1	Widely used catalysts in biodiesel production: A review
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54 Abstract

An ever-increasing energy demand and environmental problems associated with 55 exhaustible fossil fuels have led to the search for an alternative renewable source of energy. In 56 57 this context, biodiesel has attracted attention worldwide as an alternative to fossil fuel for being renewable, non-toxic, biodegradable, carbon-neutral; hence eco-friendly. Despite 58 59 homogeneous catalyst has its own merits, currently, much attention has been paid to chemically synthesize heterogeneous catalysts for biodiesel production as it can be tuned as per specific 60 requirement, easily recovered, thus enhance reusability. Recently, biomass-derived 61 heterogeneous catalysts have risen to the forefront of biodiesel productions because of their 62 sustainable, economical and eco-friendly nature. Further, nano and bifunctional catalysts have 63 emerged as a powerful catalyst largely due to their high surface area and potential to convert 64 free fatty acids and triglycerides to biodiesel, respectively. This review highlighted the latest 65 synthesis routes of various types of catalysts including acidic, basic, bifunctional and 66 nanocatalysts derived from different chemicals as well as biomass. In addition, the impacts of 67

68	different methods of preparation of catalysts on the yield of biodiesel are also discussed in
69	details.
70	
71	Highlights
72 73	• Biodiesel has attracted immense attention as a potential substitute for fossil fuels.
74 75	• Esterification and transesterification reactions to produce biodiesel are discussed.
76 77 78	• Effects of various reaction parameters in biodiesel production processes are highlighted.
79 80 81	• Review on the different characterization techniques employed in biodiesel production processes.
82 83 84	• Acid, basic and bifunctional catalysts employed in biodiesel productions are highlighted.
85	• Different sources, methods of preparation and activities of catalysts are reviewed.
86	
87	Keywords: Biodiesel, Homogeneous catalyst, Heterogeneous catalyst, Characterization, Fatty
88	acid methyl esters, Renewable energy

90 Graphical abstract:



93 **1. Introduction**

94 The exponential growth of the world's population coupled with the high standard of living has resulted in a steep increase in energy consumption.^{1, 2} The world's total primary 95 energy consumed (TPEC), which is over 150,000,000 GW h in the year 2015, is estimated to 96 rise by a triggering 57 % in 2050³. Currently, the transportation of goods and services, which 97 is the major contributor to the global economy, primarily rely on non-renewable fossil fuels. In 98 99 total primary energy consumption, 80 % of the energy consumed is associated with petroleum resources. Amongst these, 54 % is consumed in the transportation sector ⁴. It has been predicted 100 that energy consumption in the transportation section will increase with an average rate of 1.1 101 % per year. As a result, the high energy consumption of non-renewable petroleum-based fuel 102 to fulfil increasing energy demand of human society has led to an ecological imbalance, excess 103 greenhouse gas emission, acid rain, global warming and drastic decline in fossil fuel reserves. 104 These negative factors associated with excessive consumption and exhaustible nature of fossil 105 fuels compel scientific communities to look out for an alternative energy source. ^{5, 6} 106

Biofuels are an excellent source of energy and widely seen as a potential substitute for 107 fossil fuels. They are prepared from renewable sources such as plants, municipal wastes, 108 agricultural crops, agricultural and forestry by-product.⁷ Over the last few decades, biofuel such 109 110 as biodiesel has gained significant attention as an alternative fuel in the research field because of its sustainable and environment-friendly nature. Biodiesel has exhibited properties similar 111 to conventional fossil fuels (petro-diesel) and has some properties better than petro-diesel such 112 as high combustion efficiency, high flash point, high cetane number, lower CO₂ emission, 113 lower sulfur content and better lubrication.^{8,9} The high flash point of biodiesel (423 K), as 114 compared to petrodiesel (337 K), makes it non-flammable and non-explosive resulting in easy 115 and safe handling, storage, and transportation. Additionally, it can be directly used in the 116 automotive engine without any additional alteration.¹⁰ It is estimated that biodiesel demand 117 will increase to double or triple by the year 2020.¹¹ In the light of this, in the last decades, much 118 attention has been paid to research on biodiesel production with an intension make it more 119 sustainable and economical. An increasing interest in biodiesel is validated by the number of 120 research paper publications in this area as shown in Figure 1. Statistical data analysis in Figure 121 122 1 depicted the increasing trend of published research papers in the field of biodiesel. These data were collected in February 2020 from "SciFinder Database" using the keyword "biodiesel". 123 From a meagre 157 publications in the year 1993, it has exponentially increased to 3725 124 publications during its peak in 2014. 125



Figure 1: Publications per year for biodiesel during the period 1993 to Feb 2020 (Data collected from SciFinder Database)

128 **2.** (Trans)esterification

Transesterification or alcoholysis is a process to produce biodiesel in which edible/non-129 edible oils or triglyceride (TG) and alcohol have undergone nucleophilic reaction to form fatty 130 acid methyl ester (FAME) and glycerol as a byproduct.¹² The transesterification reaction is 131 illustrated in Scheme 1. Three sequential reversible reactions occurred in the transesterification 132 process; i) conversion of triglyceride to diglyceride, ii) diglyceride conversion to 133 monoglyceride, and finally, iii) monoglyceride conversion to glycerol. An ester is formed in 134 each conversion steps, thus one TG molecule produced three molecules of ester. 135 136 Transesterification reaction can efficiently convert triglyceride of vegetable oil into FAME, also called biodiesel, as depicted in Scheme 1. However, esterification reaction, a reaction 137 between carboxylic acids and alcohols to afford esters. $\frac{13-15}{13}$ is essential to converts all free fatty 138 acids (FFA) of vegetable oil into biodiesel as shown in Scheme 2. These transesterification and 139 140 esterification reactions are usually carried out in the two-pots procedure. Usually, the high FFA content of vegetable oil is first converted to esters (FAME) via esterification reaction by 141 142 employing acid catalyst followed by transesterification reaction using a basic catalyst to converts triglycerides to FAME. However, (trans)esterification reactions (or simultaneous 143 144 transesterification and esterification) in one-pot is highly desirable to convert both triglycerides and FFA of vegetable oil (with high FFAs) to FAME to reduce time and cost of biodiesel 145 production. The different routes to synthesized biodiesel are outlined in Figure 2. 146



Figure. 2: Catalyst classification for biodiesel synthesis



Scheme 1: Base-catalyzed reaction mechanism for transesterification of TGs of vegetable oil to biodiesel.



Scheme 2: Acid-catalyzed esterification of FFA content of vegetable oil to biodiesel

150 **3. Biodiesel**

American Society for Testing and Materials (ASTM) described biodiesel as a mono-151 alkyl ester produced from edible/non-edible oils or animal fats.¹⁶ Vegetable oils or animal fats 152 comprise of mainly triacylglycerol (TAG) which is an ester of fatty acids (FA) and glycerol. 153 The physicochemical properties of vegetable oils and animal fats are greatly influenced by the 154 compositions of the TAG which further often dictates the quality of biodiesel produced from 155 these resources. FA are classified broadly into two groups: i) saturated FA which has carbon-156 carbon single bond, and ii) unsaturated FA which comprises of at least one carbon-carbon 157 double bond. The FA most widely found in vegetable oils are oleic acid (18:1), palmitic acid 158 159 (16:0), linoleic acid (18:2), and linolenic acid (18:3), stearic acid (18:0), palmitoleic acid (16:1), myristic acid (14:0), arachidic acid (20:0). Besides these FA, a trace amount of 160 phospholipids, tocopherols, carotenes, sulphur compounds, and water are also found in 161 vegetable oils.^{17,18} 162

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164 **4. Feedstocks for biodiesel production**

The feedstocks for production of biodiesel are mainly edible¹⁸⁻²⁰ and non-edible 165 vegetable oils, $\frac{21-23}{21-23}$ waste cooking oils^{24,25} and animal fats including tallow, $\frac{25}{25}$ vellow grease, $\frac{26}{25}$ 166 lard, ²⁷ chicken fat²⁸⁻³⁰ and by-products from the production of Omega-3 fatty acids from fish 167 oil.^{31,32} Algae are another promising feedstocks for biodiesel which have a high potential to 168 replace edible oil due to their availability in a pond, sewage water or in shallow ocean water 169 without dislodging land used for food production.32-34 In worldwide, 31 % biodiesel is 170 produced from palm oil, 27 % from soybean oil and 20 % from rapeseed oil.³⁵ Different 171 countries used various feedstock based on their local availability. The major feedstocks used 172 in various countries are listed in Table 1. The feedstocks cost alone contributed 75 % of 173 biodiesel cost.³⁶ Thus, proper selection of feedstocks for biodiesel is necessary to reduce the 174 overall cost of biodiesel production. Ironically, utilization of edible oils (sunflower, rape, soy, 175 etc.) as feedstocks for biodiesel, called the first-generation biofuels, resulted in food versus fuel 176

problem and also disturbed the agricultural farmland allocation.^{27,37} In Malaysia, the edible 177 palm oil price has increased by 70 % due to its uses as feedstock in biodiesel industry.³⁸ In this 178 regard, to mitigate the problem associated with food versus fuel nexus and high cost of first-179 generation biodiesel, currently, non-edible oil are largely targeted as a biodiesel feedstock. 180 Another problem associated with first-generation biofuels is their remarkably higher cost than 181 fossil fuels. Hence, to bring the cost of biodiesel, utilization of non-edible oil as biodiesel 182 feedstocks is highly relevant. Non-edible oil of more than 300 species are available in South 183 Asia. India has abundant amount (approximately 1 million tons per year) of such non-edible 184 oils. Pongamia Pinnata (Karanja) and Jatropha curcas oils (JCO) were identified as the most 185 promising feedstocks by the Government of India. However, in India's biodiesel program, 186 Jatropha has got prominence over Karanja due to its less gestation period. If properly managed, 187 non-edible crops planted in different parts of the world has the potential to reduce our 188 dependence on fossil fuel for energy sources and edible oil as biodiesel feedstocks. 189

Biodiesel has been widely used biofuels in the European Union (EU) and 49 % of 190 biodiesel were produced from rapeseed oil in 2015 in EU.³⁹ With the increasing uses of waste 191 cooking oil (WCO), recycled vegetable oils and palm oils, the share of rapeseed oil in biodiesel 192 production decreased from 72 % in 2008. To reduce our dependency on edible oil and reduce 193 the price of biodiesel, EU has raised the share of WCO to 2nd position after rapeseed oil in 194 2015.⁴⁰ The top five biodiesel producer in EU are Germany, France, Spain, Netherlands, and 195 Poland. Germany is the largest biodiesel producer in EU and its production capacity increased 196 from 3.2 billion litres in 2010 to 3.8 billion litres in 2014.⁴¹ 197

Country	Feedstock	
India	Jatropha/ Pongamia Pinnata (Karanja) /	
	Soybean/ Rapeseed/ Sunflower	
Argentina	Soybeans	
Brazil	Soybeans/ Palm oil/ Castor/ Cotton oil	
France	Rapeseed/ Sunflower	
Peru	Palm/ Jatropha	
Germany	Rapeseed	
Spain	Linseed oil/ Sunflower	
Italy	Rapeseed/ Sunflower	

199 Table 1: Countrywise feedstocks used for biodiesel production.

Turkey	Sunflower/ Rapeseed
Greece	Cottonseed
Sweden	Rapeseed
Norway	Animal fats
China	Jatropha/ Waste cooking oil/ Rapeseed
	oil
Indonesia	Palm oil/ Jatropha/ Coconut
Japan	Waste cooking oil
Malaysia	Palm oil
Philippines	Coconut/ Jatropha oil
Bangladesh	Rubber seed/ Pongamia Pinnata oil
Pakistan	Jatropha oil
Thailand	Palm/ Jatropha/ Coconut oil
Iran	Palm/ Jatropha/ Castor/ Algae oil
Singapore	Palm oil
Ghana	Palm oil
Zimbabwe	Jatropha oil
Kenya	Castor oil
Mali	Jatropha oil
UK	Rapeseed/waste cooking oil
Ireland	Frying oil/ Animal fat
Canada	Rapeseed/ Animal fat/ Soybean oil
Mexico	Animal fat/ Waste Oil
USA	Soybeans/ Waste oil/ Peanut
Cuba	Jatropha curcas/ Moringa/ Neem oil
Australia	Jatropha/ Pongamia/ Waste cooking oil/
	Animal tallow
New Zealand	Waste cooking oil/ Tallow

Wide types of feedstocks such as edible plant oils, non-edible oils, waste cooking oils, animal fats, and algal oil have been considered for the synthesis of biodiesel, and are discussed below:

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205 **4. 1 Edible plant oils**

Soybean oil,⁴² sunflower oil,⁴³ rapeseed oil,⁴⁴ and palm oil⁴⁵ are widely utilized as a biodiesel feedstock in numerous nations, for example, Argentina, Brazil, Indonesia, Europe, US, Malaysia etc. At present, an estimated 95 % of the worlds' total biodiesel is produced from sunflower oil, rapeseed oil, and palm oil.⁴⁶ Various types of edible oils exploited as feedstocks for the production of biodiesel are recorded in Table 2.

211

212 Table 2. Different forms of edible oils utilized to	o produce biodiesel.
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No.	Edible of	il for	Plant source	The botanical name of the
	biodiesel pro	duction		plant source
1	Sunflower oil	1	Sunflower	Helianthus annuus
2	Rapeseed oil		Rape	Brassica napus
3	Soybean oil		Soybean	Glycine max
4	Palm oil		Mesocarp of oil palm	Elaeis guineensis
5	Coconut oil		Coconut	Cocos nucifera

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214 **4. 2 Non-edible plant oils**

Recently, non-edible plant oils have been increasingly considered as another promising 215 potential feedstock for biodiesel attributable to their high oil content and low cost. In addition, 216 unlike edible oils, it does not pose 'food versus fuel' problem as they can be grown in barren 217 and arid regions which are not suitable for agriculture. Further, non-edible oil plants can grow 218 under harsh conditions and hardly need any attention; thus, reducing the cost involved in 219 cultivation and potentially reduced the cost of biodiesel.^{47,48} Some of the commonly 220 221 investigated non-edible plant oils for biodiesel production include Jatropha curcas, Pongamia glabra (Karanja), Madhuca indica (Mahua), Azadirachta indica (neem), Moringa oleifera 222 223 (moringa seed), Calophyllum inophyllum, Salvadora oleoides (Pilu), Nicotiana tabacum (tobacco), cottonseed oil, Eruca Sativa Gars, terebinth, rubber seed oil, desert date, Acrocomia 224 225 aculeate (macaúba), Crambe abyssinica (hochst), linseed oil, rubber seed oil, Sapium sebiferum (chinese tallow), Sapindus mukorossi (soapnut), Euphorbia tirucalli (milk bush), 226 Calophyllum inophyllum (polangafish oil, Jojoba, leather pre-fleshings, apricot seed, Pistacia 227 chinensis Bunge Seed, sal oil, Moringa oleifera and croton megalo-carpus. Amongst all these 228 oil plants, Jatropha curcas, Pongamia glabra (karanja), Madhuca indica (Mahua), Azadirachta 229 indica (neem) are commercially available and most largely used in biodiesel production.⁴⁹ 230

232 4. 3 Waste cooking oil

Biodiesel production from WCO can partially substitute fossil fuels as well as can solve 233 the energy crisis and environmental pollution. Moreover, WCO is cheaper than fresh vegetable 234 oils, consequently, lessen the expense incurred for biodiesel synthesis. WCO can be grouped 235 into two classifications based on their FFA content if the FFA content is >15 %, then it is called 236 brown grease, otherwise, it is named 'yellow grease'. Annually billion tons of WCO is 237 generated throughout the world. In EU, it is estimated that around 0.7-1 MT WCO were 238 collected per year. Among 80,000 tons of WCO, around 65,000 tons were collected from UK 239 240 only, basically originating from commercial restaurants and food processing industries. Therefore, disposal of WCO is a major concern which otherwise contaminates water and 241 environment at large. Although some portions of WCO oil were used in the production of soap, 242 major parts of WCO were usually dumped into the river and landfills. In the light of this, the 243 production of biodiesel from WCO not only reduced the cost of biodiesel but also resolved the 244 disposal problem of WCO and minimized environmental pollution. 245

246

247 **4. 4 Animal fats**

Animal fats are another feedstock for biodiesel production that have the potential to reduce the cost of biodiesel. This type of feedstock includes lard, tallow and chicken fat. However, due to the presence of a high quantity of saturated fatty acids, it has some shortcomings both in chemical and physical properties such as poor cloud point, poor pour point and so forth. At the same time, its high saturation level has various advantages such as high cetane number, high oxidation stability etc. Moreover, animal fats are more favourable biodiesel feedstocks as compared to vegetable oils due to their low price.

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256 **4. 5 Algal oil**

Currently, microalgae are viewed as one of the most promising feedstocks for the industrial-scale synthesis of biodiesel. Biodiesel production from algal oil is highly sustainable as several strains of microalgae can double in size within hours; thereby have the capacity to create a large number of litres of biodiesel per hectare every year.⁵⁰ Additionally, as several microalgal strains can be grown on non-arable land in a saline water medium, their mass cultivating doesn't compete with food production.

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266 5. Characterization of catalysts and biodiesel

Several analytical techniques are employed to characterize both catalysts and FAME 267 produced. Each analytical techniques will be discussed in the upcoming sections as and when 268 relevant. As a preliminary study, Fourier transform infrared spectroscopy (FT-IR) is usually 269 employed to detect the presence of various functional groups in the catalyst, while X-ray 270 diffraction (XRD) can be employed to investigate the crystallinity and qualitative detection of 271 elements present in the catalyst. The surface morphology, particle size and the structure of the 272 catalysts can be investigated using Scanning electron microscopy (SEM) and Transmission 273 274 electron microscopy (TEM). The chemical compositions are investigated using Energydispersive X-ray spectroscopy (EDX). X-ray fluorescence (XRF) is commonly used for 275 quantitative detection of metal oxides and X-ray photoelectron spectroscopy (XPS) analyses 276 are routinely performed for the quantitative measurement of the elements present in the catalyst 277 and also provide the chemical state information of the catalyst. Surface area, pore volume and 278 pore diameter are usually measured by Brunauer-Emmett-Teller (BET) analysis, whereas the 279 thermal stability of the catalysts is analyzed using thermogravimetric analysis (TGA). The 280 acidity, as well as basicity of the catalysts, are usually investigated using NH₃ and CO₂ 281 temperature-programmed desorption (TPD) analyses. In addition, basicity and acidity of the 282 283 catalyst can be visualized by Hammett indicators tests and acid-base titration methods. Valuable information about the degree of carbonization and/or aromatization of carbonaceous 284 material used as a catalyst can be obtained using solid-state magic-angle spin-nuclear magnetic 285 resonance (MAS NMR). Likewise, the successful conversion of biodiesel feedstocks to FAME 286 287 is confirmed using different analytical techniques. Usually, NMR analysis is used as a confirmation tool to identify the formation of FAME. Despite not common, FT-IR analysis can 288 also be used to identify the FAME formation. The chemical components of FAME along with 289 their respective percentage are usually identified using gas chromatography-mass spectroscopy 290 291 (GC-MS) technique. In addition, ¹H NMR spectra can be used to give concrete information about the purity of FAME and percentage conversion of vegetable oil to FAME using Knothe 292 293 and Kenar equation (1).

% Conversion = 100 X
$$\frac{2A_{Me}}{3A_{CH_2}}$$
 (1)

Here, A_{Me} and A_{CH_2} are the integration values of methoxy protons and methylene protons of FAME respectively.

298 6. Homogeneous catalyst:

The homogeneous catalysts utilized for the transesterification reaction are classified into two groups such as i) base catalysts, for example, NaOH and KOH and ii) acid catalysts such as sulphuric, sulphonic, hydrofluoric, and hydrochloric acids.

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6. 1 Base catalyst:

Homogeneous base catalysts are most widely investigated in the transesterification of 304 vegetable oil to FAME as they are cheap and easily accessible. Till date, several homogenous 305 306 base catalysts have been utilized for the synthesis of FAME e.g., KOH, NaOH, NaOCH₃ etc. as shown in Table 3. The uses of NaOH and KOH as catalyst showed excellent catalytic 307 activities towards biodiesel production such as minimum reaction time, high biodiesel yield 308 and occurred at ambient temperature and pressure. However, this process has some certain 309 limitations like water is formed as a byproduct, which reduces biodiesel yield. Other than KOH 310 and NaOH, sodium methoxide and potassium methoxide gives better biodiesel performance as 311 water is not formed in these processes. An alkaline catalyst is not suitable for transesterification 312 of vegetable oils with high FFA content (>2 wt. %). However, it is fit for refined vegetable 313 oils with low FFA content (ranging from less than 0.5 wt. % to less than 2 wt. %). 314

Dmytryshyn et al. ⁵¹ examined the transesterification of various vegetable oil such as 315 canola oil, green seed canola oil from heat-harmed seeds, handled waste fryer oil and natural 316 317 waste fryer oil with methanol to afford FAME using KOH catalyst, and reported a biodiesel yield of 51-87 % under the optimum reaction conditions. In another study, KOH was exploited 318 319 to convert crude rubber oil and palm oil mixture to biodiesel in 98 % yield under the optimum reaction conditions. The vegetable oil was esterified using acid catalyst prior to a base-320 catalyzed transesterification process, to get low FFA content vegetable oil.⁵² Similarly, KOH 321 was utilized as a catalyst for the transformation of soybean oil to FAME in 96 % yield.⁵³ 322 Roselle oil³⁴ rapeseed oil, ⁵⁴ frying oil, ^{55,56} used olive oil,⁵⁷ palm kernel ⁵⁸ and duck tallow ⁵⁹ 323 were also successfully transesterified to FAME using KOH catalyst. Karmee et al. ⁶⁰ reported 324 the transesterification of *Pongamia pinnata* to FAME in 92 % conversion using base catalyst 325 KOH. Interestingly, the utilization of tetrahydrofuran (THF) as a co-solvent increased the 326 conversion to 95 %. 327

Meng *et al.*²³ detailed an exceptionally high activity of NaOH towards biodiesel production from WCO with high FFA in 89.8 % conversion under the optimized reaction settings. The high FFA substance of WCO was reduced by a pre-esterification process with sulphuric acid. Similarly, waste cooking/frying oil,^{61,62} canola oil,⁶³ sunflower oil,⁶⁴ palm oil⁶⁵ and cotton seed oil⁶⁶ were converted to biodiesel using NaOH as a homogeneous catalyst. Furthermore, NaOCH₃^{67, 68} was evaluated as a catalyst for transesterification of rice bran oil to FAME by Rashid *et al.*⁶⁷ where 83.3 % biodiesel yield was observed in 60 min under the optimum reaction conditions.

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Table 3: Distinctive homogeneous base catalysts utilized for biodiesel production.

No.	Catalyst	Feedstock	^a Conditions	Yield (%)	Ref.
1.	КОН	Vegetable oil	6:1, 1, 25, 40	51-87	51
2.	КОН	Crude rubber/palm oil	8:1, 2, 55, 300	98	52
3.	КОН	Soybean oil	6:1, 1, 60, 60	-96	53
4.	КОН	Roselle oil	8:1, 1.5, 60, 60	99.4	36
5.	КОН	Rapeseed	6:1, 1, 65, 120	95-96	54
6.	КОН	Frying oil	12:1, 1, 60, 120	72.5	55
7.	KOH	Waste Frying oil	6:1, 1, 65, 60	96.15	56
8.	КОН	Used olive oil	12:1, 1.26, 25, 90	94	57
9.	КОН	Palm kernel	6:1, 1, 60, 60	96	58
10.	КОН	Duck tallow	6:1, 1, 65, 180	83.6	59
11.	КОН	Pongamia pinnata	10:1, 1, 60, 90	92^{b}	60
12.	NaOH	Waste cooking oil	6:1, 1, 50, 90	89.8 ^b	23
13.	NaOH	Waste frying oil	4.8:1, 0.6, 65, 60	98	61
14.	NaOH	Waste frying oil	7.5:1, 0.5, 50, 30	96	62
15.	NaOH	Canola oil	6:1, 1, 45, 15	98	63
16.	NaOH	Sunflower	6:1, 1, 60, 120	97.1	64
17.	NaOH	Refined palm oil	6:1, 1, 60, 30	95	65
18.	NaOH	Cotton seed oil	6:1, 1, 60, 60	-97	66
19.	NaOCH ₃	Soybean oil	6:1, 0.6, 60, 60	-97	53
20.	NaOCH ₃	Rice bran	7.5:1, 0.88, 55, 60	83.3	67
21.	NaOCH ₃	Waste cooking oil	6:1, 0.75, 65, 90	96.6	68

^aMethanol-to-oil (M/O) molar ratio, catalyst loading (wt. %), temperature (°C), reaction time

339 (min).

340 ^{*b*}Conversion

341

342 6. 2 Acid catalyst:

Base catalysts are usually preferred over acid catalysts as they are more reactive and 343 low cost. However, base catalysts may react with FFA present in the feedstock during 344 transesterification, bringing about soap formation by saponification, which may consume the 345 catalyst and diminish its reactivity. Meanwhile, acidic catalyst is neutral to the FFA and 346 henceforth shows better outcomes for transesterification or esterification of vegetable oils or 347 fats having a high amount of FFA (≥ 2 wt. %). Generally, acid catalysts are utilized to bring 348 down the FFA content in WCO and animal fats by means of esterification prior to 349 transesterification using base catalyst⁵. Several acids such as H_2SO_4 , HCl, H_3PO_4 and 350 sulfonated acids were mostly utilized for the (trans)esterification of vegetable oils.³⁶ However, 351 acid-catalyzed biodiesel production has some major limitations such as slow reaction rate 352 (4000 times slower than the rate of base-catalyzed transesterification) and require high alcohol 353 to oil molar ratio.^{69–71} Moreover, it has environmental and corrosive related problems.⁶⁹ 354 Because of these demerits, acid-catalyzed biodiesel synthesis is not very popular and is less 355 examined. Some reported literature of acid-catalyzed biodiesel production and their results are 356 357 listed in Table 4.

Wang et al.⁷⁰ examined the biodiesel synthesis from WCO and reported a 90 % yield.. 358 Moreover, Miao et al.⁷² examined the conversion of soybean oil to biodiesel using 359 trifluoroacetic acid catalyst and reported 98.4 % biodiesel yield at optimal reaction conditions 360 . Similarly, various edible/non-edible oils such as WCO,⁷³ soybean oil,⁷¹ zanthoxylum 361 bungeanum⁷⁴ and tobacco seed oil⁷⁵ were used for biodiesel production using sulfuric acid. 362 Moreover, trifluoroacetic acid was utilized as a homogeneous acid catalyst for the 363 esterification/transesterification of soybean oil to biodiesel.⁷² The catalyst brought about a high 364 biodiesel yield of 98.4 % under the optimum reaction conditions. From the above discussion, 365 it was observed that an acid-catalyzed esterification/transesterification reactions usually 366 require drastic reaction conditions such as high M/O molar ratio, catalyst loading, temperature 367 368 and long reaction time as compared to base-catalyzed transesterification reactions.

