



Biodiesel: The Use of Vegetable Oils and Their Derivatives as Alternative Diesel Fuels

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Vegetable oils and their derivatives (especially methyl esters), commonly referred to as “biodiesel,” are prominent candidates as alternative diesel fuels. They have advanced from being purely experimental fuels to initial stages of commercialization. They are technically competitive with or offer technical advantages compared to conventional diesel fuel. Besides being a renewable and domestic resource, biodiesel reduces most emissions while engine performance and fuel economy are nearly identical compared to conventional fuels. Several problems, however, remain, which include economics, combustion, some emissions, lube oil contamination, and low-temperature properties. An overview on all aspects of biodiesel is presented.

The use of vegetable oils in diesel engines is nearly as old as the diesel engine itself. The inventor of the diesel engine, Rudolf Diesel, reportedly used groundnut (peanut) oil as a fuel for demonstration purposes in 1900 (*1*). Some other work was carried out on the use of vegetable oils in diesel engines in the 1930's and 1940's. The fuel and energy crises of the late 1970's and early 1980's as well as accompanying concerns about the depletion of the world's non-renewable resources provided the incentives to seek alternatives to conventional, petroleum-based fuels. In this context, vegetable oils as fuel for diesel engines were remembered. They now occupy a prominent position in the development of alternative fuels. Hundreds of scientific articles and various other reports from around the world dealing with vegetable oil-based alternative diesel fuels ("biodiesel") have appeared in print. They have advanced from being purely experimental fuels to initial stages of commercialization. Nevertheless, various technical and economic aspects require further improvement of these fuels.

Numerous different vegetable oils have been tested as biodiesel. Often the

vegetable oils investigated for their suitability as biodiesel are those which occur

abundantly in the country of testing. Therefore, soybean oil is of primary interest as biodiesel source in the United States while many European countries are concerned with rapeseed oil, and countries with tropical climate prefer to utilize coconut oil or palm oil. Other vegetable oils, including sunflower, safflower, etc., have also been investigated. Furthermore, other sources of biodiesel studied include animal fats and used or waste cooking oils. Sources of biodiesel with some emphasis on developing countries have been discussed (2).

Several problems, however, have impaired the widespread use of biodiesel. They are related to the economics and properties of biodiesel. For example, neat vegetable oils reported to cause engine deposits. Attempting to solve these problems by using methyl esters causes operational problems at low temperatures. Furthermore, problems related to combustion and emissions remain to be solved. The problems associated with the use of biodiesel are thus very complex and no satisfactory solution has yet been achieved despite the efforts of many researchers around the world. This article will briefly discuss economics and regulatory issues as well as conventional diesel fuel (petrodiesel) and then focus on research on the use of biodiesel in a diesel engine.

Economics and Regulatory Issues

Economic reasons have been one of the major obstacles in the use of biodiesel. Diesel fuel (DF) derived from vegetable oils is more expensive than petroleum-based DF. The feedstock for biodiesel is already more expensive than conventional DF. For example, in the United States, a gallon of soybean oil costs approximately two to three times as much as a gallon of conventional DF. However, in the case of conversion of vegetable oils or fats to their esters, the resulting glycerol co-product, which has a potential market of its own, may offset some of the costs.

In most European countries, however, transportation fuels are so heavily taxed that tax incentives can be applied to encourage the use of biodiesel in the form of lower or no taxes on the biofuel and higher taxes on the petroleum-based fuel (3,4). This subsidy artificially cheapens the biodiesel to make it competitive. In many developing countries, the overriding concern is to become independent of the imported commodity petroleum. In the United States, the tax mechanism is inapplicable because of the comparatively low taxes on transportation fuels. Artificially regulating the demand for fuels from specific sources by means of taxation is currently politically not feasible.

Nevertheless, biodiesel is attractive for other reasons. Besides being a renewable resource and therefore creating independence from the imported commodity petroleum and not depleting natural resources, health and environmental concerns are the driving forces overriding the economic aspects in some cases. These concerns are manifested in various regulatory mandates of pollutants, particularly CAAA (Clean Air Act Amendments of 1990) and EPACT (Energy Policy Act of 1992) in the United States, which present opportunities for alternative fuels such as biodiesel. A life-cycle analysis of biodiesel (5) has shown that it is competitive with other alternative fuels such as compressed natural gas (CNG) and methanol in the urban transit bus market.

It is generally recognized that biodiesel has lower emissions, with the exception of nitrogen oxides (NO_x), than conventional petroleum-based DF. For example, due to its lack of sulfur, biodiesel does not cause SO_2 emissions. The lower emissions have caused biodiesel to be used in urban bus fleets and to make it especially suitable for

other niche markets such as mining and marine engines. Besides environmental and health reasons with accompanying Government regulations, focusing on the use of biodiesel in niche markets is rendered additionally attractive because not enough vegetable oil is produced to supply the whole diesel market with biodiesel.

