non-edible oil having significant potential for biodiesel production. This was because it was the third most productive vegetable oil producing crop in the world, after algae and oil palm.

3 Subcritical water technique as extraction method

Extraction always involved a chemical mass transfer from one phase to another. The principles of extraction were used to advantage in everyday life, for example in making juices, coffee and others. To reduce the use of organic solvent and improve the extraction methods of constituents of plant materials, new methods such as microwave assisted extraction (MAE), supercritical fluid extraction (SFE), accelerated solvent extraction (ASE) or pressurized liquid extraction (PLE) and subcritical water extraction (SWE), also called superheated water extraction or pressurized hot water extraction (PHWE), had been introduced.

Lately, subcritical water technology had attracted many
researchers for its versatile applications in the field of environment as a green alternative process to solvent extraction.

The past decade had witnessed an intense interest in the use of subcritical water technology (SWT) in separation processes. SWT was an alternative extraction method where water instead of organic solvents were used as the extraction media. On heating within the critical point of water (Fig. 2) (temperature <374°C, pressure 22.1 MPa) under enough pressure to maintain the liquid state, water (subcritical water in this state) was reported to have distinctive properties. These were a low dielectric constant and a high ion product. Many extractable components from different biomass could be easily extracted under these conditions.

Based on the research worked published in the recent years, it had been shown that the SWE was cleaner, faster and cheaper than the conventional extraction methods.

3.1 Extraction mechanism

The SWE process could be proposed to have six sequential steps: (1) rapid fluid entry; (2) desorption of solutes from matrix active sites; (3) diffusion of solutes through organic materials; (4) diffusion of solutes through static fluid in porous materials; (5) diffusion of solutes through layer of stagnant fluid outside particles; and (6) elution of solutes by the flowing bulk of fluid. The plots the amount of compound extracted versus solvent flow rates and versus solvent volume can determine the relative importance of these steps. For example, if the rate of extraction was controlled by intra-particle diffusion or kinetic desorption, the increase in bulk fluid flow rate would have little effect on extraction rate. On the other hand, if the extraction was controlled by external film transfer diffusion, extraction rates increase with solvent flow rate.

3.2 Example of Oil Extraction Using Subcritical Water Technology (SEW)

No commercial SWE equipment was available, but the apparatus was easy to construct in the laboratory. SWE was performed in batch or continuous systems. The SWE was carried out in a laboratory-built apparatus. After the preparation step, the SCE was carried out in stainless steel pipes with Swagelok caps. The seeds were then charged into the reactor tube. The reactor was then sealed and immersed in a heating bath containing one of the heating media. The extraction was carried out in the range of 180-260°C, and the pressure inside the reactor was estimated from the steam table. After the desired extraction time, the reactor was immediately cooled down by immersing it into a cold-water bath. The extract product contained three phases, the oil phase, the aqueous phase (including oil and water), and the solid phase. The three phases were separated through simple centrifugation and vacuum filtration processes. To recover any traces of oil from the aqueous phase, we added petroleum ether to extract any oil that could be emulsified into the water phase. Then, the petroleum ether was evaporated by heating it in a furnace at 80°C. The extracted oil was then weighed and the data was recorded.

3.3 Kinetics study for Oil extraction Using Subcritical Water Technology

The oil extraction process could be viewed as an irreversible consecutive unimolecular-type first order. In which, the oil was firstly extracted from inside the seed by diffusion following a first order-like mechanism. Then the extracted oil, due to extending the extraction time under the high temperature and pressure, was subjected to a decomposition reaction following a first order mechanism. Accordingly, the oil extraction process could be simplified into the following reaction scheme

\[ A \xrightarrow{k_1} R \xrightarrow{k_2} S \]

Where, A, R and S, represent the total oil in the seed, extracted oil, and the decomposition product, respectively and \( k_1 \) and \( k_2 \) represent the specific reaction rate constants for the extraction and decomposition steps, respectively.