Table 4: Different acidic homogeneous catalysts utilized for biodiesel synthesis.

No.	Catalyst	Feedstock	^a Conditions	Yield (%)	Ref.
1.	H_2SO_4	Chicken/mutton	30:1, 1.25/2.5,	99.01±0.71/	25
		tallow	50/60, 1440	93.21±5.07	
2.	H_2SO_4	WCO	20:1, 4, 95, 600	90	70
3.	H_2SO_4	Used frying oil	3.6:1, 0.1, 65, 40	79.3	73

4.	H_2SO_4	Soybean oil	6:1, 3, 60, 2880	98	71
5.	H_2SO_4	Zanthoxylum	24:1, 2, 60, 80	98	74
		bungeanum			
6.	H_2SO_4	Tobacco seed oil	18:1, 1, 60, 25	91	75
7.	$C_2HF_3O_2$	Soybean oil	20:1, 2 M, 120,	98.4	72
			300		

- ^aMethanol-to-oil molar ratio, catalyst loading (wt. %), temperature ($^{\circ}$ C), reaction time (min).
- 372

373 7. Heterogeneous catalysts

374 Although homogeneous catalyst has its own advantages such as high reactivity and low 375 cost, its utilization in the production of biodiesel is accompanied by several shortfalls such as 376 low quality of glycerol produced, the catalyst cannot be regenerated and the lengthy process involves in the purification of biodiesel; thereby makes the whole process labour-intensive and 377 378 uneconomical. Hence, in recent years, the heterogeneous catalyst has attracted immense attention for biodiesel production as it can be tailored to match specific requirement, easily 379 380 recovered and reused for several cycles of catalytic reaction, thereby potentially bring down the labour involved and cost of biodiesel. 381

Unlike homogeneous catalysts, heterogeneous catalysts mostly appear in a solid form, thus the reaction mixture and the catalyst are in a different phase. In heterogeneous catalyzed reactions, the catalyst surface is the main site for reaction to occurs.⁷⁶ The following advantages of utilizing a solid catalyst in transesterification make the process green: i) Catalyst can be reused, ii) very minimal amount of wastewater generated during the process, iii) glycerol separation from the final mixture (glycerol, biodiesel and catalyst) is much easier, iv) high purity glycerol is obtained.

Heterogeneous catalysts have several advantages over a homogeneous catalyst such as 389 simple separation, recyclability and reusability. Moreover, solid catalysts are eco-friendly, less 390 toxic, minimum corrosion and reduced energy intake. Thus, solid catalysts provide an efficient 391 and economical pathway for biodiesel production.^{12,77,78} Heterogeneous or solid catalysts can 392 393 be grouped into two categories: i) basic and ii) acidic heterogeneous catalysts. Nowadays, researchers have developed several heterogeneous catalysts, which can promote esterification 394 395 and transesterification reactions simultaneously in one reaction vessel (one-pot). These type of 396 catalysts are mostly utilized for biodiesel synthesis from the vegetable oils or animal fats 397 having a high amount of FFA without the requirement of additional pretreatment step to reduce the FFA content.¹² 398

399 7.1 Base catalysts:

In recent years, basic heterogeneous catalysts have been most widely investigated as it 400 can overcome the constraints associated with homogeneous basic catalysts and shows excellent 401 catalytic activity under mild reaction conditions. However, these catalysts are suitable only for 402 biodiesel feedstock with low FFA content, otherwise, catalysts will react with the FFA to 403 produce soap by means of saponification reaction, which make separation of biodiesel from 404 glycerol tedious, thereby diminished biodiesel yield. Several solid base catalysts reported in 405 the literature such as alkaline metal oxide, transition metal oxides, mixed metal oxides, 406 407 hydrotalcites, zeolites, biomass-based catalyst are discussed comprehensively in this section.

408

409 **7.1.1 Alkaline earth metal oxides:**

Oxides of alkaline earth metal are one of the most widely studied catalysts for biodiesel 410 synthesis due to their insolubility in methanol and low toxicities. The basicity of alkaline earth 411 metal oxides follows the order: MgO < CaO < SrO < BaO. MgO is almost inactive towards the 412 transesterification reaction.^{79,80} Among all alkaline earth metal oxides, CaO is most widely 413 utilized in FAME production as it is highly basic, insoluble in alcohol, non-toxic, cheap and 414 easily available.⁸¹ However, it is very sensitive to FFA content and forms undesirable 415 byproducts *via* saponification and also lost its activity in the process.⁸² Despite its high activity, 416 SrO is less studied in transesterification reactions as it is very sensitive to the atmospheric 417 moisture and reacts with CO₂ and water to form SrCO₃ and Sr(OH)₂. Table 5 shows the activity 418 of various alkaline metal oxide towards biodiesel production. 419

Kouzu et al.⁸² examined the transesterification of soybean oil using CaO catalyst and 420 reported a high biodiesel yield of 95 % under the optimized reaction conditions. Granados et 421 al. ⁸³ found that CaO calcined at 700 °C showed very high activity towards biodiesel production 422 from sunflower oil and attained 94 % biodiesel yield. Furthermore, the transesterification of 423 rapeseed oil was reported by Kawashima et al.⁸⁴ where CaO was pretreated with methanol to 424 form Ca(OCH₃), which acted as an initiator for the transesterification reaction. A high biodiesel 425 yield of 90 % was observed using the optimized reaction conditions. In another work, SrO 426 catalyzed transesterification of soybean oil has been reported by Liu et al.85 The catalyst 427 showed excellent activity with a high yield of 95 % at 70 °C and 30 min time. The catalyst is 428 highly stable and can be reused for 10 successive cycles. 429

Ultrasonic-assisted biodiesel synthesis from palm oil was reported using diverse metal
 oxides such as CaO, BaO and SrO.⁸⁶ The activity of the catalyst in ultrasonic-assisted biodiesel
 synthesis was compared with the traditional magnetic stirring process and found that ultrasonic

process showed 95.2 % of yield using BaO within 60 min reaction time, which otherwise take 433 3-4 h in conventional stirring process. Similarly, ultrasonic-assisted transesterification using 434 CaO and SrO resulted in an increase in biodiesel yield from 5.5 % to 77.3 % and 48.2 % to 435 95.2 % respectively. These findings show the advantages of using ultrasonication in the field 436 of chemical synthesis particularly in the field of biodiesel synthesis. The authors also 437 investigated the influence of ultrasonic amplitude on the biodiesel synthesis from palm oil and 438 observed that 50 % ultrasonic amplitude displayed the best result in terms of biodiesel yield. 439 Catalyst reusability test revealed that the catalytic activity of BaO decreased drastically, 440 441 especially in the ultrasonic process during the reusability test, which was mainly due to catalyst leaching. The reaction set-up is depicted in Figure 3. 442





Figure 3: Schematic portrayal of experimental set up for ultrasonic-assisted transesterification reaction. Reproduced from ref. [86].

444

445 Table 5: Different alkaline earth metal oxide catalyzed biodiesel production under various446 reaction conditions.

No.	Catalyst	Feedstock	^a Conditions	Yield (%)	Ref.
1	CaO	Soybean oil	12:1, 8, 65, 180	95	82
2	CaO	Sunflower oil	13:1, 3, 60, 120	94	83
3	CaO	Rapeseed oil	3.8:1, 0.7, 60, 160	90	84
4	SrO	Soybean oil	6:1, 3, 70, 30	95	85
5	BaO	Palm oil	9:1, 3, 65, 60	95.2	86

⁴⁴⁷ ^{*a*}Methanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).

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449 **7.1.2 Transition metal oxides:**

Despite the high reactivity of alkaline earth metal oxides, they have some serious 450 drawbacks such as low reusability and high sensitivity towards moisture that reduced their 451 catalytic efficacy. To overcome these inherent drawbacks, metal oxides of Zn, Ti, Zr and Zn 452 are widely investigated in transesterification reactions as they are easily available, highly stable 453 and showed excellent catalytic activities.^{87–89} To date, numerous transition metal oxide-based 454 catalysts have been reported in the field of biodiesel synthesis from vegetable oil as depicted 455 in Table 6. da Silva et al. 90 reported Cu(II) and Co(II) impregnated on chitosan catalyst for 456 FAME synthesis from soybean oil. The adsorption process for Cu(II) on chitosan is better than 457 Co(II). However, Co(II)@chitosan showed higher biodiesel yield (94.01 %) as compared to 458 Cu(II)@chitosan (88.82 %) using the optimal reaction conditions. In another work, Jitputti et 459 al. ⁸⁷ investigated ZrO₂, ZnO, SO₄²⁻/SnO₂, SO₄²⁻/ZrO₂, KNO₃/KL zeolite and KNO₃/ZrO₂ for 460 the FAME synthesis from the crude palm kernel oil and crude coconut oil, and found that 461 SO_4^{2-}/ZrO_2 catalyst displays the highest reactivity for both the oils with biodiesel yield of 90.30 462 % and 86.30 % respectively. The decreasing order of the catalyst activity towards biodiesel 463 synthesis from crude kernel oil is $SO_4^{2-}/ZrO_2 > SO_4^{2-}/SnO_2 > ZnO > KNO_3/ZrO_2 > KNO_3/KL$ 464 zeolite > ZrO_2 and for the crude coconut oil is $SO_4^{2-}/ZrO_2 > SO_4^{2-}/SnO_2 > ZnO > KNO_3/KL$ 465 zeolite > $KNO_3/ZrO_2 > ZrO_2$. 466

Meanwhile, Baskar et al.91 used Mn-doped ZnO nanomaterial for the conversion of 467 Mahua oil to biodiesel and observed that catalyst calcined at 600 °C showed highest biodiesel 468 yield of 97 % under the optimum reaction conditions. The kinetic investigation of the reaction 469 revealed that 181.91 kJ/mol activation energy is necessary for biodiesel synthesis from Mahua 470 471 oil utilizing Mn-doped ZnO catalyst. The prepared Mn-doped ZnO catalyst was seen as a cluster and spherical in shape as depicted in Figure 4 A. FI-TR analysis was performed to 472 confirm the formation of biodiesel. Absorption bands at 1744 and 1703 cm⁻¹ demonstrated CO 473 stretching of methyl esters in Mahua oil and biodiesel respectively. The main spectrum region 474 that allows for chemical discrimination between Mahua oil and produced biodiesel is in the 475 range 1500-900 cm⁻¹ also called known as fingerprint region. Figure 4B revealed the symmetric 476 and asymmetric stretching of alkyl regions at 1376, 1463, 2852, 2922 cm⁻¹ and CO group of 477 lactones and esters at 1735 cm⁻¹. Moreover, the stretching band of CO group of typical esters 478 at around 1703 cm⁻¹ was observed in Figure 4C. In light of this FT-IR bands, the product 479 480 obtained after transesterification of Mahua oil using Mn-doped Zno catalyst was confirmed as biodiesel. 481

482	Na ₂ MoO ₄ has been synthesized and investigated as a catalyst in transesterification of
483	soybean oil by Nakagaki et al.92 The catalyst displayed high activity towards the
484	transesterification reaction and afforded biodiesel yield of 95.6 %. The high reactivity of the
485	catalyst is due to the acid sites of Mo(VI), which can polarize O-H bond easily.
486	Correspondingly, Serio et al.93 also reported the high reactivity of vanadyl phosphate-based
487	catalyst in the biodiesel synthesis from soybean oil. Regardless of the low surface area, the
488	high reactivity of the catalyst is attributed to the structural/surface morphologies. Biodiesel
489	yield of ≥ 88 % was recorded using the optimal reaction conditions. The dehydrated product of
490	the catalyst VOPO ₄ .2H ₂ O can be converted to VOPO ₄ simply by calcination at 400-500 °C.
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Figure 4: SEM image (A) and FT-IR spectrum (B, C) of Mn-doped ZnO nanomaterial. Reproduced from ref. [92].

511	Table 6: Various transition	metal oxide catalyzed	biodiesel production	under different reaction

512 conditions.

No.	Catalyst	Feedstocks	^a Conditions	Yield (%)	Ref.
1	Cu(II)@chitosan	Soybean oil	1:5 ^c , 2, 70, 180	88.82	90
2	Co(II)@chitosan	Soybean oil	1:5 ^c , 2, 70, 180	94.01	90
3	SO ₄ ²⁻ /ZrO ₂	Crude palm	6:1, 3, 200, 60	90.30	87
		kernel oil			
4	SO ₄ ²⁻ /ZrO ₂	Crude coconut	6:1, 3, 200, 60	86.30	87
		oil			
5	Mn doped ZnO	Mahua oil	7:1, 8, 50, 50	97	91
6	Na ₂ MoO ₄	Soybean oil	54:1, 3, 120, 180	95.6	92
7	Vanadyl phosphate	Soybean oil	0.88:2, 0.5, 180, 60	≥88	93

⁵¹³ ^{*a*}Methanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).

514 ^{*c*}w/w

515 **7.1.3 Zeolites**

Zeolites are crystalline aluminosilicates that possess microporous structure.⁹⁴ It can 516 exist in different structural morphology depending on their synthesis process and reaction 517 conditions such as Si/Al molar ratio, pour sizes and proton exchange levels. The wide 518 opportunity for structural modification of zeolites makes them an excellent catalyst for various 519 520 acid-base reactions. Recently, zeolites are intensively investigated in the field of biodiesel 521 production due to their shape selectivity and acidic character. Normally, zeolites are moderately active for the esterification reaction, however, by increasing the pore size and 522 523 varying the Si/Al ratio, the catalytic properties can be improved. Moreover, zeolites can incorporate various metal ions such as Na⁺, K⁺, Mg²⁺ etc., which are mainly responsible for its 524 basic nature.⁹⁵ Table 7 shows various reported zeolite catalysts employed in biodiesel 525 synthesis. 526

In 2007, NaX zeolite loaded with various concentration of KOH was synthesized and reported as a catalyst in FAME production from soybean oil.⁹⁶ Catalyst loaded with 10 % KOH followed by heating at 393 K for 3 h gave the best result with 85.6 % yield under the optimized reaction conditions. Shu *et al.*⁹⁷ prepared La/zeolite beta using La(NO₃)₃ as a precursor *via* ion exchange technique and exploited in FAME production from soybean oil. They reported that La/zeolite beta has higher stability and catalytic activity towards FAME production compared to zeolite beta catalyst. A yield of 48.9 % was obtained using La/zeolite beta under the

optimized reaction conditions such as 14.5:1 M/O molar ratio, 0.011 wt. % catalyst loading, 534 60°C and 4 h time. In the year 2008, Ramos et al. 98 studied three zeolites such as mordenite, 535 beta and X for the conversion of sunflower oil biodiesel. They examined the effect of different 536 loaded/stacked metals on such zeolites. Zeolite X showed the best catalytic activity as it has a 537 higher number of super basic sites which is absent in other zeolites. Effect of binder, sodium 538 bentonite, on the catalytic reactivity of such zeolites was tested, where X zeolite was 539 agglomerated and thus catalytic activity reduced slightly. A high yield of 93.5 % and 95.1 % 540 of FAME was obtained at 60 °C with and without binder, respectively. In another report, Wu 541 et al.⁹⁹ synthesized a series of CaO supported on zeolites such as NaY, KL and NaZSM-5 via 542 microwave irradiation and utilized in biodiesel synthesis from soybean oil. They reported that 543 supported CaO showed better result compared to the naked CaO as supported catalyst have a 544 high surface area, porosity and basic strength. Accordingly, the best result was exhibited by 545 NaY supported CaO (30 % CaO loaded on NaY) under the optimized reaction conditions. 546

Strontium nanocatalyst supported on ZSM-5 by incipient wetness impregnation method 547 was prepared and applied in biodiesel synthesis from sunflower oil.¹⁰⁰ The authors reported the 548 effect of calcination temperature and Sr/ZSM-5, Ba-Sr/ZSM-5 mass ratios. Ba-Sr/ZSM-5 (Ba 549 4 wt. % to the Sr weight and Sr 6 wt. % to the ZSM-5 weight exhibited the best performance 550 with 87.7 % yield under optimal conditions. In the meantime, Narkhede et al.¹⁰¹ synthesized a 551 series of 12-tungstosilicic acid, SiW₁₂ (10-40 wt. %) impregnated on zeolite H β and applied it 552 in biodiesel synthesis from soybean oil. Interestingly, SEM image of the 30 % SiW₁₂/H β 553 (Figure 5b) is similar with the pure zeolite H β (Figure 5a) and revealed that framework 554 structure of H β was retained even after the impregnation of SiW₁₂ and suggested that SiW₁₂ 555 was homogeneously distributed in the framework structure of H β zeolite. They reported a 95 556 557 % yield of FAME under the optimized reaction conditions.



Figure 5: SEM micrographs of (a) H β and (b) 30 % SiW₁₂/H β . Reproduced from ref 101.

In 2012, Babajide et al.¹⁰² synthesized a zeolite derived from fly ash and then ion-560 exchanged with K to form FA/K-X zeolite, which was then applied in biodiesel synthesis from 561 562 sunflower oil. They reported a high yield of 83.53 % under the optimized reaction conditions. Similarly, Manique et al.¹⁰³ prepared zeolite (sodalite) derived from coal fly ash via the 563 564 hydrothermal process and utilized in biodiesel synthesis from soybean oil. The developed sodalite has a definite surface area of 10 m²/g. They also reported a maximum conversion of 565 95.5 % soybean oil using the optimized reaction conditions . Recently, Al-Jammal et al.¹⁰⁴ 566 567 prepared zeolite derived from zeolite tuft followed by impregnation of a series of KOH (1-6 M) and heated at 80 °C for 4 h to form KOH/zeolite catalyst and finally utilized in biodiesel 568 synthesis from waste sunflower oil. The catalyst (1-4 M) KOH/zeolite exhibited a biodiesel 569 yield of 96.7 % under the reaction conditions: 11.5:1 M/O molar ratio, catalyst amount of 6 wt. 570 % w.r.t. oil, 50 °C temperature and reaction time of 2 h. 571

In the same vein, Du et al.¹⁰⁵ developed La₂O₃ impregnated on NaY zeolite catalyst 572 having a spherical shape with 3-5 mm size and utilized it in biodiesel synthesis from castor oil. 573 In addition, they explored the impact of calcination temperature in the range of 600-1000 °C 574 on biodiesel yield and observed that the catalyst calcined at 800 °C showed the best result. 575 They also revealed that the incorporation of surfactant improved the dispersion of La₂O₃ and 576 577 pore size of zeolite. The XRD pattern of the pure zeolite NaY and the catalyst La₂O₃/NaY zeolite calcined in the temperature range of 600-1000 °C is displayed in Figure 6. The XRD 578 pattern of the pure zeolite (Figure 6a) and the catalyst calcined at 600 °C (Figure 6b) and 800 579 °C (Figure 6c) are almost same and revealed that the crystallinity of the zeolite NaY does not 580 581 change upon the incorporation of La₂O₃. However, on increasing the temperature to 1000 °C,

- the XRD pattern (Figure 11e) showed no characteristic peaks of zeolite, suggests that at highcalcination temperature, the crystallinity of the zeolite is lost.
- 584



Figure 6: XRD pattern of pure zeolite (a), $La_2O_3/NaY-600$ (b), $La_2O_3/NaY-800$ (C), $S-La_2O_3/NaY-800$ (d), $La_2O_3/NaY-1000$ (e). Reproduced from ref. [105].

Table 7: Different zeolite catalyzed FAME production under various reaction conditions.

No.	Catalyst	Feedstocks	^a Conditions	Yield	Ref.
				(%)	
1	KOH@NaX zeolite	Soybean oil	10:1, 3, 65, 480	85.6	96
2	La/zeolite beta	Soybean oil	14.5:1, 0.011, 60,	48.9	97
			240		
3	Zeolite X	Sunflower oil	6:1, 10, 60, 420	95.1	98
4	CaO@NaY zeolite	Soybean oil	9:1, 3, 65, 180	95	99
5	Ba-Sr/ZSM-5	Sunflower oil	9:1, 3, 60, 180	87.7	100
6	$H_4[W_{12}SiO_{40}] @ zeolite \\$	Soybean oil	4:1, 0.2, 65, 480	95	101
	Нβ				
7	FA/K-X zeolite	Sunflower oil	6:1, 3, 60, 480	83.53.	102
8	Sodalite	Soybean oil	12:1, 4, 65, 120	95.5	103
9	KOH/zeolite	Waste	11.5:1, 6, 50, 120	96.7	104
		sunflower oil			

10	La ₂ O ₃ /NaY zeolite	Castor oil	15:1, 10, 70, 50	84.6	105
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^aMethanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).

589 7.1.4 Supported catalyst

To increase the stability and reusability of alkaline earth metal oxides, catalyst support 590 plays an important role as they can reduce the mass transfer limitation and provide a high 591 surface area with high porosity, where metals are anchored.¹⁰⁶ Till now several catalyst support 592 such as alumina, silica, ZnO and ZrO₂ had been proposed for the production of FAME. 593 Alumina is extensively employed as catalyst supports for various basic or acidic compounds 594 exploited as a solid catalyst in esterification/transesterification reactions.¹⁰⁷ Several alumina 595 supported catalysts employed in the transesterification reaction for biodiesel synthesis as 596 shown in Table 8. In 2006, Xie *et al.*¹⁰⁸ investigated the potential of KI loaded on Al₂O₃ support 597 catalyst for biodiesel synthesis from soybean oil. They prepared a series of KI@Al₂O₃ catalysts 598 599 by changing the KI amount and investigated their catalytic activities. They observed that catalyst loaded with 35 % KI and calcined at 773 K showed highest FAME conversion of 96 600 % against all other catalysts under the optimal reaction conditions. In another study, potassium 601 oxide loaded on alumina derived from various potassium salts such as KNO₃, KOH, KF, KI 602 and K₂CO₃ were compared and found that KF@Al₂O₃ showed the best result compared to other 603 604 catalysts, because of the generation of new phase K₂O on the surface of alumina and as result basicity of the catalyst increases.¹⁰⁹ In addition, Ma et al. ¹¹⁰ reported the synthesis of FAME 605 via transesterification of rapeseed oil using K@KOH@Al₂O₃ catalyst. The formation of Al-O-606 K composite enhanced the basicity of the catalyst, thereby catalytic efficiency. They 607 investigated catalytic activity by varying the amount of K and KOH and found that 7.5 and 20 608 wt. % (w.r.t. alumina) of K and KOH, respectively displayed the highest activity with 84.52 % 609 biodiesel yield. Moreover, Chen et al.¹¹¹ reported biodiesel production from soybean oil using 610 $K @\gamma - Al_2O_3$ catalyst in a rotating packed bed (RPB) reactor. The schematic representation of 611 RPB model is displayed in Figure 7. The main advantage of RPB reactor is that it provides 612 efficient mixing of three immiscible reactants such as oil, methanol and the catalyst. A high 613 614 yield of 96.4 % was reported using the optimal reaction conditions.



Figure 7: RPB experimental apparatus utilized for heterogeneously catalyzed transesterification reaction. Components: (1) CSTR reactor; (2) stirrer; (3) thermocouples; (4) sample port; (5) thermostat; (6) control valve; (7) pumps; (8) flow-meter; (9) RPB reactor; (10) stationary liquid distributor; (11) packed-bed rotator; (12) K/g-Al2O3 catalyst; (13) housing case; (14) rotor shaft; (15) motor. Reproduced from ref. [111].

Zhang et al.¹¹² synthesized KOH impregnated modified alumina catalyst for biodiesel 617 synthesis from microalgae oil. Firstly, the alumina was modified with Lanthanum and barium 618 to increase its surface area, possess desired pore volume and pore distribution and finally 619 impregnation of KOH on the modified alumina to form the desired catalyst. They reported that 620 25 % KOH (w.r.t. modified alumina) impregnated on modified alumina and calcined at 550 °C 621 for 4 h showed the best activity towards the transesterification reaction with 97.7 % biodiesel 622 yield under the ideal reaction conditions. Umdu et al.¹¹³ synthesized CaO@Al₂O₃ via the sol-623 gel method and conducted a transesterification reaction of microalgae (Nannochloropsis 624 *oculata*) oil to produce biodiesel. The catalyst has higher reactivity than the bare CaO, which 625 was almost inactive towards transesterification of the desired microalgae. The alumina was 626 loaded with 80 wt. % (w.r.t. Al₂O₃) Ca(NO₃)₂. 4H₂O and calcined at 500 °C for 6 h to form 80 627 wt. % CaO@Al₂O₃ that possessed the highest catalytic activity with 97.5 % biodiesel yield. In 628 addition, Zabeti et al.¹¹⁴ synthesized a CaO@Al₂O₃ catalyst using calcium acetate via 629 calcination at 718 °C for biodiesel synthesis from palm oil. They have used Response Surface 630

Methodology (RSM) in association with Central Composite Design (CCD) to determine the optimum reaction conditions such as M/O molar ratio, catalyst amount, reaction temperature and reaction time. Biodiesel yield of 98.64 % was obtained under the optimum reaction conditions.

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Table 8: Different aluminium supported solid catalyst for biodiesel production.

No.	Catalyst	Feedstock	^a Conditions	Yield (%)	Ref.
1	KI@Al ₂ O ₃	Soybean oil	15:1, 2, 65, 480	96	108
2	K@KOH@Al ₂ O ₃	Rapeseed oil	9:1, 4, 60, 60	84.52	110
3	K@y-Al ₂ O ₃	Soybean oil	24:1, 10.6, 60,60	96.4	111
4	KOH/La-Ba-Al ₂ O ₃	Microalgae	NR, 25, 60, 180	97.7^{b}	112
5	CaO@Al ₂ O ₃	Nannochloropsis	30:1, 2, 50, 240	97.5	113
		oculata			
6	CaO@Al ₂ O ₃	Palm oil	12:1, 6, 65, 300	98.64	114

 a Methanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).