Numerous reports exist showing that fuel economies of certain biodiesel blends and conventional DF are virtually identical. In numerous on-the-road tests, primarily with urban bus fleets, vehicles running on blends of biodiesel with conventional DF (usually 80% conventional DF and 20% biodiesel; for a list of most biodiesel demonstration programs in the United States, see Ref. 6) required only about 2-5% more of the blended fuel than of the conventional fuel. No significant engine problems were reported as discussed later.

Conventional Diesel Fuel. Diesel Engines.

In contrast to gasoline which is spark-ignited, DF after injection is ignited by the heat of compression in a diesel engine. The diesel engine is therefore also termed a compression-ignition (CI) engine. The differences in the ignition processes entail significant differences in chemical composition and physical properties of the fuels.

Conventional DF is, like gasoline, obtained from cracking of petroleum. It is a fraction boiling at an initial distillation temperature of 160° (90% range of 290-360°C) (7), also termed middle distillates because of its boiling range in the mid-range of cracking products.

The ignition quality of DF is commonly measured by ASTM D613 and reported as the cetane number (CN). Ignition quality is defined by the ignition delay time of the fuel in the engine. The shorter the ignition delay time, the higher the CN. To rank different compounds on the cetane scale, hexadecane ($C_{16}H_{34}$; also called cetane), which has a very short ignition delay, has been assigned a CN of 100. At the other end of the scale, 2,2,4,4,6,8,8-heptamethylnonane (HMN; also $C_{16}H_{34}$), which has poor ignition qualities, has been assigned a CN of 15. It should be noted that the cetane scale is arbitrary and that compounds with $CN > 100$ (although the cetane scale does not provide for compounds with $CN > 100$) or $CN < 15$ have been identified. The ASTM specification for conventional DF (ASTM D975) requires a minimum CN of 40.

The CN scale clarifies an important aspect of the composition of, or, on a more fundamental level, the molecular structure of the compounds comprising DF. Long-chain, unbranched, saturated hydrocarbons (alkanes) have high CNs and good ignition quality while branched hydrocarbons (and other materials such as aromatics) have low CNs and poor ignition quality.

Since both too high and too low CN can cause operational problems (in case of too high CN, combustion can occur before the fuel and air are properly mixed, resulting in incomplete combustion and smoke; in case of too low CN, engine roughness, misfiring, higher air temperatures, slower engine warm-up and also incomplete combustion occur), most engine manufacturers designate a range of required CN for their engines. In most cases, this range is around CN 40-50.

Conventional DF is classified into different grades by ASTM D 975. This classification is the following: No. 1 diesel fuel (DF1) comprises volatile fuels oils from kerosene to intermediate distillates. They are applicable for high-speed engines whose operation involves frequent and relatively wide variations in engine load and

stating that “the issue of including biodiesel mixtures or blends comprised of more than 20 percent biodiesel is currently under study. However, this subject is complex and will require significantly more data and information, and a separate, future rulemaking, before DOE can make a determination as to whether to include them in the definition of “alternative fuel.”

Vegetable oils.

Most vegetable oils are triglycerides (TGs; triglyceride = TG). Chemically, TGs are the triacylglyceryl esters of various fatty acids with glycerol (Figure 1).

Some physical properties of the most common fatty acids occurring in vegetable oils and animal fats as well as their methyl esters are listed in Table I. Besides these fatty acids, numerous other fatty acids occur in vegetable oils and animal fats, but their abundance usually is considerably lower. Table II lists the fatty acid composition of some vegetable oils and animal fats that have been studied as sources of biodiesel.

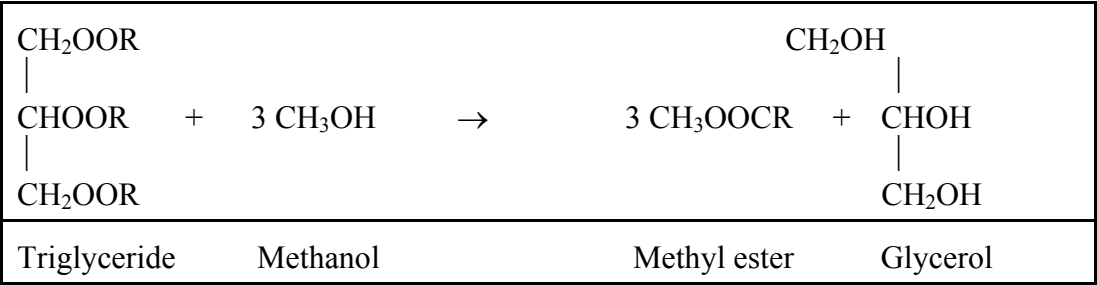


Figure 1. Structure of triglycerides and principle of the transesterification reaction (shown for methyl esters; R = (CH₂)_xCH₃ or unsaturated rests according to the fatty acids listed in Table I).