The rate equations for the three components were:

\[ \frac{dC_A}{dt} = -k_1C_A \]  (1)

\[ \frac{dC_R}{dt} = k_1C_A - k_2C_r \]  (2)
The concentration of decomposition product, respectively.

\[ \text{where } C_a, C_r, \text{ and } C_s \text{ represent the concentration of the total oil in the seed, the concentration of extracted oil, and the concentration of decomposition product, respectively.} \]

Integration of the rate equations results in:

\[ - \ln \left( \frac{C_a}{C_{ao}} \right) = kt \text{ or } C_a = C_{ao} e^{-k_1t} \tag{4} \]

Where \( C_{ao} \) represents the initial concentration of the oil in the seeds. To find the variation in concentration of \( R \), substitute the concentration of \( A \) from Eq. 1 into the differential equation governing the rate of change of concentration of the extracted oil; \( R \):

\[ \frac{dC_r}{dt} = k_2 C_r \tag{5} \]

by integrating Eq. 5, the final variation in concentration of the extracted oil, \( R \), was:

\[ C_r = C_{ao} k_1 \left( \frac{e^{-k_1t}}{(k_2 - k_1)} \right) + \left( \frac{e^{-k_2t}}{(k_1 - k_2)} \right) \tag{6} \]

Eq. 6 had been used to elucidate the concentration time course of jojoba oil extraction.40

The kinetic study of the oil extraction from spent bleaching earth (SBE) & jojoba seed under subcritical water conditions were based on consecutive steps. In these steps, the oil was firstly extracted from inside the powder by diffusion following a first order-like mechanism. Then the extracted oil, due to extending the extraction time under high temperature and pressure, was subjected to a decomposition reaction following a first order mechanism.

The experimental data correlated well with the irreversible consecutive unimolecular-type first order mechanism using Eq. 6. The values of both oil extraction rate constant and decomposition rate constant were calculated through non-linear fitting using DataFit software. Table 1 kinetic & thermodynamic study of oil extraction from spent bleaching earth (SBE) & jojoba seed. The extraction rate constants were 0.018, 0.03, 0.08, 0.1 and 0.14 min \(^{-1} \), while the decomposition rate constants were 0.016, 0.012, 0.02, 0.025 and 0.0205 min \(^{-1} \) at extraction temperatures of 180, 230, 250, 260 and 270°C, respectively for SBE & The values of both oil extraction rate constants and the decomposition rate constants for jojoba were 0.0094, 0.0187, and 0.1154 min \(^{-1} \), while the decomposition rate constants were 0.0985, 0.1074, and 0.0078 min \(^{-1} \) at extraction temperatures of 180, 200, and 240°C, respectively. The results showed that by increasing the extraction temperature, the extraction rate increases. However, the decomposition rate decreased by increasing the extraction temperatures. Such phenomena could be explained on the basis of a compensation mechanism. Such mechanism could be viewed as follows: by increasing the extraction temperature, part of the fat phase decomposes into the oil phase, resulting in an increase in the oily content and a decrease in the apparent overall decomposition step. However, by increasing the extraction time, part of the degradable products is finally decomposed into the gas phase (as mentioned above), resulting in the reduction of the overall mass balance. This point needs more investigation and analysis.

### 3.4 Thermodynamic study for Oil extraction Using Subcritical Water Technology

The oil extraction process was also analyzed thermodynamically. Arrhenius was used for describing the temperature dependence of the reaction rate. The Arrhenius equation is founded on the empirical observation and common sense, that is, the Chemical Perception which suggests that the higher the temperature the faster is a given chemical reaction will proceed (Eq. 7).

\[ K = A \exp \left[- \frac{E_a}{RT} \right] \tag{7} \]

Where \( A \) is the pre-exponential factor (sometimes called

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Spent Bleaching Earth (SBE)</th>
<th>jojoba Oil Extraction Rate, ( k ), (min (^{-1} ))</th>
<th>Spent Bleaching Earth (SBE)</th>
<th>jojoba Oil Decomposition Rate, ( k ), (min (^{-1} ))</th>
</tr>
</thead>
<tbody>
<tr>
<td>180</td>
<td>0.018</td>
<td>0.00948</td>
<td>0.016</td>
<td>0.0985</td>
</tr>
<tr>
<td>260</td>
<td>260</td>
<td>0.1</td>
<td>0.0944</td>
<td>0.025</td>
</tr>
<tr>
<td>Activation energy, ( E ) (KJ/mol)</td>
<td>46.1</td>
<td>82.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Frequency, ( A ) (min (^{-1} ))</td>
<td>3084.05</td>
<td>464483</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Standard Gibbs free energy of activation, \( \Delta G^\circ \) (KJ.mol \(^{-1} \)) at temperature:

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>( \Delta G^\circ ) (KJ.mol (^{-1} ))</th>
</tr>
</thead>
<tbody>
<tr>
<td>180</td>
<td>44.3</td>
</tr>
<tr>
<td>260</td>
<td>59.5</td>
</tr>
</tbody>
</table>

| Table 1 Kinetic & thermodynamic study of oil extraction from spent bleaching earth (SBE) & jojoba seed. |
frequency factor), $E$ is the apparent activation energy, $T$ is the absolute temperature, and $R$ is the gas constant ($8.31451 \, \text{J} \cdot (\text{mol} \cdot \text{K})^{-1}$).

On the integration of both sides of Eq. 7:

$$\ln k = -\frac{E_a}{R}T + \ln A \quad (8)$$

By plotting $\ln k$ vs. $1/T$, a linear relationship is obtained and one can determine $E_a$ from the slope ($\frac{-E_a}{R}$)/and $A$ from the y-intercept. This equation assumes that $E_a$ and $A$ are constants or nearly constant with respect to temperature.$^{50}$

The activation thermodynamic parameters were calculated from the transition-state theory using the Eyring equation. Eyring (1935) formulated his equation, which represents a theoretical construct based on transition-state theory. The Eyring equation in its thermodynamic version becomes:

$$k = (k_BT/h)e^{-\Delta G^*/RT} \quad (9)$$

$$k = (k_BT/h)e^{\Delta S^*/R}e^{-\Delta H^*/RT} \quad (10)$$

where $k$ is the rate constant, $\Delta G^*$ is the standard Gibbs free energy of activation, $\Delta H^*$ is the standard enthalpy of activation, $\Delta S^*$ is the standard entropy of activation, $k_b$ is the Boltzmann constant ($1.380658 \times 10^{-23} \, \text{J} \cdot \text{K}^{-1}$), $h$ is Planck’s constant ($6.6260755 \times 10^{-34} \, \text{J} \cdot \text{s}$), $R$ is the gas constant, and $T$ is the absolute temperature in Kelvin.

On taking a logarithm for both sides of eq. 10, one obtains:

$$\ln (k/T) = [\ln (k_b/h) + (\Delta S^*/R)] - \Delta H^*/RT \quad (11)$$

By plotting $\ln (k/T)$ vs. $1/T$, a linear relationship is obtained and one can determine $\Delta H$ from the slope which is equal to the value $(\Delta H^*/R)$ and $\Delta S^*$ is determined from the y-intercept $[\ln (k_b/h) + (\Delta S^*/R)]$.

The Gibbs free energy of activation can be determined by:

$$\Delta G^* = \Delta H^* - T\Delta S^* \quad (12)$$

In this section, the thermodynamic properties of the oil extraction process were investigated. For spent bleaching earth (SBE), the activation energy, $E_a$, and the frequency factor, $A$, were calculated from these values using Eq. 7. Accordingly, the values of $E_a$ and $A$ were 46.1 $\, \text{kJ} \cdot \text{mol}^{-1}$ and 3084.05 min$^{-1}$, respectively. To gain a deeper understanding of the extraction process, the activation thermodynamic $\Delta H^*$ and $\Delta S^*$ parameters were calculated from the transition-state theory. The standard enthalpy of activation $(\Delta H^*)$ and the standard entropy of activation $(\Delta S^*)$ were calculated using eq. 11. The value of $\Delta H^*$ was $-42.038 \, \text{kJ} \cdot \text{mol}^{-1}$ and the value of $\Delta S^*$ was $-0.1906 \, \text{kJ} \cdot \text{K}^{-1} \cdot \text{mol}^{-1}$. The value of $\Delta S^*$ gives information about the degree of order in the transition state. Low values of $\ln A$ correspond to large negative values of $\Delta S^*$, indicating unfavorable reactions, which means that the activated complex in the transition state has a more ordered or more rigid structure than the reactants in the ground state. The values of the Gibbs free energy $\Delta G^*$ of activation were calculated at different temperatures using eq. 12 were 44.3, 53.8, 57.6, 59.5 and 61.4 kJ.mol$^{-1}$ at 180, 230, 250, 260 and 270$^\circ\text{C}$, respectively. The positive values indicate a non-spontaneous reaction while the negative value indicates a spontaneous process. These results explain why high extraction temperatures and pressure were needed for improving the extraction process to overcome the non-spontaneous nature of the process.