637 ^bConversion

638 NR: Not reported

639

Apart from alumina, there are several materials which are used as catalyst support such 640 as SiO₂, ZrO₂ and activated carbon (AC) (Table 9). In 2010, Samart et al.¹¹⁵ conducted 641 transesterification reaction using CaO impregnated on mesoporous SiO₂ catalyst for FAME 642 production. They also investigated the influence of CaO amount and reported that 15 wt. % 643 644 CaO (w.r.t. SiO₂) loading showed the maximum yield of 95.2 %. In addition, the synthesis of FAME from palm oil using a CaO impregnated on bimodal meso-macroporous SiO₂ support 645 catalyst was reported by Witoon et al.¹¹⁶. They investigated the influence of CaO loading and 646 pellet size on biodiesel conversion and also compared with unimodal SiO₂ supported CaO 647 catalyst. CaO in 40 wt. % CaO@SiO₂ were highly aggregated on the surface of the mesoporous 648 SiO₂ hence increases the surface basicity; while CaO in 30 wt. % CaO@SiO₂ were highly 649 dispersed inside the mesopore of the silica support, accordingly 40 wt. % CaO@SiO₂ showed 650 higher FAME yield compared to 30 wt. % CaO@SiO2. They also reported that the catalyst 651 with pellet size 335 μ m showed a maximum yield of 92.45 %. Moreover, Wu *et al.*¹¹⁷ reported 652 catalysts consisting of three different potassium compounds (KAc, K₂CO₃ and K₂SiO₃) 653 impregnated on mesoporous SiO₂ such as AlSBA-15 and SBA-15 for the production of FAME 654

from JCO. Three potassium salts with different concentration were impregnated on AlSBA-15 and SBA-15 and found that the basicity lies in the order of 35 wt. % $K_2SiO_3@AlSBA-15 > 35$ wt. % $K_2CO_3@AlSBA-15 > 35$ wt. % KAc@AlSBA-15 and thus 30 wt. % K_2SiO_3 showed highest yield of 95.7 % under the optimized reaction conditions.

The concept of AC-based catalyst is an attempt towards the development of a novel 659 alternative to homogeneous alkaline in the form a heterogeneous catalyst. These kinds of 660 catalysts have pulled in a lot of consideration from the scientific community because the uses 661 of carbon as catalysts not only makes them reusable in the production process but also greatly 662 reduce the formation of the soap and increases glycerol purity.¹¹⁸ To date, different kinds of 663 activated carbon-based catalysts have been developed and successfully exploited in biodiesel 664 production, some of them are briefly discussed here (Table 18). Narowska et al.¹¹⁸ proposed 665 the development of a novel carbon-based catalyst to replace alkaline homogeneous catalyst as 666 a solid catalyst which has the potential to be reused multiple times, eliminating various 667 668 limitations associated with other traditional catalysts. In this context, the authors demonstrated the preparation of FAME from corn oil via transesterification utilizing KOH supported on 669 activated carbon catalyst. The result showed that the highest yield (92 wt. %) of FAME was 670 recorded using optimal reaction conditions. These finding indicated that activated carbon-671 672 supported catalysts can be promisingly employed in the transesterification of the waste corn 673 oil using methanol.

Previously, Buasri et al.¹¹⁹ reported calcium oxide impregnated on AC catalyst in the 674 synthesis of highly pure FAME from waste cooking palm oil through continuous 675 transesterification of FFA. After optimization of various reaction, a maximum FAME yield (94 676 %) was accomplished. In another study, Konwar et al.¹²⁰ also synthesized AC-supported 677 calcium oxide from *Turbonilla striatula* shell and further, their applicability as a catalyst has 678 been investigated in biodiesel synthesis from vegetable oil. It was reported that the catalyst 679 680 displayed more than 90% oil conversion under the optimized reaction conditions. Moreover, this approached is economically viable due to easy recoverability of the catalyst. The catalyst 681 was utilized for five progressive reactions cycles with minimum activity loss. 682

Hameed *et al.*¹²¹ examined a solid catalyst KF supported on AC for biodiesel synthesis from WCO. They designed a composite rotatable reactor to optimize the reaction parameters and obtained 83 % methyl ester yield. In 2010, Baroutian *et al.*¹²² studied FAME synthesis in a packed bed membrane reactor (PBMR) from palm oil using a solid catalyst of KOH supported on AC generated from palm shell (Figure 8). They also investigated the impact of reaction parameters using RSM. The highest biodiesel yield of 98.03 % was reported using the catalyst 689 with optimized reaction conditions. In addition, Li *et al.*¹²³ reported in situ synthesis of 690 K₂CO₃@KFA *via* mixing of K₂CO₃ and kraft lignin (KF) succeeded by calcination at 800 °C 691 and utilized the catalyst in biodiesel synthesis from rapeseed oil. They also investigated the 692 influence of reaction parameters on the FAME production and reported that a maximum yield 693 of 99.6 % under the optimized reaction conditions.

Further, Buasri et al.¹²⁴ conducted a synthesis process where a solution of KOH was 694 mixed with activated carbon (AC) originated from coconut shell to form KOH@AC and used 695 this catalyst in biodiesel synthesis from WCO. The authors claimed that the synthesized 696 catalyst has extraordinary catalytic reactivity and showed 86 % biodiesel yield under the 697 optimized reaction conditions. Similarly, Wan et al.¹²⁵ examined a solid base catalyst 698 CaO@AC for FAME synthesis from palm oil. RSM was utilized to investigate the impact of 699 reaction parameters on biodiesel synthesis. A maximum yield of 80.98 % was reported under 700 the optimal reaction conditions and also claimed that the catalyst can retain its activity even 701 after two cycles. Recently, Fadhil et al.¹²⁶ conducted a transesterification reaction of bitter 702 almond oil to produce biodiesel using KAc impregnated on activated carbon originated from 703 the waste of polyethyleneterphathalte. A maximum yield of 93.21 % with high purity was 704 reported. The authors claimed that the catalyst showed excellent reactivity towards biodiesel 705 706 synthesis compared to other reported solid base catalyst as the catalyst showed a very high yield in very low optimal reaction conditions. Moreover, according to the authors, the catalyst 707 708 has great stability as it can be reused for 6 cycles.



Figure 8: Schematic diagram of PBMR for FAME synthesis. Components: (1) palm oil; (2) methanol; (3) crude material siphon; (4) magnetic stirrer; (5) blending vessel; (6) flowing siphon; (7) boiling water flowing; (8) water chiller; (9) wound thermal exchanger; (10) ceramic membrane; (11) pressure check; (12) temperature indicator; (13) methanol recuperation unit; (14) siphon; (15) isolating funnel. Reproduced from ref. [122].

Liu et al.¹²⁷ examined a solid base catalyst KF/CaO/AC calcined at 500 °C for 5 h for 711 the conversion of soybean oil to biodiesel. The authors claimed that the main catalytic role was 712 played by K₂O and KCaF₃ that are present in the catalyst. The catalyst demonstrated high yield 713 of 99.9 % only in 20 min. Nonetheless, they reported that the catalyst is highly sensitive 714 towards water contents in methanol and oleic acid. Therefore it is necessary to use anhydrous 715 oil and methanol to overcome this problem. In conclusion, from all these above-mentioned 716 717 studies a collective inference can be drawn that activated carbon-based catalysts will be the next generation novel alternative to traditionally available catalysts for efficient 718 transesterification of different oils. 719

In the meantime, the application of zinc oxide supported silver nanoparticles (ZnO@Ag
 NPs) as a solid catalyst for the conversion of palm oil to FAME was reported by Laskar *et al.*¹²⁸
 The transformation of palm oil to FAME was confirmed using NMR analysis and 10

components of FAME were identified using GC-MS technique, with methyl octadecanoate
(C18:0) being the major component. Mixture with different ratio of Ag on ZnO were prepared,
where 10 wt. % ZnO@Ag was found to be the most active catalyst producing 96 % FAME
under optimum reaction conditions. In the recent past, Taslim *et al.*¹²⁹ also demonstrated the
efficacy of low-cost AC-based catalyst developed from candlenut shell (an agricultural waste)
through the impregnation of KOH for biodiesel production from WCO. The results obtained
has shown that the yield of biodiesel up to 96.65 % using optimized reaction conditions.

No.	Catalyst	Feedstocks	^a Conditions	Yield	Ref.
				(%)	
1.	CaO/SiO ₂	Soybean oil	16:1, 5, 60, 480	95.2	115
2.	CaO/SiO ₂	Palm oil	12:1, 5, 60, 240	94.15	116
	(bimodal)				
3.	K ₂ SiO ₃ @AlSBA-	Jatropha oil	9:1, 15.30, 60, 180	95.7	117
4.	KOH/AC	Corn oil	3:1, 0.75, 62.5, 60	92	118
5.	CaO/AC	WCO	25:1, NR, 60, 480	94	119
6.	CaO/AC	Vegetable oil	40:111,120, 420	>90	120
7.					
8.	KF/AC	WCO	8.85:1, 3, 175, 60	83	121
9.	KOH/AC	Palm oil	24;1, 30.3, 64.1,	98.03	122
			60		
10.	K ₂ CO ₃ @KFA	Rapeseed oil	15:1, 3, 65, 120	99.6	123
11.	KOH@AC	WCO	25:1, NR, 60, 120	86.3	124
12.	CaO@AC	Palm oil	15:1, 5.5, 190, 81	80.98	125
13.	KAc/AC	Bitter almond	9:1,2.50, 65, 150	93.21	126
		oil			
14.	KF/CaO/AC	Soybean oil	12:1, 2.1, 65, 20	99.9	127
15.	Ag@ZnO	Palm oil	10:1, 10, 60, 60	96	128
16.	KOH/AC	WCO	12:1, 3, 60, 120	96.65	129

Table 9: Different solid supported catalyst for biodiesel synthesis.

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^{*a*}Methanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).

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- 733

735 **7.1.5 Hydrotalcite:**

Recently, hydrotalcites have attracted interest as a solid catalyst in the 736 transesterification reactions due to their tunable properties and excellent performance. They 737 belong to the layered double hydroxide (LDH) family. The general formula of hydrotalcite is 738 $[M_n^{2+} M_m^{3+} (OH)_{2 (n+m)}]^{m+} [A^{x-}]_{m/x}.yH_2O$, where M^{2+} is a divalent metal e.g., Ca^{2+} , Zn^{2+} , Mg^{2+} 739 etc., M^{3+} is a trivalent metal, most frequently Al^{3+} , whereas A^{x-} is an anion with x in the range 740 of 0.1-0.5^{130,131}. Table 10 shows various reported hydrotalcite catalysts employed in the 741 biodiesel synthesis from different feedstocks. Navajas et al.¹³² prepared Mg/Al hydrotalcite 742 with composition within the range of 1.5-5 by co-precipitation method and applied it in the 743 744 conversion of sunflower oil to biodiesel. The basicity of the catalyst increased with the increase in Mg/Al molar ratio and degree of rehydration. They reported a 96 % conversion of oil to 745 FAME (92 % yield) utilizing the rehydrated hydrotalcite under the optimal reaction conditions. 746 Zeng et al.¹³³ reported Mg-Al hydrotalcite with various Mg/Al molar ratio and used 747 them as a heterogeneous catalyst for the transesterification of soybean oil. The hydrotalcite 748 749 calcined at 773 K and 3:1 Mg to Al molar ratio exhibited the highest catalytic activity with 90.5 % conversion of oil. Recently, Ma et al.¹³⁴ investigated a heterogeneous catalyst Mg-Al 750 751 hydrotalcite in the production of biodiesel from WCO. They mentioned that the catalyst with 752 Mg/Al molar ratio 3:1 and calcined at 500 °C have a high surface area, excellent crystallinity

and mesoporous structure, subsequently showed excellent activity. They also reported 95.2 % 753 FAME yield under the optimized reaction condition. In the same manner, Zeng et al.¹³⁵ 754 755 prepared Mg/Al-CO₃ with Mg/Al molar ratio of 4:1 via urea method and compared their structures and catalytic activities with those prepared by co-precipitation for the biodiesel 756 757 synthesis from microalgae oil. They studied the crystal size and surface basicity of all the prepared hydrotalcites and reported that the crystal size of the hydrotalcites prepared using urea 758 759 method is greater than as-synthesized ones. They also reported that the mixed oxide of the hydrotalcite prepared via urea method showed the highest catalytic reactivity with the 760 761 maximum conversion of 90.30 %.

Further, Mg-Al hydrotalcite loaded with 1.5 % K was prepared and used as a catalyst for the synthesis of biodiesel from palm oil.¹³⁶ A maximum 86.6 % yield was reported usingthe optimized reaction conditions. They also studied the effect of the synthesized biodiesel on six types of elastomers such as NBR, HNBR, NBR/PVC, acrylic rubber, co-polymer FKM, and terpolymer FKM, which are commonly found in the fuel system. For testing, the elastomers were immersed in B10 (10 % biodiesel in diesel) and found that only terpolymer FKM and co-

polymer FKM showed a slight change in the properties. Thus, concluded that B10 is compatible 768 with the diesel engines without any modification. In another work, Liu et al.¹³⁷ prepared Zn-769 770 Al hydrotalcite within the temperature range of 413-773 K to form dehydrated Zn-Al 771 hydrotalcite and Zn-Al mixed oxides and used both the catalysts in the transesterification 772 reaction in a fixed-bed reactor. The OH groups in the dehydrated Zn-Al is responsible for the high basicity of the catalyst. However, Mn^+-O^{2-} pairs and isolated O^{2-} anions are the main 773 basic sites in Zn-Al metal oxides. Furthermore, they compared the catalytic activity of both 774 dehydrated Zn-Al HT and Zn-Al oxides and found that the dehydrated HT calcined at 473 K 775 776 showed highest catalytic activity and stability towards biodiesel synthesis with a maximum yield of 76 % at 140 °C for 1 h. Similarly, a heterogeneous base catalyst, KF/Ca-Al was 777 developed for the biodiesel production from palm oil.¹³⁸ The catalyst was prepared from 778 layered double hydroxides of Ca-Al, where the introduction of KF enhanced the catalytic 779 activity. It was observed that 100 wt. % loading of KF decreased particle size of catalyst as 780 shown by the SEM image of KF/Ca-Al (Figure 9). The authors also reported biodiesel yield of 781 97.14 % under the optimized reaction conditions. Besides, biodiesel production from poultry 782 fats was reported by using a solid base catalyst, Mg-Al hydrotalcite.¹³⁹ The influence of 783 calcination temperature for the preparation of catalyst was investigated and disclosed that the 784 785 catalyst calcined at 550 °C showed the maximum catalytic activity. Moreover, the authors detailed that rehydration of the catalyst before the transesterification reaction and preferential 786 787 adsorption of TAGs on the surface of the catalyst reduced the catalytic activity.



Figure 9: SEM image of KF/Ca-Al. Reproduced from ref. [138].

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Helwani *et al.*¹⁴⁰ synthesized Mg-Al hydrotalcite *via* combustion method using
saccharose for biodiesel synthesis from JCO. SEM image of the catalyst calcined at 850 °C
displays a lamellar microstructure with closely packed flakes (Figure 10). The catalyst calcined

at 850 °C and recrystallized with 20 % saccharose fuel showed the best reactivity with 75.2 %
biodiesel conversion under the optimized reaction conditions. A layered double hydroxide of
zinc hydroxide nitrate was also reported for FAME synthesis from palm oil.¹⁴¹ The catalyst
showed excellent reactivity towards the transesterification reaction with 96.5 % biodiesel yield.



Figure 10: SEM image of Mg-Al HT calcined at 850 °C Reproduced from ref. [140].

798

Table 10: Different hydrotalcite catalyzed FAME production under various reactionconditions.

No.	Catalyst	Feedstocks	^a Conditions	Yield	Ref.
				(%)	
1	Mg-Al HT	Sunflower oil	48:1, 2, 60, 480	92	132
2	Mg-Al HT	Soybean oil	6:1, 1.5, 65, 240	90.5	133
3	Mg-Al HT	WCO	6:1, 1.5, 80, 150	95.2	134
4	Mg/Al-CO ₃	Microalgae oil	6.4:1, 1.7, 66, 240	90.3	135
5	K/Mg-Al HT	Palm oil	30:1, 7, 100, 360	86.6	136
6	Zn-Al HT	Soybean oil	26:1, NR, 140, 60	76	137
7	KF/Ca-Al	Palm oil	12:1, 5, 65, 300	97.98	138
8	Mg-Al HT	Poultry fat	30:1, 10, 120, 120	75	139
9	Mg-Al HT	Jatropha oil	30:1, 5, 160, 240	93.4	140
10	$Zn_5(OH)_8(NO_3)_2$	Palm oil	6:1, 2, 140, 120	96.5	141
	.2H ₂ O				

^aMethanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).

802 NR= Not reported

803 7.1.6 Mixed metals oxides:

Mixed metal oxides provide exceptionally fascinating properties especially when each 804 component differs from one another. The basic idea of synthesizing the mixed metal-oxide 805 catalysts is to increase the basic or acid strength, surface area, and stability of these catalysts 806 when compared with single metal oxides. Henceforth, a series of highly efficient, reusable, and 807 stable solid catalysts were prepared. For example, a combination of two metal oxides can show 808 acid-base properties or some unique properties irrespective of their individual properties.¹⁴² 809 The basicity of metals increases as it becomes less electronegative down the group. In the 810 meantime, the highly basic metal oxides those formed with alkaline and alkaline earth metals 811 are usually carbonated in air, and hence are inert. Hence, strong basicity can be achieved only 812 after a high temperature treatment to obtain a carbonate-free metal oxide surface, making the 813 process highly energy demanding.¹⁴³ Interestingly, mixed metal oxides with high reactivity can 814 be obtained at a much lower temperature making it highly demanded in catalysis. To date, 815 several mixed metal oxides have been reported in transesterification reactions and are listed in 816 Table 11. 817

Kawashima *et al.*¹⁴⁴ investigated various calcium-containing catalysts-CaTiO₃, 818 819 CaMnO₃, Ca₂Fe₂O₅, CaZrO₃, and CaO-CeO₂ in the biodiesel production from rapeseed oil. Among these, CaO-CeO₂ showed excellent results (approximately 90 % yield) with high 820 821 stability compared to other calcium-containing heterogeneous catalysts under the optimized reaction conditions. The catalyst can be reused for 7 times with a high yield of >80 % in each 822 time. Sun et al.¹⁴⁵ also prepared La₂O₃ loaded ZrO₂ catalyst by varying La₂O₃ amount from 7-823 28 wt. % and investigated for the synthesis of biodiesel. 21 wt. % La₂O₃ loading on ZrO₂ and 824 825 calcined at 600 °C demonstrated the highest catalytic activity towards biodiesel production from sunflower oil. The authors proposed a model for the preparation of the catalyst, where 826 827 La(NO₃)₃ was impregnated on the surface of ZrO₂ followed by drying to form a film of La(NO₃)₃, which upon calcination forms the La₂O₃/ZrO₂ composite, resulting in a decrease in 828 particle size due to t/m phase transition (Figure 11). A high oil conversion of 96 % and 84.9 829 % FAME yield was observed under optimal reaction conditions. They reported an excellent 830 activity of catalyst prepared by 21 wt. % loaded La₂O₃ and calcined at 600 °C. 831


Figure 11: Proposed model for the solid-state reaction on the catalyst surface. Reproduced from ref. [145].

Wen et al.¹⁴⁶ obtained TiO₂-MgO catalyst via the sol-gel method and employed it in 833 834 the FAME synthesis from WCO. Substitution of Ti to the Mg lattice led to defects in the surface of the catalyst, enhanced both the activity and stability of the catalyst. It was revealed that the 835 catalyst with 1:1 Ti to Mg molar ratio and calcined at 923 K is the most active one in FAME 836 synthesis. Biodiesel yield of 92.3 % was observed utilizing the catalyst MT-1-923 using the 837 838 optimal reaction conditions. Similarly, SrO/SiO₂ and SrO/CaO has been synthesized and their catalytic activity was compared with naked SrO in transesterification of olive oil by Chen et 839 al.¹⁴⁷ Although naked SrO showed very good catalytic activity and afforded 82 % yield in just 840 15 min, biodiesel yield shrunk to 68.9 % when the reaction was performed for 3 h. They 841 842 reported that the reason for the unusual decreased in biodiesel yield was due to reverse reaction between FAME and glycerol, which showed that the catalyst not only catalyzed the forward 843 reaction but also catalyzed the reverse reaction as well. On the contrary, modification of SrO 844 with SiO_2 and CaO provided an excellent activity as well as high stability. They observed that 845 around 95 % conversion was obtained at 65 °C using SrO/SiO₂ and SrO/CaO in 10 and 20 min 846 respectively. However, they reported that on decreasing the reaction temperature to 45 °C, 847 SrO/CaO showed only 20.20 % conversion as compared to SrO/SiO₂, which showed 76.9 % 848 conversion. Thus, SrO/SiO₂ displayed better reactivity towards transesterification of olive oil 849 than SrO/CaO and possessed high tolerance to the water content and FFA of biodiesel 850 851 feedstocks.

In the recent past, Madhuvilakku *et al.*¹⁴⁸ developed TiO₂-ZnO nanocatalyst and utilized in FAME synthesis from palm oil. Arrangement of deformities on the catalyst surface as a result of the substitution of Ti on Zn grid improved the reactivity and stability of the prepared catalyst. They recorded 92 % biodiesel yield was acquired under the optimized reaction conditions. Similarly, a series of ZnO-La₂O₃ catalyst have been examined in the biodiesel synthesis from waste oil by Yan *et al.*¹⁴⁹ Incorporation of La promoted dispersion of ZnO and improved acidic-basic sites, thereby increased catalytic activity towards both

transesterification and esterification reactions. The molar ratio of 3:1 Zn to La showed the 859 highest activity towards biodiesel production. A high 96 % yield was reported under the 860 optimal reaction conditions. The authors also reported that the catalyst can endure FFA and 861 water contents and thus allowed direct conversion of waste oil to FAME. In another work, 862 transesterification of palm kernel oil to produce biodiesel has also been reported using a mixed 863 metal oxide solid base catalyst CaO-ZnO.¹⁵⁰ Upon incorporation of Zn to the CaO phase, the 864 particle size of the catalyst has decreased and has reduced the calcination temperature required 865 for the decomposition of carbonates to its oxides. Lowering of calcination temperature for the 866 867 decomposition of CaCO₃ upon the incorporation of Zn can be explained by particle size reduction coupled with a loss of H₂O and CO₂ from the zinc carbonate. The schematic 868 representation for the decomposition of CaCO₃ and formation of CaO-ZnO mixed metal oxides 869 is displayed in Scheme 3. It is well known that decarbonisation is a reversible process, which 870 mostly depends on atmospheric CO₂, particle size and composition. The dissociation of CO₂ 871 normally occurs in the outer surface (Scheme 3A). Moreover, upon calcination, the evolved 872 CO₂ may form a layer on the surface of the material during the continuous disjunction of inner 873 874 particles, generated a possibility for recarbonation of CaO to CaCO₃ (Scheme 3B). However, incorporation of ZnCO₃ resulted in the formation of voids due to its decomposition to zinc 875 876 oxide. The resulting voids facilitated heat transfer to the interior particles and evaporation of the gaseous compounds. Moreover, due to the small particle size of the CaO-ZnO, the diffusion 877 878 distance of CO₂ decreased, thus calcination temperature also decreased.



Scheme 3: Proposed models for CaCO₃ decomposition to CaO (A) and mixed precipitate of Ca-Zn Reproduced from ref. [150].

Among solid base catalysts, solid ZrO_2 catalysts become popular because of their environmentally benign nature and economic viability for biodiesel production. Till date, different types of ZrO_2 catalysts have been developed for use in biodiesel production. In this line, Su *et al.*¹⁵¹ synthesized microporous solid base MgO-ZrO₂ composites and utilized them as effective heterogeneous catalysts in biodiesel synthesis. They claimed that such microporous catalysts are of great significance as the presence of porous materials in the preparation of these catalysts provided the ability to interact with atoms, ions, molecules.

Recently, Ibrahim et al.¹⁵² examined the influence of different support materials like 888 Al₂O₃, Fe₂O₃, TiO₂ and SiO₂ on physicochemical properties and efficacy of ZrO₂ solid catalysts 889 commonly used in biodiesel synthesis. From the results obtained it was revealed that ZrO₂ 890 supported on SiO₂ showed the highest conversion rate due to comparatively high surface area 891 and a high number of Lewis acid sites. In another study, Faria et al.¹⁵³ developed nanosized 892 893 catalyst mixed metal oxides SiO₂/ZrO₂ catalyst prepared via sol-gel strategy and examined its reactivity in the synthesis of biodiesel from soybean oil. It was observed that this catalyst 894 displayed promising reactivity and gave 96.2 ± 1.4 % biodiesel yield after 3 h of reaction time. 895 In addition, the catalyst can be reused for 6 progressive cycles with little drop in activity. In 896 2008, Albuquerque et al.¹⁵⁴ synthesized MgO-CaO mixed metal oxides with different Mg/M 897 (M = A) or Ca) molar ratios and used it as a highly active catalyst for the transformation of 898

sunflower oil to biodiesel in 92 % yield under the optimized reaction conditions. The highest activity towards the transesterification reaction was found for a bulk Mg:Ca molar ratio of 3.8, whereas bare CaO were found to afford a lower yield of biodiesel under the same reaction conditions. The authors attributed this interesting activity to the higher BET surface area of the MgO-CaO mixed metal oxide ($12.8 \text{ m}^2 \text{ g}^{-1}$), in comparison to CaO ($1.2 \text{ m}^2 \text{ g}^{-1}$).

904

905 **Table 11:** Various mixed metal oxide catalyzed transesterification of vegetable oil.