The most common derivatives of TGs (or fatty acids) for fuels are methyl esters. These are formed by transesterification of the TG with methanol in presence of usually a basic catalyst to give the methyl ester and glycerol (see Figure 1). Other alcohols have been used to generate esters, for example, the ethyl, propyl, and butyl esters.

Selected physical properties of vegetable oils and fats as they relate to their use as DF are listed in Table III. For esters these properties are given in Table IV. Also listed in Table III are the ranges of iodine values (centigrams iodine absorbed per gram of sample) of these oils and fats. The higher the iodine value, the more unsaturation is present in the fat or oil.

That vegetable oils and their derivatives are suited as DF is shown by their CNs (Table III) which generally are in the range suitable for or close to that of DF. The heat

Table I. Selected properties of some common fatty acids and esters.

<i>Trivial (Systematic)name^a; Acronym^b</i>	<i>Mol. wt.</i>	<i>m.p.^c (°C)</i>	<i>b.p.^{c,d} (°C)</i>	<i>Cetane No.</i>	<i>Heat of Combustion^e (kg-cal/mole)</i>
Caprylic acid	144.22	16.5	239.3		

<i>Trivial (Systematic)name^a; Acronym^b</i>	<i>Mol. wt.</i>	<i>m.p.^c (°C)</i>	<i>b.p.^{c,d} (°C)</i>	<i>Cetane No.</i>	<i>Heat of Combustion^e (kg-cal/mole)</i>
(Octanoic acid); 8:0					
Capric acid (Decanoic acid); 10:0	172.27	31.5	270	47.6 (98.0) ^f	1453.07 (25°),
Lauric acid (Dodecanoic acid); 12:0	200.32	44	131 ¹		1763.25 (25°),
Myristic acid (Tetradecanoic acid); 14:0	228.38	58	250.5 ¹⁰⁰		2073.91 (25°),
Palmitic acid (Hexadecanoic acid); 16:0	256.43	63	350		2384.76 (25°),
Stearic acid (Octadecanoic acid); 18:0	284.48	71	360d		2696.12 (25°),
Oleic acid (9Z-Octadecenoic acid); 18:1	282.47	16	286 ¹⁰⁰		2657.4 (25°),
Linoleic acid (9Z,12Z- Octadecadienoic acid); 18:2	280.45	-5	229-30 ¹⁶		
Linolenic acid (9Z,12Z,15Z- Octadecatrienoic acid); 18:3	278.44	-11	230-2 ¹⁷		
Erucic acid (13Z-Docosenoic acid); 22:1	338.58	33-4	265 ¹⁵		
Methyl caprylate (Methyl octanoate); 8:0	158.24	...	193	33.6 (98.6) ^f	1313
Methyl caprate (Methyl decanoate); 10:0	186.30	...	224	47.7 (98.0) ^f	1625
Methyl laurate (Methyl dodecanoate); 12:0	214.35	5	266 ⁷⁶⁶	61.4 (99.1) ^f	1940
Methyl myristate (Methyl tetradecanoate); 14:0	242.41	18.5	295 ⁷⁵¹	66.2 (96.5) ^f	2254
Methyl palmitate (methyl hexadecanoate); 16:0	270.46	30.5	415-8 ⁷⁴⁷	74.5 (93.6) ^f	2550
Methyl stearate (Methyl octadecanoate); 18:0	298.51	39.1	442-3 ⁷⁴⁷	86.9 (92.1) ^f	2859
Methyl oleate (Methyl 9Z- octadecenoate); 18:1	296.49	-20	218.5 ²⁰	47.2 ^g	2828
Methyl linoleate (Methyl 9Z, 12Z-octadecadienoate); 18:2	294.48	-35	215 ²⁰	28.5 ^g	2794
Methyl linolenate (Methyl 9Z, 12Z,15Z-octadecatrienoate); 18:3	292.46	-57 -52	109 ^{0.018}	20.6 ^g	2750