4 Biodiesel production process
4.1 Transesterification process

The most practical and common way of producing biodiesel was by transesterification (also called methanolysis), which was a catalyzed reaction of vegetable oil in the presence of alcohol to yield biodiesel and glycerol.$^{50}$ The process starts with a sequence of three consecutive reversible reactions, wherein triglycerides were converted to diglycerides, diglycerides were converted to monoglycerides, and monoglycerides were converted to glycerol. Each step produces an ester and, as a consequence, generates three ester molecules from one triglyceride molecule. Scheme 1 shows the transesterification chemical equation.

$$\text{TG} + \text{MeOH} \rightleftharpoons \text{Catalyst} \Rightarrow \text{DG} + \text{FAME}$$

$$\text{DG} + \text{MeOH} \rightleftharpoons \text{Catalyst} \Rightarrow \text{MG} + \text{FAME}$$

$$\text{MG} + \text{MeOH} \rightleftharpoons \text{Catalyst} \Rightarrow \text{G} + \text{FAME}$$

where TG, DG and MG refer to tri-, di-, and monoglycerides, respectively, G represents glycerol, and FAME represents fatty acid methyl esters or biodiesel.$^{51}$

There were various methods for biodiesel production from non-edible oils, and the following section gives an overview of various acid, base catalytic (homogeneous, heterogeneous) and non-catalytic processes reported so far for biodiesel production.

4.1.1 Homogeneous transesterification

Homogeneous catalysts can be base catalysts, such as sodium hydroxide (NaOH) and potassium hydroxide (KOH), or acid catalysts, such as sulphuric, sulphonic, phosphoric, and hydrochloric acids. Base catalysts were preferred over acid catalysts because they have higher catalytic efficiency, lower cost, and lower reaction temperature and pressure. However, base catalysts may react with FFA present in the feedstock during transesterification, resulting in soap formation by saponification, which may consume the catalyst and reduce its efficiency.$^{50}$

Nevertheless, two-step catalyzed transesterification (Fig. 3) had been proven effective in the production of bio-
diesel from feedstock that contains a large amount of FFA. In this method, both acidic and basic catalysts were used separately: the acidic catalyst, such as sulphuric acid, was used in the first step, the esterification of FFA, whereas the basic catalyst was used in the second step, in the triglyceride’s transesterification. However, in this method, catalysts were removed after each step, according to Bautista et al. In contrast, Guzatto et al. used a modified double-step process, which uses the basic catalyst in the first step and the acidic catalyst in the second, reducing the overall process time and catalyst concentration.

4.1.2 Heterogeneous transesterification

Just like homogeneous catalysts, heterogeneous catalysts were also of two types, namely, acidic and basic. Examples of heterogeneous acid catalysts were sulphated metal oxide, heteropolyacids, acidic ion exchange resin, and sulphonated amorphous catalysts. Additional examples of heterogeneous base catalysts include zinc oxide (ZnO), calcium oxide (CaO), strontium oxide (SrO), and Na/SiO₂.

However, heterogeneous catalysts may not be practical at an industrial scale because of their high excess alcohol requirements, high reaction temperatures, long reaction times, and high energy consumption. The type of a heterogeneous catalyst for the biodiesel production from non-edible vegetable oils depends on the FFA content in the feedstock.

The solid acid catalysts were insensitive to FFA content, esterification and transesterification occur simultaneously, avoids the washing step of biodiesel, separation of the catalyst from the reaction medium was easy, low product contamination, easy regeneration, recycling of catalyst and reduced corrosion problem even with the presence of acid species.