No.	Catalyst	Feedstocks	^a Conditions	Yield (%)	Ref.
1.	CaO-CeO ₂	Rapeseed oil	6:1, 10, 60, 600	90	144
2.	La ₂ O ₃ /ZrO ₂	Sunflower oil	30:1, 21, 200, 300	84.9	145
3.	TiO ₂ -MgO	WCO	50:1, 10, 160, 360	92.3	146
4.	SrO/SiO ₂	Olive oil	6:1, 5, 65, 10	95	147
5.	SrO/CaO	Olive oil	6:1, 5, 65, 20	95	147
6.	TiO ₂ -ZnO	Palm oil	6:1, 14, 60, 300	92	148
7.	ZnO-La ₂ O ₃	Waste oil	6:1, 2.3, 200, 180	96	149
8.	CaO-ZnO	Palm kernel oil	30:1, 10, 60, 60	>94	150
9.	MgO-ZrO ₂	Soybean oil	20:1, 3, 150, 360	99	151
10.	$ZrO_2@SiO_2$	Stearic acid	120:1, 10, 120, 180	48.6	152
11.	SiO ₂ /ZrO ₂ NP	Soybean oil	6.6:1, 2.8 mmol, 50,	96.2±1.4	153
			180		
12.	MgO-CaO	Sunflower oil	12:1, 2.5, 60, 60	92	154

^aMethanol-to-oil molar ratio, Catalyst loading (wt. %), Temperature (°C), Reaction time (min).
907

908 7.1.7 Biomass-based catalyst

909 In recent year, bio-waste derived heterogeneous catalyst gains significant attention both in the realm of catalysis and biofuel research, and are reviewed by several authors recently ^{155–} 910 ¹⁶⁰The advantages of using waste materials as a catalyst are largely due to their cheap, 911 abundant, non-toxic, ecofriendly, economic, renewable, sustainable and easily availability. 912 Many researchers utilized waste biomass as a catalyst for low FFA oil (edible oil) as well as in 913 high FFA oil (edible and non-edible oils). The biomass includes plant ashes, waste shells, 914 bones, industrial wastes and so forth. Profitably, catalysts derived from waste biomass 915 potentially make biodiesel production highly cost-effective and environmentally benign. 916

919 **7.1.7.2 Waste shells**

920 Despite several chemically synthesized heterogeneous catalysts mentioned earlier show 921 promising and comparatively high biodiesel yield, their synthesis routes are sometimes complicated, expensive, chemically wasteful, time consuming and non-economical. Therefore, 922 with the growing high demand for renewable energy, there is a need to search for an ideal 923 924 heterogeneous catalyst which is easy to synthesize, non-toxic, low cost, widely available, biodegradable and eco-friendly in nature, yet exhibits high catalytic activity in biodiesel 925 926 production. In the light of this, utilization of CaO (derived from high-temperature calcination of waste shells containing CaCO₃) has been a front runner in recent times. The use of waste 927 shells as a source of CaO not only make the whole production of biodiesel sustainable but also 928 solved the problem associated with waste disposal of huge quantities of waste shell generated 929 due to human consumption. 930

931

932 7.1.7.2.1 Eggshell

933 Various eggshell derived heterogeneous catalysts are available for the transformation of edible/non-edible oils to FAME as listed in Table 12. For the first time, CaO originated from 934 chicken eggshell calcined at 1000 °C was utilized for biodiesel synthesis by Wei et al.¹⁶¹ 935 Biodiesel yield greater than 95 % was obtained. They have calcined the eggshell at different 936 937 temperatures from 200 °C to 1000 °C and then tested their efficacy for the transformation of soybean oil to FAME. They observed that those calcined above 800 °C were the most active 938 939 catalysts, where the XRD spectra display a crystalline CaO (Figure 12). Samples calcined at 940 700 °C for 2 h contain CaCO₃ as the principal constituent and CaO as a minor one, hence 941 medium yield (90 %) were obtained. Calcinations below 600 °C did not result in the formation of CaO, hence, low catalytic activity was observed (<30 % biodiesel yield). Hence, CaO in the 942 943 catalyst is the principal basic constituent, which led to the high reactivity of the catalyst. From this experiment, it is suggested that waste shells have to be calcined at a temperature of at least 944 800 °C for 2 h to fully convert CaCO₃ to CaO, a highly basic catalyst. 945

946



Figure 12: XRD patterns of natural eggshell and the materials obtained by calcining natural eggshell in the range of 200 °C– 1000 °C. Reproduced from ref. [161].

In recent years, CaO derived from eggshell has been widely investigated in transformation 949 of various edible/non-edible oils such as soybean oil, $\frac{162-164}{162-164}$ karanja oil, $\frac{165}{160}$ WCO, $\frac{166-175}{160-175}$ palm 950 oil,^{176–179} rapeseed oil,^{180,181} sunflower oil,^{182–185} JCO,¹⁸⁶ microalgae oil,^{187–189} chicken fat,¹⁹⁰ 951 catfish oil,¹⁹¹ Helianthus annuus L oil,¹⁹² cotton oil¹⁹³ and sativa oil¹⁹⁴ for FAME production. 952 In 2014, Niju et al.¹⁷² examined a highly active modified chicken eggshell derived CaO catalyst 953 for the synthesis of FAME from WFO. The authors reported that highly reactive CaO can be 954 955 obtained from eggshells via calcination-hydration-dehydration treatment. While the FAME conversion was only 67.57 % for commercial CaO catalyst, CaO obtained from the eggshell 956 calcined at 900 °C followed by hydration and dehydration at 600 °C (Eggshell-CaO-900-600) 957 gave 94.52 % conversion under the optimized reaction conditions. Calcination followed by 958 959 hydration and dehydration greatly increased the surface area of the eggshell derived CaO as compared to those obtained with the only calcination. The high activity of the modified CaO 960

961 (Eggshell-CaO-900-600) is attributed to the high surface area (8.6401 m²g⁻¹) compared to both 962 commercial CaO ($3.0022 \text{ m}^2\text{g}^{-1}$) and eggshell derived-CaO calcined at 900 °C (eggshell-CaO-963 900) ($3.7262 \text{ m}^2\text{g}^{-1}$). The basicity of modified catalyst lies in the region $12.2 < H_{-} < 15.0$. Figure 964 13b depicted the SEM image of CaO generated from the calcination–hydration–dehydration 965 treatment of eggshells (i.e. egg shell-CaO-900-600) which shows a honeycomb-like porous 966 surface. However, in the case of the eggshell-CaO-900, rod-like structure with microporous 967 particles (size ranging from 1.29 to 2.0 µm) was observed (Figure 13a). 968



Figure 13: SEM image of a) eggshell-CaO-900. b). Eggshell-CaO-900-600. Reproduced from ref. [172].

969

In another work, waste chicken fat obtained from slaughterhouse was converted to
FAME using calcined chicken eggshell catalyst under microwave irradiation (Figure 14).¹⁹⁰
Esterification was carried out to lessen FFA content of the chicken oil below 1 mg KOH/g of
oil, followed by transesterification to yield FAME. Flow diagram of biodiesel production using
chicken eggshell as a catalyst is presented in Figure 15. Optimization of transesterification
process parameters by response surface methodology was performed.

- 976
- 977



Figure 14: Microwave-assisted synthesis of FAME using an eggshell catalyst. Reproduced from ref. [190].



Figure 15: Flow diagram of biodiesel production utilizing chicken eggshell catalyst. Reproduced from ref. [190].

981 Similarly, *Helianthus annuus* L oil was converted to FAME using eggshell derived
982 CaO.¹⁹² The preparation route of CaO starting from the shell is presented in Figure 16. Under
983 the optimized reaction conditions, 99.2 % of FAME yield was achieved. The catalyst is stable
984 up to the fourth cycle where 87.8 % yield was observed.



Figure 16: Schematic layout for eggshell originated CaO synthesis. Reproduced from ref. [192].

Earlier, Ansori et al.¹⁹⁵ reported a chicken shell derived CaO catalyzed synthesis of FAME 987 from C. inophyllum L oil under a microwave (MW) irradiation. Initially, oil FFA content was 988 pre-esterified utilizing H₂SO₄, which was then transesterified by utilizing the CaO catalyst 989 (originated from chicken shell) and reported 98.90 % FAME yield in 12.47 min. In another 990 work, Mansir et al.¹⁹⁶ examined the application of W/Mo/CaO catalyst, where tungsten and 991 molybdenum were impregnated on CaO derived from waste eggshell, for the transformation of 992 WCO via concerted esterification/transesterification to produce FAME in a one-pot process. 993 994 Moreover, the authors investigated the influence of W and Mo loading on CaO in its catalytic activity and found that catalytic activity increased when wt. % of W is higher than wt. % of 995 996 Mo over the range of 0.3-0.7 %. A maximum yield of 96.2 % was reported under the optimum reaction conditions using 0.6 W/0.4 Mo/CaO. In addition, several literatures are available for 997 998 the transesterification of WCO having FFA content in the range of 4-7.1 % to produce methyl ester using various eggshell derived CaO catalyst impregnated with acidic and basic 999 compounds. Examples of such catalysts are CaO/anthill,¹⁹⁷ CaO/Zn,¹⁹⁸ CaO/KF/Fe₃O₄,¹⁹⁹ 1000 CaO/SiO₂ based on palm empty fruit bunch (PEFB),²⁰⁰ and Mo-Zr/CaO²⁰¹ etc. 1001

In 2015, Joshi *et al.* ¹⁶⁴ synthesized various metal oxides, for example, ZnO, MnO₂, Fe₂O₃ and Al₂O₃ impregnated on CaO derived from eggshell via calcination at 900 °C and exploited these catalysts in the conversion of non-edible JCO to FAME. Among all the mixed metal oxides, the surface area and pore volume of ZnO-CaO is highest and thus showed an excellent 95.2 % JCO conversion. The authors also reported that the catalyst is very stable towards the

transesterification of JCO and can be reused for 4 cycles. Similarly, Teo et al. ²⁰² synthesized 1007 CaO NPs derived from *Gallus domesticus* eggshell via precipitation method and utilized it for 1008 1009 the conversion of JCO to give FAME with 97 % yield under the optimal reaction conditions. 1010 TEM images and particle size distribution of waste eggshell of Gallus domesticus derived nano-CaO catalyst is displayed in Figure 17 (A, B, C) which revealed that the particles were 1011 regular spheroidal shape and the average particle diameter is 16-27 nm. Figure 17 D displays 1012 1013 the basicity measurement of the catalyst and commercial CaO using CO₂-TPD technique. All CaO catalysts showed a broad desorption peak owing to the existence of strong basic strength. 1014 1015 The desorption peaks of both catalysts observed over the temperature ranging from 550 to 700 °C are attributed to the super-basic characteristics of the nanoparticles. 1016

In 2011, Olutoye et al. ²⁰³ reported a mixed metal solid catalyst, where Mg(NO₃)₂ and 1017 KNO₃ were impregnated on CaO originated from eggshell and exploited it in the 1018 transformation of palm oil to FAME. The authors made three sets of a catalyst by changing the 1019 loading amount of Mg(NO₃)₂ and KNO₃ on CaO with wt. % ratio of 6:1:1, 2:1:1 and 1:1.5:1.5 1020 and investigated their influence on the transesterification reaction and reported that the catalyst 1021 with wt. % ratio of 6:1:1 showed the maximum yield of 85.8 %. In addition, several works are 1022 reported in the literature regarding the transesterification of palm oil using chicken shell 1023 derived CaO modified solid catalyst such as CaO/SiO2^{204,205} CaO/rice husk ²⁰⁶ etc. Recently, 1024 Sulaiman et al. ²⁰⁷ successfully synthesized a mixture of calcined coconut waste and egg waste 1025 1026 for the transformation of palm oil to biodiesel. The authors employed RSM based on CCD to study the ideal reaction conditions: coconut waste/eggshell waste ratio, M/O molar ratio, 1027 1028 catalyst amount, reaction temperature and reaction time. After a successful investigation, they reported that 5:1 wt. % ratio of coconut waste/eggshell waste showed the maximum yield of 1029 1030 81 % under the optimal reaction conditions.

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Figure 17: TEM images and particle size distributions of surfactant assistant CaO nanocatalysts: after 40 min (a); after 80 min (b) and after 120 min (c) and CO_2 desorption performance commercial of CaO (a) and nano CaO catalysts: after 40 min (b); after 80 min (c) and after 120 min (d). Reproduced from ref. [202].

1040 In another work, A Li doped CaO catalyst derived from eggshell was examined for the 1041 transformation of nahor oil to produce FAME by Boro *et al.* 208 They measured the FFA content 1042 in the nahor oil and found 15 mg KOH/g. Due to this high FFA contents, a two-step process 1043 was investigated; firstly, esterification was performed using sulfuric acid to bring down FFA 1044 amount up to <1 followed by transesterification reaction using Li/CaO catalyst. They also 1045 examined the impact of Li doping on the conversion of oil to FAME and reported a maximum

94 % conversion when the Li doping was 2 wt. %. Recently, Rahman et al. ²⁰⁹ modified CaO 1046 derived from chicken eggshell with transition metals such as Zn and Cu and applied the catalyst 1047 1048 in the transformation of eucalyptus oil to FAME. The authors reported that the surface area and 1049 basicity of Zn/CaO are higher than the Cu/CaO, therefore Zn/CaO showed better result with 93.2 % FAME yield. Moreover, the impregnation of Zn on CaO improved the stability of the 1050 catalyst and can be used for 7 consecutive cycles. In another report, a magnetically recoverable 1051 1052 KF modified CaO derived from eggshell was prepared and employed in the transformation of neem oil to FAME.¹⁹⁹ The author reported that the primary advantage of the catalyst is that the 1053 catalyst circumvented saponification reaction and therefore transesterification of neem oil 1054 (FFA content 4.2 %) can proceed through the one-step process, and 94.5 % FAME can be 1055 1056 achieved.

1057 In 2010, a novel eggshell originated CaO impregnate on fly ash was reported for the transesterification of soybean oil to form FAME. The influence of CaO loading was studied by 1058 the authors and found that 30 wt. % CaO loading showed a maximum yield of 96.97 %. 1059 Moreover, CaO supported on fly ash enhanced catalyst reusability and reactivity compared to 1060 neat eggshell originated CaO.²¹⁰ In addition, a KF modified CaO originated from eggshell was 1061 examined for the transformation of soybean oil to FAME. The modified catalyst has higher 1062 basicity than the neat CaO due to the addition of KOH in the process.²¹¹ Recently, Chowdhury 1063 et al.²¹² synthesized a Na-doped CaO derived from chicken eggshell and exploited it in the 1064 transesterification of Madhuca indica oil. A two-step process was employed as the oil have 45 1065 % of FFA content. They first esterified the oil using 5 wt. % sulfuric acid to lessen FFA content 1066 1067 of the oil followed by transesterification using Na-doped CaO catalyst. To study the influence 1068 of reaction parameters on the transformation of oil to biodiesel, Taguchi approach was used, 1069 where they observed that M/O molar ratio and the reaction temperature have the highest impact and reaction time has the minimal impact on the transformation of oil to FAME. In 2014, Chen 1070 et al.²¹³ demonstrated the synthesis of FAME from palm oil using CaO catalyst derived from 1071 ostrich egg-shell via ultrasonication. They compared the production of biodiesel using both 1072 mechanical stirring and ultrasonication process and reported that the latter case showed higher 1073 yield (92.7 %). Moreover, the catalyst can be used for 8 consecutive cycles. A 1074 1075 transesterification process for soybean oil deodorizer distillate (SODD) to produce FAME was 1076 reported using CaO derived from the duck eggshell. They measured the FFA content of SODD and found 53.2 %, therefore to overcome the saponification problem the oil was pre-esterified 1077 1078 with sulfuric acid and then the transesterification was performed of the pre-esterified SODD oil using CaO catalyst to produce FAME with an overall yield of 94.6 %.² In addition, CaO 1079

1080 derived from quail eggshell were also utilized for the transformation of palm oil ²¹⁴ and JCO

1081 ²¹⁵ to biodiesel in high yield.

No.	Catalyst source	Catalyst	Feedstock	^a Conditions	Yield (%)	Ref.
1.	Chicken eggshell	CaO	Soybean oil	9:1, 3, 65, 180	>95	161
2.	Chicken eggshell	CaO	Soybean oil	10:1, 7, 57.5,	93	216
				120		
3.	Chicken eggshell	CaO	Soybean oil	8:1, 10, 65, 180	90	163
4.	Chicken eggshell	CaO	Soybean oil	14:1, 4, 60, 180	91	164
5.	Ostrich eggshell	CaO	Karanja oil	8:1, 2.5, 65, 150	95	165
6.	Chicken eggshell	CaO	WCO	22.5:1, 3.5, 65,	91	166
				330		
7.	Chicken eggshell	CaO	WCO	12:1, 1.5, 65,	94	167
				120		
8.	Chicken eggshell	CaO	WCO	4:1, 2, 65, 120	NR	168
9.	Chicken eggshell	CaO	WFO	9:1, 3, 65, 180	95.05	169
10.	Chicken eggshell	CaO	WCO	12:1, 1.5, 60, 60	96.23	191
11.	Chicken eggshell	CaO	WCO	24:1, 4, 60, 240	100	217
12.	Chicken eggshell	CaO	WCO	12:1, 5, 65, 60	94.52^{b}	172
13.	Chicken eggshell	CaO	WCO	10:1, 1.5, 60, 50	96.07	173
14.	Chicken eggshell	CaO	WCO	6:1, 3, 60, 30	97.50	174
15.	Chicken eggshell	CaO	WCO	9:1, 5, 65, 165	87.8	175
16.	Chicken eggshell	CaO	WCO	15:1, 6, 65, 420	75.92	218
17.	Chicken eggshell	CaO	Palm oil	18:1, 10, 60, 90	>90	176
18.	Chicken eggshell	CaO	Palm oil	18:1, 15, 900 W,	96.7	177
				4		
19.	Chicken eggshell	CaO	Palm oil	12:1, 10, 60, 120	94.1	178
20.	Chicken eggshell	CaO	Palm oil	6:1, 5, NR, 30	95	179
21.	Chicken eggshell	CaO	Rape seed oil	9:1, 3, 60, 180	96	180
22.	Chicken eggshell	CaO	Rapeseed oil	9:1, 4, 60, 60	95.12	181
23.	Chicken eggshell	CaO	Sunflower oil	9:1, 3, 60, 180	96	182

24.	Chicken eggshell	CaO	Sunflower oil	11:1, 5, 60, 3	83.2	183
25.	Chicken eggshell	CaO	Sunflower oil	9:1, 3, 60, 240	97.75	219
26.	Chicken eggshell	CaO	Sunflower oil	12:1, 2, 60, 180	100	185
27.	Chicken eggshell	CaO	JCO	81, 2, 65, 150	90	186
28.	Chicken eggshell	CaO	Microalgae	10:1, 1.39, 70,	92.03	187
			Chlorella	180		
			vulgaris			
29.	Chicken eggshell	CaO	Microalgae	10:1, 1.7, 70,	86.41	188
				216		
30.	Chicken eggshell	CaO	Micro algae/	10:1, 1.61, 75,	90.44	189
			S. armatus	240		
31.	Chicken eggshell	CaO	Chicken fat	13:1, 8.5, 57.5,	90.41	190
				300		
32.	Chicken eggshell	CaO	Catfish oil	12:1, 1.5, 60, 60	87.77	191
33.	Chicken eggshell	CaO	Helianthus	8:1, 2.5, 65, 120	99.2	192
			annuus L oil			
34.	Chicken eggshell	CaO	Cotton oil	9:1, 3, 60, 180	98.08	193
35.	Chicken eggshell	CaO	C. sativa oil	12:1, 1, 65, 120	97.2	194
36.	Chicken eggshell	CaO	С.	9:1, 3.88, MW,	98.90	195
			inophyllum L	12.47		
			oil			
37.	Chicken eggshell	CaO/W/Mo	WCO	15:1, 2, 70, 120	96.2	196
38.	Chicken eggshell	CaO/anthill	WCO	6:1, 5, 60, 120	70	197
39.	Chicken eggshell	CaO/Zn	WCO	20:1, 5, 65, 240	96.74	198
40.	Chicken eggshell	CaO/KF/Fe ₃	WCO	15:1, 6, 65, 120	97	199
		O_4				
41.	Chicken eggshell	CaO/SiO ₂	WCO	14:1, 8, 60, 90	96	200
		based on				
		PEFB				
42.	Chicken eggshell	Mo-Zr/CaO	WCPO	15:1, 3, 80, 180	90.1	201
43.	Chicken eggshell	ZnO/CaO	JCO	12:1, 5, 65, 60	98.2	164
44.	Chicken eggshell	CaO NPs	JCO	6:1, 2, 90, 120	98	202

-	45.	Chicken eggshell	Ky(MgCa) _{2x}	Palm oil	16:1, 5.53, 65,	88	203
			O ₃		273		
	46.	Chicken eggshell	CaO/SiO ₂	Palm oil	15:1, 9, 65, 480	80.21	204
	47.	Chicken eggshell	CaO/SiO ₂	Palm oil	15:1, 3, 60, 120	87.5	205
	48.	Chicken eggshell	CaO/Rice	Palm oil	9:1, 7, 65, 240	91.5	206
			husk				
	49.	Chicken eggshell	CaO/Cocon	Palm oil	24:1, 5, 65, 180	81	207
			ut waste				
	50.	Chicken eggshell	Li/CaO	Nahor oil	10:1, 5, 65, 240	94	208
	51.	Chicken eggshell	CaO/Zn	Eucalyptus	6:1, 5, 65, 150	93.2	209
				oil			
	52.	Chicken eggshell	CaO/KF/Fe ₃	Neem oil	15:1, 6, 65, 120	97	199
			O_4				
	53.	Chicken eggshell	CaO/fly ash	Soybean oil	6.9:1, 1, 70, 300	96.97	210
	54.	Chicken eggshell	CaO/KF	Soybean oil	12:1, 2, 65, 120	99.1	211
	55.	Chicken eggshell	Na/CaO	Madhuca	9:1, 5, 60, 120	81.1	212
				<i>indica</i> oil			
	56.	Ostrich eggshell	CaO	Palm oil	9:1, 8, 60, 60	92.7	213
	57.	Duck eggshell	CaO	SODD	10:1, 10, 60, 80	94.6	2
	58.	Quail eggshell	CaO	Palm oil	12:1, 1.5, 65,	98	214
					120		
	59.	Quail eggshell/	CaO	Jatropha oil	18:1, 4, MW, 5	94	215
		crab shell					

^aMethanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).

1085 ^bConversion

1086 NR=Not reported

1087 WCPO= Waste cooking palm oil

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1089 7.1.7.2.2. Mollusk shell and other seashells

Mollusk shell and other seashells derived solid catalyst has been widely investigated in the transformation of edible/non-edible oils to produce biodiesel, and are listed in Table 13. Examples are a basic solid catalyst developed by impregnation of KI on the calcined oyster shell which was utilized in the transformation of soybean oil to FAME.¹¹⁶⁻¹¹⁹ The authors reported that impregnation and calcination increase the surface area to an extent of 32 fold and 1095 therefore increases the catalytic activity. The main disadvantages of the catalyst are the reusability factor and higher loading of KI.²²⁰ In addition, there is various literature where neat 1096 CaO derived from oyster shell were utilized for transformation of soybean oil to FAME ²²¹ and 1097 microwave-assisted (800 W) biodiesel synthesis from jatropha oil.²²² Recently, a basic 1098 1099 heterogeneous catalyst was developed from the river snail shell by calcination at 800 °C for 4 h. The catalyst was employed for the transesterification of WCO for biodiesel production. They 1100 1101 performed KOH titration and found that FFA content in the WCO is 0.3 %, therefore direct transesterification was carried out and 98.19 % yield was achieved under the optimal reaction 1102 conditions.²²³ Elsewhere other reports are also available where CaO derived from calcined river 1103 snail were used for the transesterification of various edible/non-edible oils, for example, palm 1104 oil,²²⁴ soybean oil²²⁵ and WFO.²²⁶ 1105

In 2016, Liu et al.²²⁷ developed a solid catalyst, where KBr was loaded on calcined 1106 snail shell and kaoline mixture followed by activation of the catalyst *via* calcination at 500 °C 1107 for 4 h and applied the catalyst in the transformation of soybean oil to FAME. They also 1108 investigated the effect of loading of KBr and wt. % ratio of snail shell/kaoline mixture on 1109 biodiesel yield and found that the catalyst showed a maximum yield of 98.5 % when the KBr 1110 loading and wt. % ratio of snail shell/kaoline were 40 wt. % and 4:1, respectively. Mixing of 1111 1112 snail shell and kaoline together provides the catalyst extra stability compared to their pure form ²²⁷. In addition, Laskar et al. ²²⁸ developed a solid basic catalyst CaO derived from a calcined 1113 snail shell for the conversion of soybean oil to biodiesel. Under the ideal reaction states, 98 % 1114 biodiesel yield was achieved. It is reported that at 400-600 °C calcination temperature, CaCO₃ 1115 1116 of snail shell transformed to calcite. When calcination temperature was further increased to 700 and 800 °C, a minor and major component of CaO was achieved, which was later completely 1117 1118 transformed into CaO at 900 °C calcination temperature. Figure 18 reveals that 100 % 1119 transformation of CaCO₃ into CaO can be achieved above 800 °C calcination temperature. 1120



Figure 18: XRD spectra of normal and calcined (400-1000 °C) snail shells. Reproduced from ref. [228].

In another work, El-Gendy et al.²²⁹ reported CaO catalyst originated from snail shell 1122 1123 calcined at 800 °C and utilized it in the transesterification reaction. RSM was utilized to 1124 investigate the influence of reaction parameters on the biodiesel production and reported that 1125 96.76 % yield was observed under the optimized reaction conditions. Similarly, various literature is available for the transesterification of WCO to FAME using CaO derived from 1126 snail shell collected from different sources.^{230,231} Very recently, Krishnamurthy et al.²³² 1127 developed a solid catalyst, CaO nanoparticles derived from snail shell via the hydrothermal 1128 method and investigated its application in the transesterification of *H. wightiana* oil to produce 1129 FAME. However, a high FFA content (7.57 %) in the oil led the authors to follow a two-steps 1130 process: 1) pre-esterification and 2) transesterification for the production of FAME. RSM was 1131 utilized to examine the impact of reaction parameters on FAME synthesis, reported 96.92 % 1132 vield under the optimal reaction conditions. In a similar vein, CaO derived from snail shell was 1133 also investigated for the transformation of A. africana seed oil ²³³ and showed 85 % FAME 1134 yield. 1135

A calcined mussel/cockle/scallop shell derived CaO was developed for the transformation of palm oil for FAME production. The authors reported high catalyst reactivity catalytic activity with great stability towards the transesterification of palm oil with 95 % conversion.²³⁴ In the meantime, Hadiyanto *et al.*²³⁵ developed a solid catalyst, modified CaO 1140 (derived from green mussel shell) with activated carbon (C) followed by impregnation on NaOH and utilized the catalyst in the transformation of palm oil. The wt. % C/CaO ratio of 2:3 1141 showed the maximum yield of 95.12 % under the optimal reaction conditions. Similarly, KOH 1142 impregnated mussel shell derived CaO was examined for castor oil transformation to biodiesel. 1143 The authors made a comparison between non-impregnated and KOH impregnated catalysts and 1144 revealed that the KOH impregnated catalyst displayed higher reactivity as well as basicity and 1145 reported 91.7 % FAME yield using KOH impregnated catalyst.²³⁶ Moreover, calcined mussel 1146 shell derived catalyst were widely examined for the transformation of vegetable oils, for 1147 example, soybean oil, ^{237,238} Chinese tallow oil, ²³⁹ Camelina sativa oil ²⁴⁰ etc., for biodiesel 1148 production. 1149

Syazwani et al.²⁴¹ examined CaO, originated from angel wing shell (AWS) calcined at 1150 900 °C for 2 h, for the conversion of N. oculata micro-algae oil to FAME. The catalyst 1151 possessed high reactivity with great stability and can be reused for 3 consecutive cycles. 1152 1153 Furthermore, a bifunctional catalyst was developed for conversion of palm fatty acid distillate (PFAD) to FAME. The angel wing shell was calcined to form CaO followed by sulfonation to 1154 1155 afford the catalyst. The authors reported that the catalyst surface area increased to two-fold after the modification, as a result, the catalyst showed excellent activity towards the 1156 1157 esterification of PFAD. Unfortunately, the catalyst was reusable only for two cycles as blocking of active sites occurred in each reaction cycles. Therefore to enhance the reusability of the 1158 1159 catalyst, pretreatment of the catalyst such as washing and re-calcination are necessary before each reaction cycles.²⁴² In 2015, Asikin-Mijan *et al.*²⁴³ developed a waste clam shell derived 1160 1161 CaO using hydration-dehydration treatment and investigated its catalytic application in the conversion of palm oil to FAME. They also examined the effect of hydration-dehydration time 1162 1163 on biodiesel conversion and found that higher is the hydration time higher is the catalytic 1164 activity as hydration for longer-term enhanced the formation of Ca(OH)₂ and thus increased the basicity, reduced the crystallinity and also enhanced the surface area. They reported that 1165 the rehydration for 12 h showed the maximum 98 % FAME yield under optimized reaction 1166 conditions. Similarly, investigation of naked CaO catalyst, derived from a calcined short-1167 necked clamshell, recorded 93 % biodiesel yield under the optimal reaction conditions.²⁴⁴ In 1168 addition, CaO derived from various calcined clamshell were utilized for the transformation of 1169 diverse edible/non-edible oils, for example, palm oil, ^{245,246} WFO ²⁴⁷ etc., to produce biodiesel. 1170

1171 A solid ethanol-treated catalyst CaO, derived from calcined abalone shell was examined 1172 for production of FAME from palm oil. The authors investigated the impact of ethanol 1173 treatment at different temperature (RT, 100 °C and 160 °C) and found that the catalyst treated with ethanol at 100 °C showed the maximum yield of 96.2 % as ethanol treatment provides high basicity, high surface area and lowered the catalyst crystallinity. Moreover, comparison of modified CaO with naked CaO showed that modified CaO has higher reusability and provided higher biodiesel yield.²⁴⁸ In addition, there are several reports available in literature regarding the transesterification of palm oil to FAME utilizing CaO based solid catalyst originated from various waste shells such as *T. jourdani* shell,²⁴⁹ *A. cristatum* shell,²⁵⁰ cockle shell ²⁵¹ and obtuse horn shell ²⁵² etc.

In 2009, Xie et al. 253 synthesized a solid catalyst via three-step: i) incomplete 1181 carbonization of a biont shell at 500 °C, ii) KF impregnation and iii) catalyst activation at 300 1182 °C. The developed catalyst was utilized for conversion of rapeseed oil to FAME. They reported 1183 that the catalyst displayed excellent reactivity due to the formation of a higher amount of active 1184 sites during the reaction between incomplete carbonized shell and KF. The effect of KF loading 1185 was also examined and found that 25 % KF loading is optimal and showed 97 % FAME yield 1186 under the optimized reaction conditions. Correspondingly, Boro et al. ²⁵⁴ demonstrated the 1187 synthesis of CaO catalyst by calcination of Turbonilla striatula shell and utilized it for 1188 1189 transformation of mustard oil to FAME. The effect of calcination temperature was examined and observed that the catalyst calcined at 900 °C displayed maximum 93.3 % FAME yield. In 1190 1191 addition, CaO derived from calcined Turbonilla striatula was modified with Ba in the range of 0.5-1.5 wt. % and utilized it for the transformation of WCO to biodiesel. Due to the high acid 1192 value 22 mg KOH/g, the oil was pretreated with sulfuric acid to reduce the acid value <1. Then 1193 the pretreated oil was transesterified with Ba/CaO catalyst. The authors also examined the 1194 1195 effect of Ba loading and found that 1 % of Ba doped showed >98 % biodiesel yield.²⁵⁵ In addition, Chicoreus brunneus shell was calcined above 800 °C to convert CaCO3 to CaO 1196 1197 followed by hydration/dehydration to form a solid base catalyst and examined it for the transformation of rice bran oil. Calcination and hydration provide the catalyst high porosity, 1198 enhances the basicity, catalytic activity and reusability.²⁵⁶ In addition, shrimp shell originated 1199 catalysts has also been utilized for transformation of various edible/non-edible oils to FAME. 1200 Yang *et al.*²⁵⁷ synthesized a catalyst *via* a three steps processes; (i) inadequate carbonization 1201 of shrimp shell, (ii) reaction with KF and (iii) activation of the catalyst under the heating 1202 condition for the rapeseed oil transformation. The authors examined the impact of 1203 carbonization temperature, KF amount and activation temperature and found that 89.1 % 1204 biodiesel was achieved under the reaction states: carbonization temperature of 450 °C, KF 1205 amount of 25 wt. % and an activation temperature of 250 °C. The excellent catalyst reactivity 1206 is attributable to the formation of active sites during the reaction between incomplete 1207

carbonized shrimp shell and KF. Moreover, a solid catalyst, CaO nanoparticles with a diameter
of 66 nm derived from *Polymedosa erosa* shell *via* calcination-hydration-dehydration process
was developed for the transformation of JCO to FAME in a two-step procedure: 1) preesterification and 2) transesterification. The influence of reaction parameters on the oil
conversion was examined by RSM technique and displayed 98.54 % FAME yield.²⁵⁸

In the recent past, Sivakumar et al. ²⁵⁹ developed a solid catalyst derived from Scylla 1213 Tranquebarica crab shell calcined at 750 °C for sunflower oil transformation to FAME. The 1214 developed catalyst displayed similar reactivity to that of commercial CaO and reported a very 1215 high conversion of 94.2 % under the optimal reaction conditions. Similarly, Shankar et al. ²⁶⁰ 1216 prepared a solid catalyst where CaO (derived from crab shell calcined at 900 °C) impregnated 1217 on Na-ZSM-5 followed by activation at 550 °C for 10 h and utilized it for production of FAME 1218 from neem oil. The impact of CaO loading was examined and found that 15 wt. % CaO 1219 impregnation showed a maximum 95 % biodiesel formation. Moreover, various reports are 1220 available for the transesterification of edible/non-edible oils such as palm oil ²⁶¹ and karania 1221 oil ²⁶² utilizing CaO originated from calcined crab shells. 1222



No.	Catalyst source	Catalyst	Feedstock	^a Conditions	Yield	Ref.
					(%)	
1.	Oyster shell	CaO/KI	Soybean	10:1, 1mmol/g,	79.5	220
				50, 240		
2.	Oyster shell	CaO	Soybean oil	6:1, 25, 65, 300	73.8	221
3.	Oyster and	CaO	Jatropha oil	15:1, 4, MW, 6	93	222
	Pyramidella					
	shells					
4.	River snail shell	CaO	WCO	9:1, 3, 65, 60	92.5^{b}	223
5.	River snail shell	CaO	Palm oil	12:1, 5, 65, 90	98.5	224
6.	River snail shell	CaO	Soybean oil	9:1, 3 ^{<i>c</i>} , 65, 180	98	225
7.	River snail shell	CaO	WFO	6.03:1, 2, 60,	87.28	226
				420		
8.	Snail shell	CaO/KBr/k	Soybean oil	6:1, 2, 65, 120	98.5	227
		aolin				
9.	Snail shell	CaO	Soybean oil	6:1, 3, RT, 420	98	228

Table 13: Various mollusk and seashells derived solid catalyst for biodiesel production.

10.	Snail shell	CaO	WFO	6:1, 3, 60, 60	96	229
11.	Snail shell	CaO	WCO	9:1, 9, 60, 180	84.14	230
12.	Snail shell (S. canarium)	CaO	WCO	12:1, 3, 65, 240	83.5	231
13.	Snail shell	Nano-CaO	<i>H. wightiana</i> oil	12.4:1, 0.892, 61.6, 145.154	98.93	232
14.	Snail shell	CaO	A. africana seed oil	6:1, 1.5, 55, 65	85	233
15.	Mussel/cockle/sc allop shell	CaO	Palm oil	9:1, 10, 65, 180	95	234
16.	Mussel shell	C/CaO/Na	Palm oil	0.5:1, 7.5, 65,	95.12	235
	(Perna varidis)	OH		180		
17.	Mussel shell	CaO/KOH	Castor oil	6:1, 2, 60, 180	91.17	236
18.	Mussel shell	CaO	Soybean oil	24:1, 12, 60, 480	94.1	237
19.	Mussel shell	CaO	Soybean oil	9:1, 4, 65, 180	>98 ^b	238
20.	Fresh water	CaO	Chinese tallow	12:1, 5, 70, 90	97.5	239
	mussel shell		oil			
21.	Mussel/clamp/oys	CaO	Camelina	12:1, 1, 65, 120	95/93	240
	ter		sativa oil		/91	
22.	Angel wing shell	CaO	N. oculata	150:1, 9, 65, 60	84.11	241
			(Microalgae)			
			oil			
23.	Angel wing shell	CaO-SO ₄	PFAD	15:1, 5, 80, 180	98 ^b	242
24.	Clamshell	CaO	Palm oil	9:1, 1, 65, 120	98	243
25.	Short necked	CaO	JCO	20:1, 4, 65, 360	93	244
	clam (O.					
	orbiculata) shell					
26.	Clamshell (<i>M</i> . <i>mereterix</i>)	CaO	WFO	6.03:1, 3, 60, 180	> 89	245
27.	White bivalve clamshell	CaO	WFO	18:1, 8, 65, 180	95.84	246

28.	Venus clam	CaO	Palm oil	15:1, 5, 65, 360	97	247
	(Tapes belcheri					
	<i>S</i> .)					
29.	Abalon shell	CaO	Palm oil	9:1, 7, 65, 150	96.2	248
30.	T. jourdani shell	CaO	Palm oil	3:1, 10, 80, 420	99.33 ^b	249
31.	A. cristatum shell	CaO	Palm oil	8:1, 3, 60, 360	93	250
32.	Cockleshell	CaO	Palm oil	0.54:1, 4.9, reflux, 180	99.4	251
33.	Obtuse horn shell	CaO	Palm oil	12:1, 5, reflux, 360	86.75	252
34.	Biont (turtle) shell	CaO/KF	Rape seed oil	9:1, 3, 70, 180	97.5	253
35.	<i>Turbonilla</i> striatula shell	CaO	Mustard oil	9:1, 3, 65 ±5, 360	93.3	254
36.	<i>Turbonilla</i> striatula shell	CaO/Ba	WCO	6:1, 1, 65, 120	> 98 ^b	255
37.	Chicoreus brunneus shell	CaO	Rice bran oil	30:1, 0.4, 65, 120	93	256
38.	Shrimp shell	CaO/KF	Rape seed oil	9:1, 2.5, 65, 180	89.1 ^{<i>b</i>}	257
39.	P. erosa seashells	Nano-CaO	Jatropha oil	5.15:1, 0.02, RT, 133.1	95.8	258
40.	Crab shell (S. tranquebarica)	CaO	Sunflower oil	12:1, 8, 95, 75	94.2	259
41.	Crab shell	CaO/Na- ZSM-5	Neem oil	12:1, 15, 75, 360	95	260
42.	Crab shell (S. serrata)	CaO	Palm oil	0.5:1, 5, 65, 150	98.8	261
43.	Crab shell	CaO	Karanja oil	8:1, 2.5, 65, 120	94	262

¹²²⁴ ^{*a*}Methanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).

^bConversion

1226 ^{*c*}w/w

1227 PFAD = palm fatty acid distillate

1229 **7.1.7.2** Ashes of biomass

In recent years, the application of waste plant ashes as a highly active heterogeneous 1230 catalyst has drawn increasing attention in the realm of biodiesel production. A huge amount of 1231 alkali or alkaline earth elements mostly K, Ca and Mg present in the ashes of waste plant 1232 biomass acted as a highly basic catalyst in the transesterification reaction to produce biodiesel 1233 from vegetable oil with low FFA. In case of vegetable oil with high FFA, reduction of FFA to 1234 <1% (by acid-catalyzed esterification) before transesterification reaction is mandatory to elude 1235 catalyst consumption in soap formation, which otherwise leads to low biodiesel yield. Usually, 1236 1237 biomass is collected, washed and dried either in oven or sunlight, burnt in the open air or burnt in the air followed by calcination to produce a highly basic ash catalyst as shown in Figure 19. 1238 Different basic ash catalysts utilized and their efficacy in the synthesis of biodiesel are 1239 presented in Table 14. In a pioneering work, Chouhan et al. ²⁶³ reported the use of amphibian 1240 plant L. perpusilla Torrey ash as a solid catalyst in biodiesel synthesis from JCO. The plant 1241 biomass was subjected to calcination at 550 ± 5 °C for 2 h to obtain the ash catalyst. The 1242 crystallinity of the catalyst was affirmed by XRD patterns. Impact of catalyst loading revealed 1243 1244 that 5 wt. % (w.r.t. oil) is enough to obtain a high 89.43% biodiesel yield under the optimal reaction conditions. Nevertheless, reusability study demonstrated that the catalyst lost its 1245 1246 reactivity in each progressive reaction cycles owing to leaching of the reactive elements in the catalyst. Thereby, the catalyst was recycled up to 3 cycles only. 1247



Figure 19: Flowchart for the synthesis of ash catalyst derived from plant biomass.

In another work, oil palm ash was seen as an active catalyst for biodiesel synthesis from 1250 WCO by Chin et al.²⁶⁴ Figure 20 depicted the SEM micrograph of the palm ash, which 1251 displayed the porous nature of the ash catalyst, while Table 15 listed the elements exist in the 1252 1253 palm ash determined from the EDX analysis. It was observed that the palm ash consisted of a large amount of potassium, while a relatively low quantity of aluminum, zinc, and magnesium 1254 1255 was also found. Besides, it was seen that the K₂O was the primary driver for the high basicity and catalytic activity of the catalyst towards biodiesel synthesis. CCD was utilized to 1256 1257 investigate the impact of the optimized reaction conditions in biodiesel synthesis such as M/O ratio, reaction time and temperature and catalyst loading. Accordingly, the predicted and 1258 experimental biodiesel yields were found to be 60.07 % and 71.74 % respectively. 1259



Figure 20: SEM micrograph of palm ash. Reproduced from ref ²⁶⁴.

Table	15.	EDA	data	IOr	compositions	OI
palm a	sh by	y. Ref.	[264]].		

Elements	Atomic wt. %
Potassium (K)	40.59
Magnesium (Mg)	0.76
Silicone (Si)	2.63
Aluminum (Al)	0.50
Zinc (Zn)	0.33
Oxygen (O)	29.36
Carbon (C)	14.56

1260

Chlorine (Cl)	7.07
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In the meantime, Boey et al.²⁶⁵ reported a solid base, derived from boiler ash (BA) via 1263 calcination, catalyzed biodiesel synthesis from palm oil. BA effectively transformed palm oil 1264 to FAME at moderate reaction conditions and delivered 90% FAME yield. Ironically, the ash 1265 is intolerant to the presence of moisture and FFA at 1 wt. % in the feedstock. Betiku et al. ²⁶⁶ 1266 reported a process for biodiesel synthesis from *Thevetia peruviana* oil by utilizing calcined 1267 Musa paradisiacal (plantain) peel ash catalyst. The dried powdered plantain peels were 1268 calcined at 500 °C for 3.5 h to produce plantain peels ash. Biodiesel yield of 95.2% was 1269 acquired using the optimized reaction conditions. In addition, Etim et al. ²⁶⁷ utilized ripe 1270 plantain fruit peel as a solid catalyst in biodiesel synthesis from Azadirachta indica oil. At the 1271 1272 onset, pre-esterification of the oil was performed to diminish the FFA contents from 5.81 wt. % to 0.90 wt. % utilizing M/O molar ratio of 2.19 v/v and 6 wt. % of Fe₂(SO₄)₃. Finally, the 1273 1274 pre-esterified oil was transformed to FAME via transesterification reaction catalyzed by plantain fruit peel ash. Coconut husk ash catalyst was also reported for biodiesel synthesis from 1275 JCO.²⁶⁸ The husks were subjected to calcination at various temperatures ranging from 250-500 1276 °C and identified that catalyst produced at 350 °C calcination temperature was found to be the 1277 1278 most reactive one for biodiesel synthesis giving 99.86 % yield within 30 min at the moderate 1279 reaction temperature. XRD patterns of the catalysts are presented in Figure 21 which revealed the presence of several components of ash such as KCl, K₂Si₂O₅, K₂SO₄, K₂S₃, KAlO₂, 1280 K₄CaSi₃O₉, FeCa₂Al₂BSi₄O₁₅OH, etc. 1281





Figure 21: XRD patterns of calcined coconut husk calcined at different temperatures. Reproduced from ref ²⁶⁸.

1283

Cocoa pod husks (CPHs) was used as a solid catalyst for biodiesel synthesis from 1284 soybean oil by Ofori-Boateng et al.²⁶⁹ The authors examined the reactivity of MgO 1285 impregnated CPH (MgO@CPH) and bare CPH in biodiesel synthesis under the optimal 1286 reaction states and achieved 98.7% and 91.4% biodiesel yields respectively. Moreover, the 1287 1288 synthesized fuel satisfies the European biodiesel quality norm (EN 14112). In another study, the production of biodiesel from yellow oleander (Thevetia peruviana) seed oil using banana 1289 (Musa balbisiana Colla) peel ash was reported.²⁷⁰ K, Na, CO₃, Cl present in the ash are 1290 responsible for the high basicity, thus the reactivity of the catalyst. Oil transformation of 96 % 1291 was demonstrated in just 3 h time under room temperature. The produced biodiesel conform to 1292 standards set for ASTM D6751, EN 14214 and so forth. BET surface area measurement of the 1293 catalyst revealed that the surface area is $1.487 \text{m}^2/\text{g}$. The biodiesel was free from sulfur and has 1294 displayed a high cetane number. Meanwhile, Musa balbisiana Colla underground stem 1295 (MBCUS) ash was examined as a solid base catalyst for biodiesel synthesis from high FFA 1296 containing JCO by Sharma et al.²⁷¹ Characterization of the ash catalyst revealed that it is 1297 comprised of oxides and carbonates of various alkali and alkaline earth metals, which leads to 1298 the high basicity of the catalyst and surface area is 39 m^2/g . It was reported that the catalyst is 1299 very effective during the biodiesel synthesis process at 275 °C and internal pressure (4.2 MPa) 1300 1301 and resulted in 98.0 % biodiesel yield.

Betiku et al. ²⁷² led an investigation on the application of banana (Musa'Gross Michel') 1302 peel waste as a catalyst for biodiesel synthesis from *Bauhinia monandra* (Napoleon's plume) 1303 seed oil (BMSO), with a motive to develop a low-cost fuel. The burnt ash of the banana peel 1304 1305 was further calcined at 700 °C for 4 h to produce a highly active catalyst. They have utilized RSM model to determine the optimal reaction conditions for biodiesel synthesis using the ash 1306 catalyst. The RSM plot of M/O molar ratio and catalyst loading on Bauhinia monandra 1307 (Napoleon's plume) methyl ester (BMME) yield is shown in Figure 22a. It is observed that 1308 1309 BMME yield improved from 0 to >90 wt. % as the M/O molar ratio expanded from 7:1 to 14:1 and catalyst loading increased from 1.5 to 3.5 wt. %. This might be ascribed to the increase in 1310 active sites number as a result of increased in catalyst loading. Besides, BMME yield 1311 diminished marginally when the catalyst loading was above 3.5 wt. % (Figure 22a). In addition, 1312 the plot revealed a direct connection between the M/O molar ratio and catalyst loading on the 1313

1314 biodiesel yield. As the two parameters increases, biodiesel yield also increased (Figure 22a). The transformation of the pre-esterified oil to biodiesel was done inside the time span of 33.79-1315 76.21 min. The extended reaction time, somewhere in the range of 33.79 and 55 min, favoured 1316 biodiesel yield; after 55 min, the yield diminished. Figure 22b displays the impact of reaction 1317 time and catalyst loading on biodiesel yield. It is observed from the surface plot that rise in 1318 catalyst loading and reaction time leads to an increase in biodiesel yield. Moreover, the plot 1319 1320 displayed that 90 wt. % biodiesel yield is reached using 4.5 wt. % catalyst loading within 80 min reaction time. In addition, Figure 22c illustrates the surface plot to examine the impact of 1321 1322 M/O molar ratio and reaction time on biodiesel yield. It is observed from the plot that increases in two parameters such as M/O molar ratio and reaction time leads to a rise in biodiesel yield. 1323 1324 It is seen from the figure that increases in M/O molar ratio from 7:1-14:1 improved the biodiesel yield from 33% to 100 %. Therefore, the highest biodiesel yield was recorded at 14:1 M/O 1325 molar ratio and 80 min reaction time. 1326 1327



Figure 22: 3-D plots of biodiesel yield. Reproduced from ref. [272].

1329 Meanwhile, Onoji *et al.*²⁷³ built up a novel technique to utilize rubber seed shell (RSS) 1330 ash calcined at 800 °C as a solid base catalyst for the transformation of rubber seed oil to 1331 biodiesel. The high FFA content of the RSS (9.01 ±0.07%) was pre-esterified using H₂SO₄ to 1332 >1% FFA. Reusability study of the catalyst revealed that >80 % biodiesel yield was noticed 1333 after 4 successive reaction cycles. The surface area and pore size of the calcined RSS was found 1334 to be 2.29 nm and 352.51 m²/g, respectively. Similarly, Gohain *et al.*²⁷⁴ studied the application 1335 of *Musa balbisiana* Colla peel ash catalyst to produce biodiesel from WCO. It was observed 1336 that the calcination procedure improved the mesoporous and microporous morphology of the catalyst and upgraded its surface area bringing about the higher catalytic activity. The external 1337 morphology of the catalyst examined by SEM analysis revealed aggregation of the particles 1338 and porosity in the range of micro and meso. Moreover, 100 % conversion of WCO to biodiesel 1339 was confirmed by ¹H NMR spectra (Figure 23b) utilizing Knothe and Kenar equation (1). The 1340 ¹H NMR spectrum of WCO (Figure 18a) displays two peaks at 4.1 and 5.3 ppm because of the 1341 glyceridic protons (Figure 23a). The presence of peak of methoxy protons at \sim 3.6 ppm and 1342 vanishing of the signs of glyceridic peak close to 4-4.2 ppm (Figure 18b) confirmed the 1343 1344 formation of biodiesel.

1345



Figure 23: ¹H NMR Spectrum of (a) WCO and (b) Biodiesel. Reproduced from ref. [274].

1346

In the recent year, Pathak *et al.* ²⁷⁵ utilized *Musa acuminata* peel ash (MAPA) catalyst for biodiesel synthesis from soybean oil at room temperature. Catalyst characterization reported the existence of various alkali and alkaline earth metals that enhance the catalyst basicity and reactivity of the ash catalyst. K (14.27 %), C (47.51 %) and O (30.27 %) are the primary/main elements exist in MAPA as revealed by the XPS data (Figure 24). The authors reported 98.95 % biodiesel yield under the optimized reaction conditions.

- 1353
- 1354



Figure 24: XPS survey (a), C 1s (b), O 1s (c), and K 2p (d) spectra of MAPA. Reproduced from ref. [275].

Sharma et al. ²⁷⁶ investigated the reactivity of wood ash catalyst calcined at different 1356 1357 temperature for biodiesel synthesis from JCO. Ester conversion in the range of 97-99% could be achieved with wood ash catalysts. Wood ash calcined at 800 °C afforded 98.7% oil 1358 conversion under the ideal reaction conditions. Uprety *et al.*²⁷⁷ studied the application of wood 1359 ash derived from birch bark and fly ash blazed at 800 °C for 4 h synthesis of biodiesel from 1360 1361 palm oil. Birch bark ash gave FAME yield of 88.06 ± 0.72 , whereas, fly ash from wood pellet afforded 99.92 \pm 0.01% yield. Recently, the application of banana peduncle ash as an efficient 1362 1363 solid base catalyst for the synthesis of biodiesel from Ceiba pentandra oil (CPO) was investigated.²⁷⁸ Based on the response surface methodology (RSM) study, the ideal reaction 1364 conditions for the transformation of CPO into FAME was found to be 1.978 wt. % catalyst 1365 loading, 60 min response time, 9.20:1 M/O molar ratio with a maximum predicted FAME yield 1366 of 99.36 % which was assessed experimentally as $98.69 \pm 0.18\%$. The same research team also 1367 investigated the utilization of *Musa acuminata* peduncle for biodiesel preparation from CPO.²⁷⁹ 1368 The authors calculated the surface area and pore diameter of the calcined ash catalyst from 1369 BET analysis data and reported 45.99 m^2/g and 9.77 nm respectively. Moreover, the catalyst 1370 consists of diverse minerals along with potassium as primary components, which leads to the 1371

higher reactivity of the catalyst (Figure 25). High conversion of 98.73 ± 0.50 % FAME was

1373 observed under the optimum reaction conditions.

1374



Figure 25: EDS images (a) uncalcined and b) calcined banana peduncle. Reproduced from ref. [279].

1375

In 2019, Mendonça et al. ²⁸⁰ reported the utilization of calcined (800 °C for 4 h) waste 1376 cupuaçu seeds as a solid base catalyst in the synthesis of biodiesel from soybean oil and ethanol. 1377 Similarly, Nath et al. ²⁸¹ utilized a solid base catalyst derived from waste Brassica nigra plant 1378 for the efficient preparation of biodiesel. The SEM-EDX analysis of the catalyst revealed the 1379 existence of potassium (56.13 %) and calcium (26.04 %) in huge amount, which may be 1380 considered as key ingredients for the high basicity of the catalyst. The authors also measured 1381 the surface area pore volume of the catalyst via BET analysis and came about 7.308 m^2/g and 1382 $0.011 \text{ cm}^3/\text{g}$ respectively. The catalyst possessed excellent reactivity in transforming the 1383 1384 soybean oil to FAME and displayed 98.79% FAME yield in a short time frame of 25 min under the optimum states. Betiku et al. ¹⁵⁶ prepared ash catalyst from kola nut pod husk and used it 1385 1386 to convert Kariya seed oil (KSO) to biodiesel, namely Kariya oil methyl esters (KOME), via transesterification process. A maximum of 98.67 ± 0.01 wt. % of FAME yield was observed. 1387

Moreover, the reusability examination of the catalyst suggests that it can be reused for 4 1388 progressive cycles. Recently Changmai et al. 282 converted soybean oil to biodiesel using 1389 orange peel ash in 98 % yield. XRF analysis showed the presence of oxide of potassium (51.64 1390 %) and calcium. Hammet indicator strategy was employed to examine the catalyst basicity and 1391 it was seen as 9.8 < H < 12.2. The author's measured catalyst pore volume and surface area from 1392 BET analysis and found 0.428 cc/g and 605.60 m²/g, respectively. Moreover, GC-MS analysis 1393 1394 (Figure 26) reveals the existence of six components in the synthesized FAME; methyl palmitate (11.63 %), methyl oleate (25.32 %) and methyl linoleate (54.34 %) are the major components. 1395 1396



Figure 26: GC-MS spectrum of biodiesel from soybean oil. Reproduced from ref. [282].