4.1.3 Supercritical transesterification

A supercritical methanol transesterification process (Fig. 4) had several advantages over the conventional process in terms of the elimination of catalysts, easier separation of products, faster reaction rate, elimination of FFA and water contents on the ester yield, and elimination of soap formation. However, supercritical transesterification requires...
high reaction temperature and pressure, a large excess of alcohol, high energy consumption, and high capital cost. This supercritical process allowed feedstock containing certain amount of water and FFA. High reaction rate and without using any catalyst were also its advantages over the traditional biodiesel production method.\(^{57,58}\)

### 4.1.4 Enzymatic catalyzed transesterification

Biocatalysts (enzymes) had been investigated as the means to overcome the challenges presented by chemical catalysts.\(^{59}\) Lipase is the most commonly used enzyme in biodiesel production, was an effective catalyst that converts all of the FFA content in the WCO to fatty acid methyl ester. However, enzyme-catalysed transesterification (Fig. 5) was not yet used at an industrial scale for several reasons, such as the high cost of lipase, inhibition of lipase by methanol, glycerol adsorption on lipase, and long reaction time.

Enzyme-catalyzed reactions are less energy intensive, do not promote side reactions and were more environmentally friendly.\(^{60,61}\)

### 4.1.5 Microwave assisted transesterification

Microwave assisted transesterification (Fig. 6) was another alternative energy tool which could be used for the production of biodiesel. Due its advantages like reduced reaction time, better yield and environmentally benign compared to conventional heating, it was becoming the obvious choice of many researchers. Microwave transfer energy had a tendency to oscillate polar ends of molecules or ions continuously. As a result the collision and friction between the moving molecules generate heat. Heat was therefore directly deposited into the reaction media and results in rapid temperature increase throughout the sample. Several studies on micro wave assisted transesterification had been reported using different catalyst system and they observed that it offers fast, low oil: methanol ratio, easy operation, reduced energy consumption and minimum by products.\(^{1,37}\)

### 4.1.6 Ultrasound assisted transesterification

The demand for biodiesel in the world was sharply increasing. Thus, increasing the production rate for biodiesel in order to meet the demand seems to be essential. Therefore, new accelerating technologies were of great interest among researchers in this area.

Ultrasonication provides the mechanical energy for mixing and the required activation energy for initiating the transesterification reaction. Ultrasonication increases the chemical reaction speed and yield of the transesterification of vegetable oils / animal fats (Fig. 7).\(^{62,63}\)

### 4.1.7 Two-step subcritical water and supercritical methanol transesterification

Esterification had commercial utility beyond biodiesel synthesis. Esters were used as solvents in many applications such as cosmetics and in the lubricant industry. Esterification was usually catalyzed by strong mineral acids
(e.g., sulfuric or hydrochloric acid), which were corrosive and environmentally hazardous. Therefore, there was a need to develop a more environmentally friendly process to produce esters. Noncatalytic esterification was an interesting possibility\textsuperscript{64,65}. Scheme 2 and Scheme 3 show hydrolysis and esterification reactions in two-step synthesis of biodiesel\textsuperscript{40}.

Moreover, the reaction did not require a large excess of alcohol, and it could tolerate the presence of modest amounts of water in the feed stream.

5 Conclusion

Currently, utilization of the non-edible oils as feedstocks for biodiesel production had become more attractive. The not suitability for food, high oil content and the low cost of cultivation were the main advantages offered using such feed stocks. Subcritical water technology seemed to be the greener alternative method for the different oil seeds extraction. The short extraction time and the elimination of using solvents were the major benefits of using such technology. The most practical processes for biodiesel production was by transesterification reaction which was a catalyzed reaction of vegetable oil in the presence of alcohol to yield biodiesel and glycerol. Also the noncatalytic esterification was an interesting possibility because the reaction did not require a large excess of alcohol.

Acknowledgment

The authors would like to acknowledge the Egyptian Science and Technological Development Fund (STDF) for supporting the present study under the umbrella of the research project entitled "Innovative Green Extraction Technology for Medicinal Plants" with Project ID 5767.

References

3) Mofijur, M.; Atabani, A.; Masjuki, H.; Kalam, M.; Masum, B. A study on the effects of promising edible and
29) Kansedo, J.; Lee, K. Process optimization and kinetic study for biodiesel production from non-edible sea mango (Cerbera odollam) oil using response surface
36) Sapna Eipeson, V.; Manjunatha, J.; Srinivas, P.; Sindhu Kanya, T. Extraction and recovery of karanjin: A value addition to karanja (Pongamia pinnata) seed oil. Industrial Crops and Products 32, 118-122 (2010).