1397

Waste Sesamum indicum plant ash catalyst was also successfully utilized for the 1398 transformation of sunflower oil to biodiesel.²⁸³ The measured surface area of the catalyst is 1399 3.66 m^2/g as obtained from the BET analysis data. A high 98.9 % biodiesel yield was 1400 accomplished. They reused the catalyst up to the 3rd cycle which yields 94.2 % biodiesel. In 1401 addition, Mendonça et al.²⁸⁴ utilized waste tucumã peels ash catalysts for biodiesel synthesis 1402 1403 from soybean oil. The catalyst characterization by XRF showed it was mostly composed of 1404 oxides of potassium, calcium and magnesium.. Because of its heterogeneous and non-leachable nature, the catalyst derived from *tucumã* peels could be reused at least 5 times. In another study, 1405 Tectona grandis leaves ash catalyst was developed and utilized for the transformation of WCO 1406 to FAME by Gohain et al.²⁸⁵ The measured surface area and pore size of the catalyst are 1407 116.833 m²/g and 112.210 Å, respectively as calculated from BET data. 100 % oil 1408 1409 transformation to FAME was accomplished at room temperature using the optimized reaction 1410 conditions. Furthermore, cocoa pod husk derived solid base catalyst was employed in the transformation of neem seed oil to FAME.²⁸⁶ A two-step process was employed for the 1411 conversion of neem seed oil to FAME, i) pretreatment of the oil was performed using $Fe_2(SO_4)_3$ 1412 catalyst to reduce the FFA content from 28.76 % to 0.39 % and ii) transesterification of 1413 pretreated oil using the calcined bio waste-derived catalyst. The authors also studied the effect 1414 of reaction parameters using Box-Behnken design (BBD) and CCD of RSM was utilized to 1415 determine the optimized reaction conditions. Similarly, a walnut shell derived catalyst was 1416 developed for the transformation of sunflower oil to biodiesel.²⁸⁷ The catalyst was prepared 1417 from walnut shells via air combustion, thereby bringing down the cost involved in the 1418 calcination process to afford ash. The authors reported a 98 % FAME yield within a brief time 1419 frame of 10 min. Recently, the transformation of sunflower oil to synthesize FAME using 1420 calcined sugar beet generated from agro-industry waste was reported.²⁸⁸ The catalyst has a high 1421 amount of highly basic CaO and showed very high reactivity towards the transesterification 1422 process to afford about 93 % FAME yield. 98.39 % soybean oil transformation to FAME under 1423 room temperature was reported using *M. acuminata* trunk ash catalyst recently.²⁸⁹ 1424

1425 Most biomass ash catalysts are usually applied for transesterification reactions of different biodiesel feedstocks and different reaction conditions. These make a comparison of 1426 1427 the effectiveness of such catalysts under the same reaction condition impossible. Hence, to have a better insight into the activities of catalysts under the same reaction conditions and 1428 feedstock, Odude *et al.*²⁹⁰ examined the transformation of pre-esterified palm kernel oil (PKO) 1429 to FAME utilizing two diverse catalysts viz. calcined banana peel ash (CBPA) and calcined 1430 1431 cocoa pod husk ash (CCPHA) under same reaction conditions. RSM technique was utilized for the optimization of both CBPA and CCPHA catalyzed transformation process of PKO to 1432 1433 FAME. CCD was utilized to acquire the best possible combination of M/O ratio, catalyst 1434 loading and reaction time for the highest conversion of oil to FAME as portrayed in Figure 27. 1435 The observed FAME yields under the optimized conditions utilizing catalysts CBPA and CCPHA were 99.5 and 99.3 wt. %, respectively. The created models when exposed to statistical 1436 assessment demonstrated that CBPA catalyzed transformation model was better than CCPHA-1437 catalyzed transformation model. In the meantime, *Carica papaya* stem²⁹¹ and *Musa balbisiana* 1438 underground stem ²⁹² was also reported as a solid catalyst to convert *Scenedesmus obliquus* 1439 and Mesua ferrea oil respectively, to FAME. 1440



Figure 27: Contour and surface plots for PKOME synthesis. Reproduced from ref. [290].

Table 14: Different plant ash catalysts in biodiesel productions

No.	Catalyst source	Feedstock	^a Conditions	Yield (%)	Ref.
1.	L. perpusilla Torrey	JCO	9:1, 5, 65 ± 5, 300	89.43	263
2.	Oil palm ash	WCO	18:0, 5.35, 60, 30	71.74	264

3.	Oil palm ash/ boiler ash (BA)	Palm olein	15:1, 3, 60, 30	90	265
4.	Musa paradisiacal	Thevetia	3.3:1, 3, 60, 60	95.2	266
	(plantain) peels	<i>peruviana</i> oil			
5.	Ripe plantain fruit	Azadirachta	1:0.73, 0.65, 65, 57	99.2	267
	peel	<i>indica</i> oil			
6.	Coconut husk	JCO	12:1, 7, 45, 30 min,	99.86	268.
7.	Cocoa pod husks	Soybean oil	6:1, 1, 60, 60/120	98.7/ 91.4	269
8.	Musa balbisiana	Thevetia	20:1 ^d , 20, RT, 180	96 ^b	270
	Colla	peruviana			
	peel	seed oil			
9.	Musa balbisiana	JCO	9:1, 5, 275, 60	98	271
	Colla underground				
	stem				
10.	Musa'Gross Michel'	Napoleon's	7.6:1, 2.75, 65,	98.5	272
	peel	plume seed oil	69.02		
11.	Rubber seed shell	Rubber seed	$0.20:1^e$, 2.2, 60, 60	83.06	273
		oil			
12.	Musa balbisiana	WCO	6:1, 2, 60, 180	100^{b}	274
	Colla peel				
13.	M. acuminata peel	Soybean	6:1, 7, RT, 240	98.95	275
14.	Wood (Acacia	JCO	12:1, 5, 65, 180	98.7 ^{<i>b</i>}	276
	nilotica) stem				
15.	Birch bark/ fly ash	Palm oil	12:1, 3, 60, 180	88.06 ± 0.72/	277
				99.92 ± 0.01	
16.	Musa spp "Pisang	Ceiba	9.20:1, 1.978, 65,	98.69 ± 0.18	278
	Awak" peduncle	<i>pentandra</i> oil	60		
17.	Musa acuminata	Ceiba	11.46:1, 2.68, 65,	98.73 ± 0.50^{b}	279
	peduncle	<i>pentandra</i> oil	106		
18.	Theobroma	Soybean oil	10:1, 10, 80, 480	98.36 ^b	280
	grandiflorum seeds				
19.	<i>Brassica nigra</i> plant	Soybean oil	12:1, 7, 65, 25	98.79	281

20.	Kola nut pod husk	Kariya seed	6:1, 3, 65, 75	98.67±0.01	156
		oil (KSO)			
21.	Orange peel	Soybean oil	6:1, 7, RT, 420	98 ^b	282
22.	Sesamum indicum	Sunflower oil	12:1, 7, 65, 40	98.9	283
	plant				
23.	Tucumã peels	Soybean oil	15:1, 1, 80, 240	97.3 ^{<i>b</i>}	284
24.	Tectona grandis	WCO	6:1, 2.5, RT, 180	100^{b}	285
	leaves				
25.	Cocoa pod husk	Azadirachta	0.73:1 ^e , 0.65, 65, 57	99.3	286
		<i>indica</i> oil			
26.	Walnut shell	Soybean oil	12:1, 5, 60, 10	98	287
27.	Sugar beet waste	Sunflower oil	4.5:1, 1, 75, 60	93 ^b	288
28.	<i>M. acuminata</i> trunk	Soybean oil	6:1, 14, RT, 360	98.39 ^b	289
29.	Banana peel/ cocoa	Palm kernel	0.80:1 ^e , 4, 65, 65	99.5/99.3	290
	pod husk	oil			
30.	Carica papaya stem	Scenedesmus	9:1, 2, 60, 180	93.33 ^b	291
		obliquus			
31.	Musa balbisiana	Mesua ferrea	9:1, 5, 60, 275	95^b	292
	underground stem	oil			

^{a*}Methanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).
 ^bConversion

1446 ^{*d*}mL/g

1447 $e_{\rm V/V}$.

1448

1449 **7.2 Acid catalysts**

Acid can catalyze both transesterification and esterification reactions without soap 1450 formation. ²⁹³ Hence, unlike base catalysts, an acid catalyst has the potential to afford biodiesel 1451 from poor quality oil with high FFA and high water content. In the transesterification reaction, 1452 1453 alkaline catalysts are superior in promoting methoxide anion formation from methanol. In contrary to it, acidic catalysts are less active in methoxide anion formation, but could activate 1454 the carbonyl bonds via H⁺ addition (Brønsted acidic sites) or via coordination of the carbonyl 1455 oxygen with the coordinatively unsaturated metal ion sites (Lewis acidic sites), and thereby 1456 promote transesterification. Hence, an increase in the number of either Brønsted or Lewis 1457 acidic sites promotes faster FAME formation via transesterification. Delightfully, 1458
heterogeneous acid catalysts are endorsed as a potential alternative to homogenous acids catalysts as it possesses certain advantages like it can be easily separated and reused, less corrosive and less toxic. ²⁹⁴ In recent years, several research groups have studied the feasibility of solid acid catalysts for esterification/transesterification processes and proposed economical and environment-friendly approaches for biodiesel production.^{295–297}

1464

1465 **7.2.2 Ion exchange resin**

It is a well-known fact that several catalysts have been employed for FAME production 1466 1467 from various feedstocks. But due to the certain disadvantages of conventional catalysts, researchers are always in search of an ideal catalyst which should overcome all associated 1468 limitations such as catalyst should be active at lower temperatures, exhibits high catalytic 1469 efficacy in terms of conversion of FAME, easy availability, low cost, easy downstream 1470 processing and reusability. Such an ideal catalyst can be considered as potential and 1471 economically viable candidates for biodiesel production.²⁹⁸ In this context, one of them is ion 1472 exchange resin which meets most of the specifications of an ideal catalyst. Thus, many 1473 1474 research groups have studied the role of ion exchange resins as solid catalysts in FAME product.^{299,300} Resin is the insoluble solid material that can retain and discharge ions 1475 simultaneously.³⁰¹ Resins are broad categories into cationic and anionic resin based on 1476 functional groups and degree of cross-linkages. It possesses specific functional groups 1477 responsible for the permutation of ions.³⁰¹ Having one of the important properties, the resin-1478 based catalyst undergoes easy recovery from liquid mixtures by simple methods and are active 1479 1480 at low temperature.

Since last few years, the cationic resins have gained considerable attention due to the 1481 1482 advantages like functioning at soft reaction conditions, non-corrosive nature, more numbers of active sites and lower residual waters production.^{302,303} The cationic resin catalysts possess 1483 numerous active acid sites that play a crucial role in FAME production via 1484 esterification/transesterification reactions.^{304,305} Various ion exchange resin catalyst utilized for 1485 FAME production together with ideal reaction conditions are listed in Table 16. In 2007, 1486 Shibasaki-Kitakawa *et al.* ³⁰² in a comparative study reported that cation exchange resins 1487 1488 showed less efficacy than anion exchange resins towards the conversion of triacylglycerols to biodiesel. Moreover, while evaluating the conversion rates of various commercial resins such 1489 1490 as Diaion PA308, PA306, PA306S and Diaion HPA25, it was observed that highly porous resin-like Diaion HPA25 showed low conversion rate and it is believed this might be due to 1491 resistance of the resin towards the water. According to Ren et al. ³⁰³ transformation of soybean 1492

oil to FAME reduced from 95.2 % to 87.7 % in the existence of D261 anion-exchange resin when the water content is enhanced from 0.0 % to 1.0 % by mass of oil. Similarly, in another study Deboni *et al.* 304 also reported lowering of reaction rate due to the presence of water inside the resins.

Generally, ion exchange resins are utilized for purification and softening of water at 1497 room temperature. Recently, Kansedo et al. ³⁰⁵ compared the catalytic efficiencies of different 1498 ion exchange resins like Amberlyst 15, Dowex DR-2030 and DR-G8 for the transformation of 1499 FFA into FAME via esterification of the sea mango oil (hydrolyzed) at RT. The results 1500 1501 revealed that Amberlyst 15 showed maximum efficacy with the highest FAME production compared to Dowex DR-2030 and Dowex DR-G8. However, Java et al. ³⁰⁶ utilized ion 1502 exchange resins catalyst at a moderately lower temperature (50 °C to 80 °C) for biodiesel 1503 production which is analogous to those of homogenous catalytic process. Further, Umer and 1504 co-worker investigated the transformation of Lagenaria Vulgaris seed oil to biodiesel 1505 exploiting Amberlyst 15 resin and calcium oxide (egg cell) catalyst. The authors reported 93.2 1506 % yield of biodiesel when Amberlyst 15 ion exchange resin was used as a catalyst with the 1507 loading of 5 % w/w and M/O ratio of 40 % w/w for 40 min of and reaction time at 60 °C.³⁰⁷ 1508 Similarly, Kansedo and Lee³⁰⁸ investigated the esterification of hydrolyzed sea mango oil 1509 1510 utilizing different cationic ion exchange resins, and over 80 % yield of FAME was recorded using Amberlyst 15 catalyst at a comparatively lower temperature within 1 hr of reaction time 1511 1512 and with catalyst loading less than 5 % w/w.

Recently, Deboni et al. ³⁰⁴ reported 99 % yield of methyl and ethyl esters from soybean 1513 oil with methanol and ethanol respectively using optimal reaction conditions. Whereas, Guzhan 1514 et al. ³⁰⁹ recorded 63 % yield of FAME from canola oil using Amberlyst-26 under the optimized 1515 reaction conditions.. Moreover, in another study, a yield of about 67 % was observed for canola 1516 oil and methanol with almost similar reaction conditions.^{301,309} The conversion of tallow fat 1517 with methanol showed the yield of methyl and ethyl esters around 95 % using Amberlyst-A26 1518 OH with reaction conditions like tallow fat with methanol molar ratio of 6:1, resin loading of 1519 2 mol/L at 65 °C temperature for about 8.5 hours.³¹⁰ 1520

Hartono *et al.* ³¹¹ investigated the catalytic efficacy of heterogeneous catalyst obtained from a different source like Lewatit macroporous resin, Amberlite gel resin and natural zeolite from Bayah to transform WCO to biodiesel. Authors reported the 85.94 % yield of biodiesel production by Lewatit macroporous anion exchanger with 6 M NaOH. Whereas, Amberlite gel with 6 M HCl displayed 65.22 % biodiesel generation. Previously, Shibasaki-Kitakawa *et al* ³¹² reported the usefulness of the anion-exchange resin from their catalytic and adsorption

abilities for the transformation of WCO to FAME. In their another study, Shibasaki-Kitakawa 1527 et al. ³¹³ also developed ion-exchange resin catalysts based continuous process for the 1528 production of biodiesel. FFA conversions rate was estimated for different catalysts with 1529 1530 reactions conditions like mole ratio of M/O (6:1), temperature (63.83 °C), reaction time (2 h) and catalyst load (20 wt. %). The maximum FFA conversions of 79.7 % were recorded for 1531 NKC-9. Whereas, for 001 x 7 and D61 catalysts it was found to be only 32.2 % and 10.3 % 1532 respectively.³¹⁴ Jalilnejad-Falizi et al. ³¹⁵ achieved the highest FFA conversions by ion 1533 exchange resins (PD206-Na⁺ and PD206-H⁺) under the optimal reation conditions. All the 1534 1535 above-mentioned reports are enough to summarize that ion exchange resins can be employed 1536 as one the potential heterogeneous catalysts in biodiesel production.



No.	Catalyst	Feedstocks	^a Conditions	Yield (%)	Ref.
1	D261 anion-exchange	Soybean oil	9:1, 50.15, 56	95.2^{b}	303
	resin				
2	Amberlyst A26 OH	Acid soybean	9:1, 2, 50, NR	78	304
	anion	oil			
	exchange resin				
3	Amberlyst-15	Hydrolyzed sea	6:1, 30,30, NR	>90	305
		mango oil			
4	Basic anion exchange	Pongamia oil	9:1, 75, 60	85	306
	resin.				
5	Amberlyst 15 ion	Lagenaria	40:1, 5, 60, 40	93.2	307
	exchange resin	vulgaris seed oil			
6	Amberlyst	Hydrolyzed sea	3:12, 100, 60	>80	308
		mango oil			
7	Amberlyst-26	Canola oil	6:1, 3, 45, 90	67	309
8	Amberlyst-A26 OH	Tallow fat	6:1, 2 mol/L, 65,	95	310
			360		
9	Amberlite gel resin	WCO	7:1, 60, 120	85.94	311
10	Cation-exchange resin	Rice bran oil	6:1, 20, 63.83, 120	79.7	313
11	Purolite-PD206	Corn oil	18:1, 65, 2880	79.45	315

Table 16: Different ion exchange resin catalyst used for the production of biodiesel.

¹⁵³⁸ ^{*a*}Methanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).

75

1539 ^bConversion

1540 NR: Not reported

- 1541 PFAD: palm fatty acid distillate
- 1542
- 1543 7.2.3 Sulfated catalyst

Among solid acid catalysts, sulfated catalysts have attracted considerable attention for 1544 transesterification due to their super-acid property. Sulfated inorganic metal oxides are reported 1545 to be chemically stable, and have super acidity comparable to 100 % sulfuric acid, remarkable 1546 acid-base and redox properties.³¹⁶ Different kinds of sulfated catalysts such as sulfated zirconia, 1547 tin oxide, zirconia-alumina, etc. have been successfully exploited in the production of 1548 biodiesel; but among these, sulfated zirconia is most widely studied catalyst (Table 17). 1549 Various reports are available on the transformation of oil to FAME utilizing sulfated zirconia 1550 catalyst, but there are some studies which presented certain drawbacks of these catalysts which 1551 include low catalytic activities, drastic reaction conditions, and reusability issues. Moreover, 1552 lack of uniform pore size and low surface area are the other factors which restrict their wide 1553 1554 uses in catalyzing bulky oil molecules of biodiesel feedstocks. In this context, several attempts have been made to modify sulfated zirconia catalysts with an intention to increase their catalytic 1555 1556 efficacy.

Xia et al. ³¹⁷ demonstrated the synthesis of mesoporous materials which has the 1557 1558 potential to improve the activity of sulfated zirconia catalyst owing to their promising and outstanding properties like high surface area, uniform and controllable pore size. According to 1559 Alexander *et al.* ³¹⁸ the modification of sulfated zirconia catalyst enhanced the total acidity 1560 which basically increased catalyst active sites. In another study, Guoliang et al. ³¹⁹ proposed 1561 1562 that change in phase structure of sulfated zirconia can also increase its catalytic activity and therefore, they developed tetragonal sulfated zirconia which showed enhanced catalytic activity 1563 in the FAME synthesis procedure. Moreover, some of the studies proposed the modification of 1564 sulfated zirconia on MCM-41 (Mobil Composition of Matter No. 41) support for the generation 1565 of methyl tert-butyl ether to improve its catalytic performance, the results obtained revealed 1566 that the catalytic performance of the prepared supported sulfated zirconia catalyst was 2.5-3.0 1567 times greater than neat sulfated zirconia.^{317,320} Similarly, Muthu et al. ³²¹ reported the 1568 preparation of FAME from neem (Azadirochta indica) oil using sulfated zirconia catalyst. It 1569 1570 was revealed that catalyst is highly stable to oils with high FFA concentration. The strong acid sites of this catalyst showed a considerable impact on its reactivity in the transformation of 1571 neem oil. 1572

Recently, Lam et al. 322 developed SO42-/SnO2 catalyst by impregnation method and 1573 exploited it for the conversion of WCO to biodiesel. Further, the authors studied the bi-metallic 1574 1575 impact of the catalyst, in which SnO₂ was blended in with SiO₂ and Al₂O₃, at various weight ratios to increase the activity of SnO₂. The finding confirmed that SO_4^{2-}/SnO_2 -SiO₂ weight 1576 ratio of 3 showed exceptionally high reactivity with 92.3 % biodiesel yield using optimal 1577 reaction conditions. Similarly, Pereira et al. ³²³ demonstrated the application of SnSO₄ catalyst 1578 for the esterification of oleic acid (as model feedstock) and acid soybean oil having high 1579 contents of FFA. It was found that model feedstock containing 70 wt. % of FFA showed 92 % 1580 FAME yield using excess ethanol, 5 wt. % SnSO₄ at 100 °C for 3 h. Moreover, it was also 1581 reported that the catalyst is stable up to ten cycles without any significant decrease in biodiesel 1582 vield. Moreover, one of the studies involved the application of sulfated tin oxide modified with 1583 SiO₂ (SO₄²⁻/SnO₂-SiO₂) catalyst to produce FAME from JCO.³²⁴ The sulfated titania-based 1584 solid superacid catalysts are another kind of sulfated catalysts. Li et al. ³²⁵ prepared three 1585 different titania-based solid superacid catalysts and these were exploited for the transformation 1586 of rape seed oil to FAME at 353 K with a 12:1 molar ratio of M/O under atmospheric pressure. 1587 It was found that all three prepared catalysts showed a significant yield of biodiesel due to their 1588 stronger surface acidities. Moreover, Alaba et al. ³¹⁶ reviewed that apart from these there are 1589 1590 various other sulfated metal oxides such as titania, silica and combination of both also showed remarkable performance. It was also proved thorough investigation led by several researchers 1591 who have applied sulfated silica as catalysts for esterification and transesterification.^{326,327} In 1592 this context, Gardy and co-workers demonstrated a facile preparation of sulfated doped TiO₂ 1593 1594 catalyst that has been utilized efficiently in petroleum refinery. The authors reported that the synthesized catalyst has better reactivity than other sulfated metal oxides, primarily because of 1595 1596 the acidic properties of TiO₂ particles, which was subjected to sulfonation to enhance its acidity. The catalyst displayed great efficiency in the synthesis of FAME from WCO. ^{328,329} 1597 1598

No.	Catalyst	Feedstocks	^a Conditions	Yield (%)	Ref.
1	SO_4^{2-}/ZrO_2	Neem oil	9:1, 1, 65, 120	95	321
2	SO_4^{2-}/SnO_2 -SiO ₂	WCO	15:1, 3, 150, 180	92.3	322
3	SnSO ₄	Soybean oil	3.5:1, 5, 100,180	92	323
4	SO_4^{2-}/SnO_2 -SiO ₂	Jatropha oil	15:1, 3, 180, 120	97	324
5	SO4 ²⁻ /TiO ₂	Rapeseed oil	12:1, NR. 80, 720	51	325

 Table 17: Different types of sulfated catalyst reported for biodiesel production.

6	Ti(SO ₄)O	WCO	9:1, 1.5, 75, 180	97.1	328
7	TiO ₂ /PrSO ₃ H	WCO	15:1, 4.5, 60, 540	98.3	329

^aMethanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).
NR: Not reported

1601

1602 7.2.1 Mixed metal oxides

A wide range of acidic mixed metal oxide catalysts has been utilized to overcome the 1603 problem associated with high FFA content in low-cost biodiesel feedstock employed in FAME 1604 production (Table 18). Suzuta et al. ³³⁰ reported utilization of Fe₂O₃-SiO₂ catalyst in the 1605 conversion of JCO to FAME. The catalyzed reaction displayed 95.6 % FAME yield under the 1606 1607 optimized reaction conditions. When Fe loading was raised from 0.07 to 2.1 wt. %, the acidity 1608 of the catalyst increased drastically. Fe-oxide species scattered over the SiO₂ surface were recognized as the active sites. In the meantime, ZnAl₂O₄/ZnFe₂O₄ catalyst was also examined 1609 for the transformation of oil such as sunflower oil, WCO and JCO.³³¹ During the reaction, the 1610 Zn 3d electrons of ZnAl₂O₄ and ZnFe₂O₄ spinels were likely to take part in the electronic 1611 1612 excitation, thereby Zn 3d electrons are probably going to undertake a vital job to enhance the catalyst reactivity. In 2012, Xie et al. ³³² synthesized SnO₂-SiO₂ catalyst by loading 8 wt. % Sn 1613 1614 onto SiO₂ followed by calcination (550 °C) and exploited it in the transformation of soybean 1615 to FAME yielding 81.7 % under the optimal reaction conditions.

Impregnation followed by calcination (600 °C) was used to synthesize Fe-Mn-1616 MoO₃/ZrO₂ catalyst, which could provide a high 95.6 \pm 0.15 % yield of FAME.³³³ It is 1617 interesting to observe that ZrO_2 and MoO_3/ZrO_2 gave a lower FAME yield of 48.6 ± 1.14 and 1618 73.0 ± 0.25 % respectively. The high activity of Fe-Mn-MoO₃/ZrO₂ catalyst is attributed to 1619 high surface area (49.5 m^2g^{-1}) and availability of huge active sites (2411 μ molg⁻¹) in the 1620 catalyst. Moreover, catalyst reusability examination revealed that it is stable up to 6 progressive 1621 1622 reaction cycles of transesterification of WCO without loss in its efficiency. On the other hand, enhanced catalytic activity was observed in a mixed metal oxide of WO₃/SnO₂ in the soybean 1623 oil transformation in comparison with the individual WO₃ and SnO₂ species.³³⁴ The bonding 1624 1625 of WO₃ with SnO₂ was believed to upgrade WO₃/SnO₂ acidity. The catalyst is highly stable and was reused up to 4 times without much depreciation in the biodiesel yield. 1626

Further, Xie *et al.* ³³⁵ studied 30 wt. % WO₃ loading on AlPO₄ catalyst and recorded a good 72.5 % conversion to biodiesel under the optimized reaction condition. The high catalyst reactivity was attributed to the existence of WO₃ that enhanced the surface acid sites. Similarly, Amani *et al.* ³³⁶ reported a series of Mn_{3.5x}Zr_{0.5y}Al_xO₃ catalyst for the transformation of WCO

to FAME. The Mn_{1.4}Zr_{0.35}Al_{0.6} O₃ catalyst demonstrates better catalyst reactivity as far as 1631 FAME yield (>93 %) than the $Mn_{1,4}Zr_{0.35}O_3$ catalyst (52.8 %). The bonding between metals in 1632 the crystal structure efficiently influenced the catalyst reactivity. It was observed that the 1633 amphoteric component of the Al developed the surface region of the catalyst and framed a 1634 complex structure with other metal oxides, though Mn alternated the morphology and catalyst 1635 basic site density. In the meantime, Zhang et al. ³³⁷ reported Zr-Mo mixed metal oxide 1636 functionalized with various carboxylic acids, for example, lauric acid, stearic acid, palmitic 1637 acid and myristic acid for biodiesel production from oleic acid. The modification of Zr-Mo 1638 1639 metal oxide using such monofunctional carboxylic acids enhances the catalyst acidity and surface area, thus upgraded the rate of the reaction. They also reported that among all, stearic 1640 acid-functionalized Zr-Mo metal oxide showed the best result with the maximum oleic acid 1641 conversion of 94. 2 %. Catalyst reusability test revealed that the catalyst is stable up to 6 1642 progressive cycles. Similarly, WCO was utilized for the FAME production using ferric-1643 manganese doped tungstate molybdena nanoparticles (FMWMo).³³⁸ The dopants Fe-Mn 1644 enhances the surface area, density of acidic sites and the stability towards the esterification of 1645 WCO. A maximum yield of 92.3±1.12 % methyl ester was achieved under the optimized 1646 reaction conditions. 1647

1648

No.	Catalyst	Feedstocks	^a Conditions	Yield (%)	Ref.
1.	Fe ₂ O ₃ -SiO ₂	Jatropha oil	218:1, 15, 220, 180	95.6	330
2.	$ZnA_{12}O_4/ZnFe_2O_4$	Sunflower	9:1, 5, 180, 600	>90	331
		oil, WCO,			
		Jatropha oil			
3.	SnO ₂ -SiO ₂	Soybean oil	24:1, 5, 180,300	81.7	332
4.	Fe-Mn-MoO ₃ /ZrO ₂	WCO	25:1, 4, 200, 300	95.6 ± 0.15	333
5.	WO ₃ -SnO ₂	Soybean oil	30:1, 5, 110, 300	79.2	334
6.	WO ₃ (30 wt.	Soybean oil	30:1, 5, 180, 300	72.5	335
	%)/AlPO ₄				
7.	$Mn_{1.4}Zr_{0.35}Al_{0.6}O_{3}$	WCPO	14:1, 2.5, 150, 300	>93	336
8.	Zr-Mo	Oleic acid	10:1, 4, 180, 120	94.2^{b}	337
9.	FMWMo	WCO	25:1, 6, 200, 480	92.3±1.12	338

Table 18: Different types of solid acid catalysts for FAME production.

^aMethanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).

1650 7.2.4 Sulfonated carbon-based catalyst

In the last few decades, various carbon materials with different shape, size, and 1651 structures have been developed by several research groups and utilized as low-cost catalysts 1652 for diverse industrial processes including transesterification.³³⁹ Currently, sulfonated carbons 1653 i.e. SO₃H-functionalized acidic carbon materials are considered as a new group of the metal-1654 1655 free solid acid catalyst described by their original carbon structure and Brønsted acidity equivalent to concentrated H₂SO₄. Sulfonated carbon acid catalysts can be easily prepared by 1656 processes like incomplete carbonization of aromatic compounds in concentrated H₂SO₄ ³⁴⁰ or 1657 sulfonation of incompletely carbonized natural organic matter, such as sugar ^{341–343} and 1658 cellulosic materials.^{344,345} Sulfonation can also be achieved by treating carbon material with a 1659 sulfonating reagent such as gaseous SO₃, ClSO₃H, p-toluenesulfonic acid, 4-benzenediazonium 1660 sulfonate or SO₃H-containing aryl diazoniums etc.³⁴⁶⁻³⁴⁹ These materials possess promising 1661 features such as biogenic, environment-friendly, lower production costs, distinctive surface 1662 1663 chemistry, high chemical and thermal stability.

The acid-catalyzed chemical reactions such as saccharification, esterification, 1664 transesterification and acetylation are vital operations commonly used for the valorization of 1665 biomass or their components to useful products in various food, fuel and chemical industries.³⁵⁰ 1666 1667 The functionalized acidic carbons from inexpensive sources including natural organic carbon matter such as sugars, carbohydrates, cellulosic materials, lignin have been achieved by several 1668 researchers.^{351–353,341} Besides this, agro waste such as husk, straw, seed cover, cow manure, 1669 corn cob ^{342,343,354,355,} carbonaceous waste from industries (char, oil pitch, coke, glycerol, etc.) 1670 ^{356,357,346,348} and polymer resins ^{349,358,359} were also used. Various carbon supports (e.g. zeolite-1671 templated carbons, mesoporous carbons, active carbon, etc) ^{352,353,360,361} and more recently 1672 nanostructured carbons such as graphene, graphene oxide, carbon nanotubes, and carbon 1673 dots) $^{362-367}$ have been exploited for the same purpose. 1674

1675 Over the last few years, there is growing interest from researchers towards the application of sulfonated carbon-based catalysts due to their noteworthy efficacies mentioned 1676 earlier. Many reports are available which demonstrated the efficient nature of sulphonic acid-1677 functionalized catalyst in biodiesel production using various feedstocks.^{362,356,367} One of the 1678 1679 reports presented the synthesis of organosulfonic acid (i.e. propylsulfonic and arenesulfonic groups) functionalized mesoporous silicas through a simple one-step process. The synthesized 1680 novel catalysts that possessed propylsulfonic groups and arenesulfonic groups were further 1681 evaluated for their catalytic efficacy in the esterification of fatty acid with methanol to produce 1682 methyl esters and the authors also compared the efficacy of these heterogeneous catalysts with 1683

a variety of commercially available catalysts such as sulfuric acid, p-toluene sulfonic acid, 1684 Nafion NR50, Amberlyst-15, etc. The obtained results indicated that the organosulfonic acid-1685 functionalized mesoporous silica catalysts showed the highest reactivity compared to all the 1686 above mentioned commercial solid acid catalysts in fatty acid esterification process. Moreover, 1687 it was also recorded that the efficiency of these catalysts was largely depended on important 1688 factors such as the median pore diameter of the catalyst and the acidic strength of the 1689 1690 organosulfonic acid group present over this catalyst. Considering these findings, it can be proposed that there is a huge potential to developed catalysts using organic-inorganic 1691 mesoporous materials.³⁶³ In general, the activity of carbon-based catalysts upon fatty acid 1692 (C16-C18) esterification to produce biodiesel primarily depends on three primary factors: (i) -1693 SO₃H group density, (ii) total acid density, and (iii) porosity. Different sulfonated carbon-based 1694 acid catalyst utilized for FAME production are listed in Table 19. Numerous reported catalysts 1695 demonstrated promising outcome in the (trans)esterification of biodiesel feedstocks with high 1696 FFA and afforded > 85% FAME yield. In the meantime, several investigations had been 1697 conducted using model acids (e.g. palmitic acid, oleic acid, etc. which are the major 1698 1699 components of vegetable oil as reactant) that mainly focused on esterification reaction.

In a pioneering work towards the preparation of biomass-based sulfonated carbon catalyst, 1700 Toda et al. ³⁶⁴ synthesized sulfonated carbon catalyst by partial carbonization of sugar followed 1701 by sulfonation in fuming H₂SO₄. The prepared catalyst consists of sheets of indistinctive carbon 1702 1703 having a high amount of sulfonic groups along with hydroxyl and carboxyl as a minor group (Figure 28). The highly active bio-based carbon catalyst was utilized for transformation of oleic 1704 1705 and stearic acid to FAME via esterification. Apart from -SO₃H group, presence of -OH and -1706 COOH groups in the catalyst greatly enhance the catalytic activity and make it highly water 1707 tolerant. The successful incorporation of -SO₃H group and formation of carbonized materials are can be easily confirmed by using FT-IR and ¹³C MAS NMR analysis respectively, as 1708 depicted in Figure 33.³⁶⁸ FT-IR spectra (Figure 29a) displayed two bands at 1040 and 1377 cm⁻ 1709 ¹ (in SO₃H), ascribed to SO₃ and O=S=O stretching vibration, respectively, suggests the 1710 existence of -SO₃H groups. ¹³C MAS NMR (Figure 29b) depicted three major peaks at 130, 1711 155, and 180 ppm, ascribed to polycyclic aromatic carbon atoms, phenolic OH, and COOH 1712 groups, respectively. 1713

In another work, Hara *et al.* ³⁵⁶ examined sulfonated carbon catalyst in biodiesel synthesis.
The findings showed that amorphous carbon material containing sulfonic acid groups enhances
the catalytic performance, thus displayed extraordinary reactivity in
esterification/transesterification reactions in comparison to the ordinary solid acid catalyst.



Figure 28: Synthesis of sulfonated carbon catalyst from sucrose and D-glucose.

Reproduced from ref. [364].

1718



Figure 29: FT-IR (a) and 13C MAS NMR (b) spectrum for the sulfonated carbon catalyst originated from cellulose. Reproduced from ref. [368].

1719

Likewise, Nakajima et al.³⁶⁵ synthesized amorphous cellulose-originated carbon solid 1720 1721 acid (CCSA) catalyst and exploited it in the transformation of oleic acid to FAME, and 1722 observed a 99.9 % yield under the optimized conditions. The carbon material displayed much higher catalytic activity in esterification reaction in comparison to ordinary solid acid catalysts 1723 1724 examined, such as niobic acid, Amberlyst-15 and Nafion NR50. Interestingly, those CCSA catalyst prepared at a lower carbonization temperature before subjected to sulfonation gave a 1725 1726 lot better biodiesel yield as compared to those prepared at higher carbonization temperature. This is attributed to the huge amount of -OH and -COOH groups in the former which enhanced 1727 1728 its acidic nature, thereby its catalytic activities (Figure 30). The catalyst reactivity remains intact after 10 progressive cycles. 1729



Figure 30: Schematic structures of SO₃H-bearing CCSA materials carbonized at below 723 K (A) and (B) above 823 K. Reproduced from ref. [365].

1730

1731 Simultaneous carbonization and sulfonation in a one-pot synthesis of solid acid catalyst directly from biomass have also been explored by various experts as it is straight forward, cost 1732 and time-efficient. Malins et al. 366 synthesized C-SO₃H via simultaneous carbonization-1733 sulfonation and utilized it for FAME production. The C-SO₃H catalysts with the highest density 1734 of SO₃H groups (0.81 mmol Hb/g) were prepared using optimal reaction conditions. It was 1735 noted that under optimized reaction conditions 96.5 % of FAME was recorded. Interestingly, 1736 1737 the catalyst has great stability and can be easily recovered and reused for subsequent reaction cycles. Moreover, in the comparative study of esterification reactions of rapeseed oil fatty 1738 acids, the prepared catalyst exhibited similar reactivity to Amberlyst-15. 1739

Another recent report proposed a synthesis of the heterogeneous sulfonated catalyst using activated carbon to overcome several problems like drastic reaction conditions such as very high temperature, pressure, longer reaction time and expensive overall process cost. The above-mentioned activated carbon catalyst was prepared from corncobs as a precursor and utilized in the microwave-assisted conversion of soybean oil with ethanol to FAME. In this study, about 88.7 % yield of pure biodiesel was reported at 0-600 W of microwave power.
Moreover, the catalyst was reused up to 5 cycles.³⁵⁵ Figure 31 represents the schematic
illustration of the application of activated carbon-based catalyst in the transesterification of
various oils using methanol.





Figure 31: Schematic representation of transesterification of various oils using activated carbon-based catalysts.

1750

In 2009, Yuan *et al.* ³⁶⁹ examined the application of solid acid catalyst originated from 1751 sulfonated activated carbon (H₂SO₄/C) for catalyzing transesterification of castor oil and 1752 methanol as feedstock. Melero et al. 370 synthesized sulfonic acid-modified mesostructured 1753 (SAM) catalyst and studied their efficacy in crude vegetable oils transformation to FAME. The 1754 1755 results obtained noted that this catalyst has the ability to yield 95 wt. % pure FAME and oil transformation close to 100 %. Despite the presence of FFAs, this catalyst displayed 1756 significantly high activity toward simultaneous esterification and transesterification reactions. 1757 Similarly, Zuo et al. ³⁷¹ developed various sulfonic acid functionalized mesoporous SBA-15 1758 catalysts and tested their catalytic activity in the microwave-assisted conversion of soybean oil 1759 and 1-butanol to biodiesel. The authors observed that the catalytic efficacy of these catalysts 1760 mainly depends on the acid strength and not on the number of acid sites. Further, propyl-SO₃H 1761 and arene-SO₃H functionalized SBA-15 catalysts found to have comparatively better reactivity 1762

1763 in the transesterification process. However, perfluoro-SO₃H functionalized SBA-15 catalyst displayed leaching of the active sites in each progressive cycles, thus reactivity decreased. Shah 1764 *et al.* ³⁷² demonstrated esterification of FFAs in acid oil (which is a byproduct of oil refining) 1765 using sulfonic acid-functionalized silica (SiO₂-Pr-SO₃H) catalyst to prepare biodiesel. Further, 1766 the authors optimized various reaction conditions such as temperature, reaction time, catalyst 1767 concentration, and M/O molar ratio, etc. which usually affects the conversion to FAME. A high 1768 1769 conversion (i.e. 96.78 % conversion after 8 h was reported at optimized conditions) can be 1770 achieved using these solid acid catalysts.

Moreover, in recent past Varyambath et al. 373 developed different sulfonic acid-1771 functionalized organic knitted porous polyaromatic microspheres (OPPSO₃H) utilizing pyrene, 1772 1773 anthracene, and naphthalene as monomers via Friedel-Crafts alkylation, followed by crosslinking reactions. Further, these heterogeneous catalysts were utilized for the 1774 transformation of long-chain fatty acids and triglycerides to biodiesel. These solid acid 1775 catalysts were found to be very promising for biodiesel synthesis as they showed excellent 1776 surface acidity. In addition, several other sulphonic acid-functionalized catalysts were 1777 successfully developed and exploited in the production of biodiesel. In this context, Shagufta 1778 et al. 374 reviewed all such sulphonic acid-functionalized in catalysts esterification and 1779 1780 transesterification reactions. This review can be consulted for more detail information.

Yu et al.³⁷⁵ studied biodiesel production by exploiting coal-based acid catalysts and 1781 reported an oleic acid conversion of 97.6 % under the optimal reaction conditions. Similarly, 1782 Tang and Niu ³⁷⁶ investigated the synthesis of carbon-based solid acid catalysts from bamboo 1783 1784 through partial carbonization and sulfonation approach. The microstructure of catalyst was 1785 activated by phosphoric acid impregnation. The catalyst afforded biodiesel yield 97.3 % at 1786 optimum conditions which decreased to 83.7 % in fourth reaction cycles. In addition, biodiesel production from oleic acid was reported using sulfonated activated carbon from bamboo.³⁷⁷ A 1787 1788 sulfonated carbonaceous material synthesized via single-step hydrothermal sulfonation of glucose has also been used as a catalyst for esterification of waste cooking oil to produce 1789 biodiesel.³⁷⁸ FESEM images of the carbonaceous material (C) (Figure 32a) and the sulfonated 1790 carbonaceous material (C-SO₃H) (Figure 32b) showed microsphere and microsphere with an 1791 attached sulfonic group on the surface respectively. The catalyst showed great stability with 1792 93.4 % FAME yield under the optimized reaction conditions. 1793

1794



Figure 32: FESEM images of a) C and b) C-SO3H. Reproduced from ref. [378].

1795

Guan et al. ³⁷⁹ synthesized sulfonated multi-walled carbon nanotube (S-MWCNT) for 1796 the conversion of triglyceride to FAME in 97.8 %. The high catalytic reactivity is because of 1797 high surface area (198.9 m² g⁻¹), high porosity (10-15 nm) and high acid sites. Similarly, 1798 sulfonated carbonaceous material from starch was utilized as a solid catalyst for esterification 1799 of PFAD.³⁸⁰ A novel, efficient, inexpensive and environment-friendly acid catalyst was 1800 1801 synthesized from coconut meal residue (CMR). CMR-DS-SO₃H catalyst was prepared by a one-step direct in-situ carbonization in concentrated H₂SO₄ and reported for the transformation 1802 of waste palm oil (WPO) to biodiesel. The prepared sulfonated catalyst has acid density 3.8 1803 mmol/g, surface area 1.33 m²/g and means pore volume 0.31 cm³/g. The results obtained 1804 recorded a high yield of 92.7 % biodiesel from WPO.³⁸¹ Moreover, Wang et al. ³⁸² investigated 1805 the application of monodispersed hollow carbon/silica solid acid catalyst HS/C-SO₃H, which 1806 was prepared by chemical activation approach, in the esterification of oleic acid with methanol 1807 to produce biodiesel. 1808

Besides this, another kind of sulfonated functionalized carbon material i.e. 1809 sulfonated ordered mesoporous carbon (SOMC) catalyst showed promising biodiesel 1810 production (73.59 % yield).³⁸³ Recently, sulfonated acid catalyst obtained from corncob (SO₄²⁻ 1811 /corncob), has been reported as an excellent catalyst for conversion of oleic acid to obtain 1812 methyl oleate in good yield (> 80 % after 8 h at 60 °C).³⁸⁴ Mahdavi and Darabi ³⁸⁵ prepared 1813 1814 sulfonated carbon catalyst by treatment of sucrose and concentrated H₂SO₄ at high temperature (sulfonation and carbonization approach). The synthesized C-SO₃H catalyst was further utilized 1815 1816 for the conversion of oleic acid to FAME in 93.04 % yield. Moreover, a solid acid catalyst generated from sulfonation of microcrystalline cellulose powder was successfully applied for 1817 oleic acid esterification, showed 99.9 % biodiesel yield under the optimized reaction 1818

1819 conditions.³⁸⁶ In another investigation, waste cooking oil was transformed to produce biodiesel 1820 utilizing an environmentally benign sulfonated carbon microsphere catalyst.³⁸⁷ The catalyst 1821 with surface area 86 m²/g and acidity 1.38 mmol/g was developed by consecutive hydrothermal 1822 carbonization and sulfonation of xylose. Using this catalyst, biodiesel yield of 89.6 % was 1823 recorded at optimal reaction conditions. Catalyst reusability report revealed that in each cycle 1824 biodiesel yield was reduced by 9 %. Furthermore, sulfonated carbon-based solid acid catalyst 1825 was also utilized for the transformation of PFAD ³⁸⁸ and *Mesua Ferrea* Linn oil ³⁸⁹ to biodiesel.

To bring down the cost of biodiesel production, several sulfonated raw biomass has been prepared and investigated for their catalytic activities. In this line, a sulfonated solid-acid catalyst obtained from coconut shells ($SO_4^{2-}/coconut$ shell) reported 88.03 % biodiesel yield.³⁹⁰ In the same vein, oil palm trunk/ sugarcane bagasse,³⁹¹ corn straw,³⁹² bamboo,³⁹³ Jatropha curcas seed,³⁹⁴ bio-glycerol,³⁹⁵ glycerol,³⁹⁶ microalgae residue,³⁹⁷ oil cake waste,^{398,399} de-oiled waste cake, ⁴⁰⁰ de-oiled canola meal-SO₃H, ⁴⁰¹ pine chip char ⁴⁰² and biochar ^{403,404} are reported as a catalysts for FAME production.

1833

No.	Catalyst	Feedstock	^a Conditions	Yield (%)	Ref.
1.	Sulfonated	Oleic acid	10:1 ^{<i>f</i>} , 7.4, 80, 240	NR	364
	sugar				
2.	Sulfonated	Oleic acid	2.92:1 ^c , 17.2, 95,	99.9	365
	carbon		240		
3.	ACPhSO ₃ H	Rapeseed oil	20:1, 10, 65, 420	95	366
4.	Sulfonated AC	Soybean oil	6:1, 20, 75, 20	88.7	355
5.	H_2SO_4/C	Castor oil	12:1, 5, 65, 60	94	369
6.	SAM	Vegetable oil	10:1, 6, 180, 120	95	370
7.	SO ₃ H/SBA-15	Soybean oil	6:1, 5, 190, 30	90	371
8.	SiO ₂ -Pr-	Acid oil	15:1, 4, 100, 480	96.78 ^b	372
	SO ₃ H				
9.	OPPSO ₃ H	Soybean oil	50:1 ^c , 10, 70, 600	93 ^{<i>b</i>}	373
10.	Coal based	Oleic acid	10:1, 8, 240, 67	97.6 ^{<i>b</i>}	375
	solid acid				

Table 19: Different sulfonated carbon-based acid catalyst used for biodiesel production.

11.	Sulfonated carbon-based	Oleic acid	10:1,10,65, 120	97.3	376
	solid acid				
12.	Sulfonated	Oleic acid	7:1 ^{<i>f</i>} , 12, 180, 85	96 ^b	377
	activated		, , ,		
	carbon				
13.	C-SO3H	Waste cooking	20:1, 10, 60, 180	93.6	378
		oil			
14.	Sulfonated	Triglycerides	10:1 ^{<i>f</i>} , 3.7, 60, 150	97.8 ^{<i>b</i>}	379
	multiwalled				
	carbon				
	nanotube				
15.	ICS-SO3H	Palm fatty acid	10:1, 2, 180, 75	90.4	380
		distillate			
16.	CMR-DS-	Waste palm oil	12:1, 5, 65, 72	92.7	381
	SO ₃ H				
17.	HS/C-SO ₃ H	Oleic acid	5:1, 3.5, 80, 300	96.9 ^{<i>b</i>}	382
18.	SOMC	Oleic acid	10:1, 3.5, 80, 600	73.59 ^b	383
19.	SO4 ²⁻ /corncob	Oleic acid	15:1, 5, 60, 480	>80	384
20.	C-SO ₃ H	Oleic acid	10:1, 1.5, 67, 120	93.04	385
21.	C-SO ₃ H	Oleic acid	16:1, 17, 95, 240	99.9	386
22.	C-SO ₃ H	WCO	10:1, 10, 110, 240	89.6	387
23.	C-SO ₃ H	PFAD	15:1, 2.5, 80, 240	95.3 ^{<i>b</i>}	388
24.	C-SO ₃ H	Mesua Ferrea	40:1, 5, 120, 1440	97.79	389
		Linn oil			
25.	Coconut shell-	Palm oil	30:1, 6, 60, 360	88.03	390
	SO ₃ H				
26.	Oil palm	Waste Oil	1.17 mL/min, 12,	80.6/83.2	391
	trunk/		130, 240		
	Sugarcane				
	bagasse-SO ₃ H				
27.	Corn straw-	Oleic acid	3:1, 3, 60, 240	92	392
	SO ₃ H				

28.	Bamboo-SO ₃ H	Oleic acid	7:1 ^{<i>f</i>} , 2, 90, 360	98.4	393
29.	Jatropha	JCO	12:1, 7.5, 60, 60	99.13 ^b	394
	curcas		99.13		
	Seed-SO ₃ H				
30.	Bio-glycerol	Karanja oil	45:1, 20, 160, 240	99.5	395
31.	Glycerol	Palmitic acid	9.7:1 ^{<i>c</i>} , 10, 65, 240	99 ^b	396
32.	Microalgae	Oleic acid	NR,5, 80, 720	98 ^b	397
	residue				
33.	Oil cake	JCO/ M.	43:1, 5, 80, 480	99	398
	waste-SO ₃ H	ferrea			
		L. oil			
34.	Oil cake	Oleic acid	12:1, 20, 60, 120	94 ^{<i>b</i>}	399
	waste-SO ₃ H				
35.	De-oiled waste	Oleic acid	20:1, 3, 64, 600	97 ^b	400
	cake				
36.	De-oiled	Oleic acid	60:1, 7.5, 65, 1440	93.8 ^b	401
	canola				
	Meal-SO ₃ H				
37.	Pine chip char	Palmitic acid	6:1, 5,55-60, 300	97	402
38.	Biochar	Canola oil	15:1 ^{<i>f</i>} , 5, 65, 1440	92	403
39.	Biochar	Canola oil,	30:1, 5, 315, 180	48	404
		oleic acid			

^aMethanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).

^bConversion

1836 ^{*f*}Ethanol to oil molar ratio

1837 NR: Not reported.

1838

1839 **7.3 Enzyme catalyst**

In recent years, enzyme catalysts are widely examined for the production of biodiesel as they produce high-quality biodiesel, improve the product separation process, mild reaction conditions and most importantly their ecological benignness (Table 20). ^{405,406} Besides, they do not form soap with FFA contrary to the alkaline catalyst and hence can be utilized in the biodiesel production in industrial scale.

In biocatalyst mediated reactions, usually enzymes can be used in free form or can be 1845 immobilized on a matrix, i.e. immobilized lipase.⁴⁰⁷ The free enzymes are more sensitive 1846 towards the pH, temperature and impurities of reactants which may create obstacle in 1847 bioprocesses. However, these problems can be overcome by immobilizing enzyme onto 1848 different types of support materials.⁴⁰⁸ The commonly adopted immobilization methods for 1849 biological processes include entrapment, adsorption and covalent bonding. Among these 1850 1851 techniques, entrapment found to be effective offering greater advantages such as ease of process scale up, higher stability of enzyme and longer enzymatic activity retention.^{409, 410} 1852 Mostly lipase enzymes obtained from microbial sources have been used for biodiesel 1853 production ⁴¹¹ proposed the method for large scale production of bacterial or fungal lipases due 1854 to their extracellular nature. Moreover, lipases obtained from diverse plant sources are also 1855 considered as the potential substitute for catalysing the transesterification process.⁴¹² The 1856 advantages associated with lipase catalyst over the other catalyst used in biodiesel production 1857 are superior quality and higher yield of biodiesel, free from soap formation, lower reaction 1858 temperature and can work on variety of feedstock.⁴¹³ 1859

Compared to homogeneous and heterogeneous catalysts, enzymatic catalysts are less 1860 studied hence scanty of literature is available when compared with reports on above-mentioned 1861 1862 two catalysts. However, the high cost of the free lipase catalyst along with the limited longterm use has led to the exploitation of immobilized lipase catalyst to reduce the cost of the 1863 1864 catalyst and its reusability. Apart from that, immobilized lipase catalyst showed greater tolerance to pH variation, high thermal stability and high substrate selectivity.^{414,415} To date, 1865 large number of literature are available in the field of biodiesel production using both free ^{416–} 1866 ⁴¹⁸ and immobilized ^{419–422} enzyme catalyst. 1867

Recently, Jayaraman *et al.* ⁴²³ demonstrated the lipase enzyme mediated 1868 transesterification of waste cooking oil (WCO) and reported 88% of biodiesel yield. Marín-1869 Suárez et al. 424 demonstrated the lipase-catalyzed transesterification of low quality fish oil 1870 through the process optimization, moreover, the reusability of enzyme was also studied. 1871 Authors evaluated the efficacy of commercially available immobilized enzymes such as 1872 Liposome RM IM, Lipozyme TL IM and Novozym 435⁴²⁵ for biodiesel production from waste 1873 fish oil. The results obtained revealed that Novozym 435 showed maximum catalytic activity 1874 resulting in highest yield of FAME i.e. 82.91 wt. % and enzyme can be reused for about ten 1875 successive cycles. In another study, it was reported that immobilized lipase (Epobond P. 1876 *cepacia*) employed in transesterification of waste vegetable oil reported to achieve ester yield 1877 of 46.32%.⁴²⁶ Similarly, the Candida cylindracea lipase immobilised on the functionalised 1878

1879 activated carbon was tested as a catalyst in the transesterification of Jatropha curcas oil. It was found that free fatty acid yield of 78% was achieved at optimized reaction conditions and 1880 1881 further biocatalyst was found stable up to four consecutive cycles of transesterification.⁴²⁷ Beside, lipase obtained from plant source like rice bran lipase produced 83.4 wt.% FAME yield 1882 1883 from rice bran oil under optimized conditions.⁴²⁸ Moreover, Muanruksa and Kaewkannetra ⁴²⁹ examined the biodiesel production from 1884 sludge palm oil (SPO) via two steps of extraction and enzymatic esterification. The 1885 immobilised Rhizopus oryzae lipase on alginate-polyvinyl alcohol (PVA) beads were used for 1886 conversion of FFAs from SPO to fatty acids methyl esters (biodiesel). It was found that at 1887 optimum condition, the maximum biodiesel yield of 91.30% was achieved and biocatalyst 1888 showed higher stability and catalytic efficiency up to 15 cycles. It is reported that enzymatic 1889 transesterification reaction to produce biodiesel is the slowest pathway among all the known 1890 transformations. Taking into account, application of ultrasonic in the enzyme-catalyzed 1891 transesterification improves the reaction rate and hence reduces the reaction time.^{414,422} Thus, 1892 it can be a promising technique for the industrial-scale production of biodiesel in a very short 1893 1894 time.

1895

Table 20: Different	enzyme	catalysts 1	reported for	the prod	uction of	biodiesel
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S1.	Catalyst	Feedstock	^a Conditions	Yield	Ref.
No					
1	Lipase immobilized on	<mark>Hybrid non</mark>	<mark>6:1°,10, 50,1440</mark>	<mark>~78</mark>	<mark>407</mark>
	biosupport beads	edible oils			
2	Lipase	WCO	3:1, 1.5, 65, 240	88	423
3	Thermomysis	Rubber seed	4:1, 5, NR, 65	92.83	416
	Lanugonosus Lipase	oil			
4	CalleraTM Trans L	Soybean oil	4.51:1, 1.45, 35,	96.9	417
	lipase		1440		
5	Lipase@AC	Sardine oil	9:1, 10, 30, 600	94.5	418
6	Lipase@APTES-	Aspergillus	4:1, 300 ^b , 45, 240	84	419
	Fe ₃ O ₄	lipid			
7	Lipase@ZIF-67	Soybean oil	6:1, 10, 45, 3600	78	420
8	Lipase@[bmim][PF ₆]	Food	6:1, 40, 50, 840	72	421
		compost			

9	Lipase@[bmim][NTf ₂]	Food compost	6:1, 40, 50, 840	48	421
10	Lipase@Immobead	Blended non-edible oils	7.64:1, 3.55, 36, 120	94	422
<mark>11</mark>	Novozym 435 lipase	Waste fish <mark>oil</mark>	35.45:1 ^d , 50, 35, <mark>480</mark>	82.91 wt %	<mark>424</mark>
<mark>12</mark>	Novozym 435 lipase	BSFL fat	14.64:1 [°] , 17.58,39.5,720	<mark>96.97</mark>	<mark>425</mark>
<mark>13</mark>	Immobilized lipase (Epobond- <i>Pseudomonas cepacia)</i>	Waste vegetable oil	3:1 ^d ,3, 37,90	<mark>46.32</mark>	<mark>426</mark>
<mark>14</mark>	Immobilized <i>Candida</i> cylindracea lipase	<i>Jatropha</i> curcas oil	HR,8,40,1440	<mark>78</mark>	<mark>427</mark>
<mark>15</mark>	Immobilised <i>Rhizopus</i> oryzae lipase	sludge palm oil (SPO)	3:1,5,40,240	<mark>91.30</mark>	<mark>428</mark>
<mark>16</mark>	Lipase (from rice bran)	<mark>Rice Bran</mark> oil	<mark>6:1, NR,40, 17280</mark>	<mark>83.4</mark> wt %	<mark>429</mark>

^aMethanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time
(min).NR: Not reported.

1898 ^{*b*}miligram

1899

1900 **7.4 Bifunctional solid catalysts**

1901 Despite the high reactivity of the basic solid catalyst towards biodiesel production, they 1902 are not an effective catalyst for transesterification of oils having a high amount of FFA as such 1903 catalyst are highly sensitive to the FFA, which leads to soap generation and thus interfere in 1904 the separation process of glycerol from biodiesel. On the other hand, solid acid catalysts are insensitive to the FFA content and esterify waste oils or low-cost oils without any requirement 1905 1906 of pretreatment. However, water formed during the course of the reaction may lead to the decomposition of triglycerides to diglycerides, resulting in the formation of more FFA and 1907 catalyst leaching.⁴³⁰ Taking these difficulties into account, developing a new type of solid 1908

1909 catalysts that possess dual characteristics such as solid acidic character to tackle the FFA and solid basic character for easy transesterification of triglycerides to FAME has been a recent 1910 interest in the realm of biodiesel research. To date, numerous bifunctional catalysts are reported 1911 for the FAME production (Table 21), which will be discussed in this section. Farooq et al. 78 1912 developed a bifunctional Mo-Mn/y-Al₂O₃-MgO catalyst and utilized it for the simultaneous 1913 esterification/transesterification of WCO having FFA content of 3.27 mg KOH/g. The authors 1914 1915 investigated the effect of MgO loading (5-20 wt. %) on its catalytic activity and found that 15 wt. % MgO loading showed highest catalytic activity with 91.4 % biodiesel yield under the 1916 1917 ideal reaction conditions. Moreover, the catalyst showed excellent stability towards the biodiesel production from WCO as it is stable up to 8 progressive reaction cycles without any 1918 major loss of its activity. In another study, Cu/Zn/y-Al₂O₃ catalyst was utilized for the 1919 simultaneous esterification/transesterification of WCO for production of FAME via RSM.⁴³¹ 1920 The effect of Cu/Zn wt. % ratio and calcination temperature on the catalytic reactivity was also 1921 examined and found that 10:90 Cu/Zn wt. % ratio and 800 °C calcination temperature showed 1922 88.82 % FAME yield. The authors also studied the structure and particle size of the synthesized 1923 catalyst via TEM micrographs (Figure 33). Figure 33a displayed that the average diameter of 1924 the particles lies in between 4-6 nm. The lattice fringes measured from Figure 33b, c and d are 1925 1926 0.201, 0.282 and 0.242 nm, matched with the hkl planes (400), (220) and (311) of alumina respectively. The lattice fringes in Figure 33e is 0.240 nm fitted with the hkl plane (200) of 1927 1928 CuO and lattice fringes 0.281 nm (Figure 33f) fitted with the ZnO plane (100). Similarly, biodiesel production from WCO was reported using diverse bifunctional solid catalysts such 1929 as Mg/MCM-41, 432 γ -Al₂O₃-CeO₂. 433 KAcZX 434 and Sr/ZrO₂ 435 etc. 1930



Figure 33: TEM micrograph for Cu/Zn(10:90)/ γ -Al₂O₃-800 °C (a) HRTEM displayed the lattice fringes of (b) Al₂O₃ (400), (c) Al₂O₃ (220), (d) Al₂O₃ (311), (e) CuO (200) and (f) ZnO (100). Reproduced from ref. [431].

1931

Nizah *et al.* ⁴³⁶ synthesized a bifunctional catalyst Bi₂O₃-La₂O₃ *via* wet impregnation 1932 procedure and employed it for one-pot esterification/transesterification of JCO having FFA 1933 1934 content of 6.1 mg KOH/g. The authors investigated the influence of Bi₂O₃ impregnation on 1935 La₂O₃ support by varying the wt. % of Bi₂O₃ in the range of 1-7 wt. % and found that 5 wt. % 1936 Bi₂O₃ impregnated on La₂O₃ showed maximum biodiesel yield of 94 %. The high catalyst reactivity is attributed to the well dispersion of Bi₂O₃ on La₂O₃ support, that directly enhanced 1937 the surface area and thus increases selectivity and rate of the reaction. Similarly, biodiesel 1938 production from JCO having a high amount of FFA was reported by using a bifunctional solid 1939 catalyst CaO-La₂O₃.⁴³⁷ The esterification/transesterification was performed in a high-1940 temperature reactor (Figure 34). The effect of Ca/La atomic ratio on the catalytic activity was 1941 1942 examined and observed that 0.8 atomic ratio of Ca/La showed maximum biodiesel yield of 98.76 % under the optimized reaction conditions. The high catalytic reactivity is because of 1943 well dispersion of CaO on the surface of La₂O₃, that led to an increase in catalyst surface area. 1944

1945 Moreover, the synthesized catalyst is chemically stable and can be used for 4 consecutive 1946 cycles.



Figure 34: Schematic diagram of a high-temperature reactor. Reproduced from ref. [437].

1947

Another study revealed the synthesis of mixed metal oxide Mn@MgO-ZrO₂ via co-1948 1949 precipitation and impregnation method and utilization of the catalyst in the FAME production from kernel oil.⁴³⁸ The efficiency of the catalyst in FAME production was tested by changing 1950 the Mg/Zr ration from 0.2-05 and found that 0.4 Mg/Zr has the optimal active sites followed 1951 by impregnation of 4 wt. % Mn to MgO-ZrO₂ composite to enhance its reactivity and displayed 1952 96.4 % biodiesel yield. The high catalyst reactivity is due to a large number of active sites and 1953 the mesoporous nature of the catalyst. Jeon et al.⁴³⁹ synthesized heteropolyacid (HPA) 1954 functionalized ZIF-8 (zeolite imidazole framework-8) to form a bifunctional catalyst for the 1955 production of biodiesel from rapeseed oil in a batch reactor. The catalyst possesses core-shell 1956 nanostructure as displayed by the TEM micrograph (Figure 35), where the rhombic 1957 dodecahedral ZIF-8 core was surrounded by thin-wrinkled HPA shell, thus enhances the 1958 1959 surface area and catalyst reactivity. Moreover, the effect of concentration of HPA for the functionalization was also tested by varying the amount of HPA such as 0.05, 0.1, 0.3 and 0.5 1960 and found that 0.1 g HPA functionalized ZIF-8 showed maximum FAME conversion of 98.02 1961

1962 % under the optimized reaction conditions. Similarly, another bifunctional catalyst 1963 organotriphosphonic acid-functionalized ferric alginate (ATMP-FA) was developed for the 1964 oleic acid esterification to produce biodiesel.⁴⁴⁰ The reaction conditions are optimized by using 1965 Box–Behnken model of RSM. Moreover, the catalyst is very stable towards the esterification 1966 reaction and can be reused for 5 consecutive cycles.



Figure 35: TEM image of HPA-ZIF-8. Reproduced from ref. [439].

1967

1968

1969 Recently, a solid bifunctional catalyst originated from bio-waste angel wing shell (AWS) via two-step processes-i) calcination of angel wing shell and ii) sulfonation of the 1970 calcined angel wing shell to produce sulfonated angel wing shell (AWS/SO₄²⁻) was reported 1971 for esterification of PFAD to produce biodiesel.⁴⁴¹ The sulfonation procedure increases the 1972 surface area of bare AWS from 3.88 to 6.53 m^2g^{-1} , thus enhanced the catalytic reactivity. The 1973 authors tested the influence of sulfuric acid concentration by varying the sulfuric acid amount 1974 from 3-11 M and found that sulfonation with 7M sulfuric acid showed 98 % FAME yield. The 1975 authors also checked the reusability of the catalyst and observed blockage of the active sites of 1976 the catalyst after 2nd consecutive cycles; which necessitate pretreatment of the spent catalyst to 1977 increase its reusability. In addition, a coordinated polymer of Zn, $[Zn(4,4'-bipy)(OAc)_2]_n$ was 1978 tested for the soybean oil transformation to FAME.⁴⁴² The catalyst showed excellent reactivity 1979 1980 and showed 98 % FAME yield under the optimized reaction conditions. The authors reported that the high reactivity of the catalyst is attributed to the bipyridine present in the catalyst. In 1981 another study, the conversion of canola oil to FAME was reported using potassium 1982 impregnated titania (K/TiO₂).⁴⁴³ The addition of K on the surface of titania increases the surface 1983

energy from 86 to $102 \text{ m}^2/\text{g}$, thus enhanced the catalytic activity. The authors investigated the effect of K loading on catalytic activity and found that 20 wt. % K loaded titania was optimum and showed 100 % conversion of canola oil to biodiesel.

No	Catalyst	Feedstocks	^a Conditions	Yield	Ref.
				(%)	
1	Mo-Mn/y-Al ₂ O ₃ -	WCO	27:1, 3, 100, 240	91.4	78
	15 % MgO				
2	Cu/Zn(10:90)/y-	WCO	18:1, 6, 65±5, 180	88.82	431
	Al ₂ O ₃ -800 °C				
3	Mg/MCM-41	WCO	8:1, 10, 80, 180	94	432
4	γ -Al ₂ O ₃ -CeO ₂	WCO	30:1, 7, 110, 270	81.1	433
5	KAcZX	WCO	48:1, 6, 120, 180	80.8	434
6	Sr/ZrO ₂	WCO	29:1, 2.7, 115.5, 169	79.7	435
7	Bi ₂ O ₃ -La ₂ O ₃	JCO	15:1, 2, 150, 240	94	436
8	CaO-La ₂ O ₃	JCO	25:1, 3, 160, 180	98.76	437
9	Mn@MgO-ZrO ₂	Kernel oil	15:1, 3, 90, 240	96.4	438
10	HPA@ZIF-8	Rapeseed oil	10:1, 4, 240, 300	98.02^{b}	439
11	AWS/SO42-	PFAD	15:1, 5, 80, 180	98	441
12	[Zn(4,4′-	Soybean oil	3.2/5 (v/v), 2, 180, 120	98	442
	bipy)(OAc) ₂] _n				
13	K/TiO ₂	Canola oil	36:1, 6, 70, 180	100^{b}	443

Table 21: Different bifunctional solid catalyst reported for biodiesel production.

^aMethanol-to-oil molar ratio, catalyst loading (wt. %), temperature (°C), reaction time (min).
 ^bConversion

1989

8. Biodiesel production process

Biodiesel can be produced by (trans)esterification, thermal cracking and pyrolysis.^{444–} ⁴⁴⁷ Amongst all these methods, transesterification is generally utilized for the synthesis of biodiesel.⁴⁴⁷ The generalized diagram for biodiesel production process is presented in Figure 36 which consists of synthesis and purification steps.⁴⁴⁸ Alkali, acid and enzyme are routinely exploited as a catalyst for the transesterification reactions. These catalysts had their own merits and demerits as compiled in Table 22.⁴⁴⁹ Till now, homogeneous base catalysts such as NaOH, KOH are normally utilized for biodiesel synthesis in industrial scale. In the meantime, owing 1998 to their capacity to catalyze both esterification/transesterification reactions, a homogeneous 1999 acid catalyst such as H_2SO_4 and HCl are generally picked for feedstock having high FFA such 2000 as non-edible vegetable oil, WCO and animal fats. Recently, the heterogeneous catalyst has 2001 attracted interest to a great extent for biodiesel synthesis because of their easy recyclability and 2002 reusability for successive reaction cycles.

2003



Figure 36: Representative diagram for biodiesel production. Reproduced from ref. [447].

2004

2005

Table 22: Points of interest and detriments of different catalyst utilized fortransesterification/esterification reaction (Reproduced from ref. [449]).

Catalyst types	Examples	Advantages	Disadvantages
Homogeneous			
Alkali	NaOH,	• High reactivity	• Inappropriate for high
	КОН	• Faster reaction rate	FFA in feedstocks
		• Minimum cost	• Deactivates in presence
		• Encouraging kinetics	of moisture and FFA.
		• Moderate working	• Requirement of high
		conditions	amount of waste water

					side reaction.
				•	Non-recyclable
				•	Corrosive in nature
Acid	H ₂ SO ₄ , HCl,	•	Non-reactive to	•	Slow reaction rate
	HF etc.		moisture and FFA	•	Long reaction time
			content in oil.	•	Equipment corrosion
		•	Catalyzed simultaneous	•	Higher reaction
			esterification/transester		temperature and pressure
			ification reactions.	•	High alcohol/oil
		•	Avoid formation of		requirement
			soap.	•	Weak catalytic activity
				•	Catalyst is difficult to
					recycle
Heterogeneous					
Alkali	CaO, SrO,	•	Non-corrosive	•	Slow reaction rate
	MgO,	•	Environmentally		compared to
	mixed oxide		benign		homogeneous one
	and	•	Recyclable	•	Low FFA requirement in
	hydrotalcite	•	Fewer disposal		the feedstock (< 1 wt. %)
			problems	•	Highly sensitive to water
		•	Easy separation		and FFA
		•	Higher selectivity	•	Saponification as side
		•	Longer catalyst life		reaction
				•	Soap formation
				•	High volume of
					wastewater
				•	Leaching of active
					catalyst sites
				•	Diffusion limitations.

• Complex and expensive synthesis route

Saponication occurs as a

•

					synthesis
Acid	ZrO, TiO,	•	Insensitive to FFA and	•	Moderate reaction rate
	ZnO, ion-		water content in the oil	•	Long reaction time
	exchange	•	Catalyzed simultaneous	•	Higher reaction
	resin,		esterification and		temperature and pressure
	sulfonic		transesterification	•	High alcohol/oil
	modified		reactions		requirement
	mesostructu	•	Recyclable, eco-	•	Weak catalytic activity
	red silica		friendly	•	Low acidic site
	etc.	•	Non-corrosive to	•	Low micro porosity
			reactor and reactor	•	Leaching of active
			parts		catalyst sites
				•	Diffusion limitations
				•	Complex and expensive
					synthesis route
				•	High cost of catalyst
					synthesis

High cost of catalyst

•

2006

2007 9. Catalyst comparison

It is seen from the literature that the reactivity of both homogeneous base and acid 2008 catalysts are very high compared to heterogeneous catalysts.^{61,70} Despite the high reactivity, 2009 homogeneous catalysts have some serious shortfalls such as low quality of glycerol produced, 2010 the catalyst cannot be regenerated and the lengthy process involves in the purification of 2011 2012 biodiesel; thereby makes the whole process labour-intensive and uneconomical.⁷⁶ To overcome 2013 these shortfalls, solid catalysts have been widely investigated. Alkaline earth, basic metal oxides and supported solid base catalyst shows excellent activity towards biodiesel production. 2014 however, their low stability, high sensitivity against the FFA limits its industrial application,¹⁴³ 2015 2016 whereas, their acid counterparts are not efficient towards the transesterification reactions. Recently, mixed metal oxides are gaining immense attentions in the field of biodiesel 2017 production due to their generally high surface area, excellent thermal and chemical stability, 2018 tailored acid-base properties and hence can be used prominently utilized for 2019 (trans)esterification of vegetable oil having high FFA.¹⁴⁵ 2020

2021 Literatures revealed that enzyme-based catalyst have various advantages over other catalyst such as environmentally benign, operate at mild reaction conditions and display high 2022 specificity.⁴³² Unfortunately, due to their sensitivity towards heat, poor operational stability 2023 2024 and narrow pH range, the use of such catalyst for industrial scale production of biodiesel is not 2025 a wise choice.⁴³³ However, immobilized lipase has various advantages compared to free lipase such as cost-effective, high thermal stability and greater tolerance to pH change.⁴³⁵ Thus, it has 2026 a scope for utilization in industrial scale biodiesel production. Besides, the present study 2027 suggests that bio-waste derived catalyst can be used potentially in the industrial scale 2028 production of biodiesel as they are easily available, cost-effective and most importantly 2029 environmentally benign.¹⁶¹ The main limitation is their reusability due to the leaching of the 2030 active sites.¹⁶⁵ Apart from that, metal free carbon based solid acid catalyst is also a promising 2031 candidate for industrial scale production of biodiesel as these materials possess promising 2032 features such as biogenic, environment-friendly, lower production costs, distinctive surface 2033 chemistry, high chemical and thermal stability.³⁸³ Bifunctional catalyst has been a recent 2034 interest in the realm of biodiesel research as it possess dual characteristics such as solid acidic 2035 character to tackle the FFA and solid basic character for easy transesterification of triglycerides 2036 to FAME and hence can be utilized for the (trans)esterification of diverse oil systems. Apart 2037 2038 from that, bifunctional catalyst are highly reusable, thermostable and insensitive to the moisture.⁴³⁸ Thus, bifunctional solid catalyst can be utilized in the successful production of 2039 2040 industrial scale biodiesel.

2041

2042 **10. Conclusion and outlook**

The exponential growth in the human population around the globe and industrial globalization tremendously increases the demand for petroleum fuels like diesel for various purposes. However, considering the limited resources of fossil fuels, searching for a novel, renewable and sustainable alternative fuel was required. In this context, researchers focused on FAME production from different renewable sources as an effective way. A variety of methods have been proposed for biodiesel production however, among all the existing methods, transesterification is considered as the foremost choice.

Transesterification reaction involves the use of a basic catalysts such as homogeneous and heterogeneous catalysts. The use of homogeneous catalysts is found to be promising as far as rate of biodiesel production is concerned; but it is associated with certain limitations. The homogeneous catalysts based transesterification reaction involves the consumption of high energy, moreover, the treatment of wastewater generated is essential due to the presence of 2055 unreacted chemicals. These limitations created the need for the development of efficient catalysts which was completed in terms of heterogeneous catalysts. These catalysts attracted a 2056 2057 great of attention from the scientific community all over the world because of its several 2058 advantages over homogeneous catalysts such as the simple realization of continuous reactors, production of cleaner glycerol, and the absence of both the alkaline catalyst neutralization step 2059 2060 and the necessity to replace the consumed catalyst. Due to these advantages, heterogeneous 2061 catalysts have opened up the chance for another powerful pathway for FAME production. 2062 However, the reactivity of the solid catalyst is dependent on several variables which mainly 2063 involve the oil type, alcohol to oil molar ratio, temperature, type of reactor, etc. therefore, selection of these variables at an optimum level is a crucial step. The heterogeneous catalysts 2064 are considered comparatively promising because only external-surface active species of porous 2065 solid support involved in the reaction and these catalysts can be recovered in some cases. 2066 However, in case of certain catalysts like CaO, leaching was reported which adversely 2067 2068 influences the reaction and hence, researchers are looking at nanotechnology as new hope.

Nanotechnology is the most emerging branch of science having promising applications in 2069 2070 catalysis. Moreover, it reported to have the ability to fabricate the catalyst surface in order to meet the prerequisites of explicit applications and beat the different issues related to both 2071 2072 homogeneous and heterogeneous catalysts. Nanocatalysts can act as an interface between homogeneous and heterogeneous catalysts having the possibility to develop promising solid-2073 2074 acid or solid-base catalysts which can be easily recovered using conventional filtration and centrifugation techniques. The development and use of magnetic nanoparticle-supported 2075 2076 catalysts is a path-breaking research because such catalysts can be easily recovered by using a 2077 simple magnetic field and reused for progressive reaction cycles which helps to reduce the 2078 overall process cost involved in biodiesel production which is the ultimate aim.

2079 It is well proven that the application of biological catalyst (enzyme) is more effective over 2080 all kinds of chemical catalysts, but the involvement of expensive enzyme increases the overall cost of the FAME production process. In this context, immobilization of such enzymes on the 2081 surface of various magnetic nanoparticles was found to be a novel concept because of the easy 2082 recovery of the immobilized enzyme along with magnetic nanoparticles and its reusability. 2083 2084 Moreover, it also solves the problem of leaching of the enzymes during the reaction due to immobilization. Although nanocatalysts reported to have promising applications, the 2085 toxicological concerns associated with nanoparticles is a topic of debate because there are 2086 2087 mixed opinions from the scientific community.

The present study revealed that the properties of the catalyst such as basicity and acidity play a pivotal role in the biodiesel production. Several literatures suggest the basicity of the catalyst directly proportional to the transesterification activity.^{171, 195} Similarly, acidity of the catalyst decides the esterification activity of the catalyst.^{383, 390} Higher the acidity of the catalyst, higher is the esterification activity. Apart from basicity and acidity, catalytic activity of the solid catalyst depends on its surface area and porosity. Literatures revealed that high surface area of the catalyst enhances the rate of biodiesel production.^{184, 225}

It is believed that several newly introduced catalysts will take a central position in the near 2095 2096 future and help to produce biodiesel through eco-friendly and economically viable processes. The development of novel heterogeneous catalysis having both acid and basic sites on its 2097 surface will have promising future in biodiesel production technologies because they have 2098 ability to overcome the issues usually caused because of the utilization of homogeneous 2099 catalysts. The application of bifunctional solids can be a novel way in heterogeneous catalysts 2100 mediated biodiesel production, because they showed the capability to accomplish simultaneous 2101 esterification and transesterification reactions in one-pot. In addition, the development and 2102 2103 application of nanocatalysts will be a milestone in biodiesel production. These nanocatalysts will be the next generation catalysts which will help to develop most effective, sensitive, 2104 2105 sustainable and economically viable technology for the FAME production in the near future. Although recent advances in the developments various homogeneous, heterogeneous and 2106 2107 nanocatalyst showed promising future for biodiesel industries or biorefineries, more efforts are required to develop even more effective and cheap catalysts which will help to overcome the 2108 2109 present issues all the above-mentioned catalysts and increase the efficiency of biodiesel 2110 production sustainably.

2111

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