	<i>F a t t y A c i d C o m p o s i t i o n (Wt.-%)</i>							
Babassu	44-45	15-17	5.8-9	2.5-5.5	12-16	1.4-3		
Canola			4-5	1-2	55-63	20-31	9-10	1-2
Coconut	44-51	13-18.5	7.5-10.5	1-3	5-8.2	1.0-2.6		
Corn			7-13	2.5-3	30.5-43	39-52	1	
Cottonseed		0.8-1.5	22-24	2.6-5	19	50-52.5		
Linseed			6	3.2-4	13-37	5-23	26-60	
Olive		1.3	7-18.3	1.4-3.3	55.5-84.5	4-19		
Palm		0.6-2.4	32-46.3	4-6.3	37-53	6-12		
Peanut		0.5	6-12.5	2.5-6	37-61	13-41		1
Rapeseed		1.5	1-4.7	1-3.5	13-38	9.5-22	1-10	40-64
Safflower			6.4-7.0	2.4-29	9.7-13.8	75.3-80.5		
Safflower, high-oleic			4-8	2.3-8	73.6-79	11-19		
Sesame			7.2-9.2	5.8-7.7	35-46	35-48		
Soybean			2.3-11	2.4-6	22-30.8	49-53	2-10.5	
Sunflower			3.5-6.5	1.3-5.6	14-43	44-68.7		
Tallow (beef)		3-6	25-37	14-29	26-50	1-2.5		

a) These oils and fats may contain small amounts of other fatty acids not listed here. For example, peanut oil contains 1.2% 20:0, 2.5 22:0, and 1.3% 24:0 fatty acids (*181*).

oleate was 47.2, the lowest of these 18:1 methyl esters. The double bond of methyl petroselinate is closer to one end of the molecule. It also has the longest uninterrupted alkyl chain of these compounds, which may play a role because alkanes have higher CNs as discussed above. This complements the observations in Ref. 16. Another possibility is benzene formation by a disproportionation reaction from cyclohexane, which in turn would arise from cleavage of methyl oleate (*17*). The low CN of benzene would account for the lower CN of methyl oleate. The other 18:1 compounds would not form cyclohexane due to the different positions of the double bond.

Table III. Fuel-related properties and iodine values of various fats and oils.^a

<i>Oil or Fat</i>	<i>Iodine Value</i>	<i>CN</i>	<i>HG (kJ/kg)</i>	<i>Viscosity (mm²/s)</i>	<i>CP (°C)</i>	<i>PP (°C)</i>	<i>FP (°C)</i>
Babassu	10-18	38					
Castor	82-88	?	39500	297 (38°)	---	-31.7	260
Coconut	6-12						
Corn	103-140	37.6	39500	34.9 (38°)	-1.1	-40.0	277
Cottonseed	90-119	41.8	39468	33.5 (38°)	1.7	-15.0	234
Crambe	93	44.6	40482	53.6 (38°)	10.0	-12.2	274
Linseed	168-204	34.6	39307	27.2 (38°)	1.7	-15.0	241
Olive	75-94						
Palm	35-61	42					
Peanut	80-106	41.8	39782	39.6 (38°)	12.8	-6.7	271
Rapeseed	94-120	37.6	39709	37.0 (38°)	-3.9	-31.7	246
Safflower	126-152	41.3	39519	31.3 (38°)	18.3	-6.7	260
High-oleic safflower	90-100	49.1	39516	41.2 (38°)	-12.2	-20.6	293
Sesame	104-120	40.2	39349	35.5 (38°)	-3.9	-9.4	260
Soybean	117-143	37.9	39623	32.6 (38°)	-3.9	-12.2	254
Sunflower	110-143	37.1	39575	37.1 (38°)	7.2	-15.0	274
Tallow	35-48	-	40054	51.15 (40°)	-	-	201
No. 2 DF		47	45343	2.7 (38°)	-15.0	-33.0	52

a) Iodine values combined from Refs. 176 and 181. Fuel properties from Ref. 11. All tallow values from Ref. 177 (No CN given in Ref. 177, calcd. cetane index 40.15).

The combustion of the glyceryl moiety of the TGs could lead to formation of acrolein and this in turn to the formation of aromatics (16), although no acrolein was found in precombustion of TGs (18). This may be one reason why fatty esters of vegetable oils perform better in a diesel engine than the oils containing the TGs (16). On the other hand, as discussed above, benzene may arise from the oleic moiety also.

Table IV. Fuel-related physical properties of esters of oils and fats.^a

<i>Ester</i>	<i>CN</i>	<i>HG</i> (kJ/kg)	<i>Viscosity</i> (mm ² /s)	<i>CP</i> (°C)	<i>PP</i> (°C)	<i>FP^b</i> (°C)
<i>Methyl</i>						
Cottonseed ^c	51.2	-	6.8 (21°)	-	-4	110
Rapeseed ^d	54.4	40449	6.7 (40°)	-2	-9	84
Safflower ^e	49.8	40060	-	-	-6	180
Soybean ^f	46.2	39800	4.08 (40°)	2	-1	171
Sunflower ^g	46.6	39800	4.22 (40°)	0	-4	-
Tallow ^h	-	39949	4.11 (40°)	12	9	96
<i>Ethyl</i>						
Palm ⁱ	56.2	39070	4.5 (37.8°)	8	6	19
Soybean ^f	48.2	40000	4.41 (40°)	1	-4	174
Tallow ^j				15	12	
<i>Propyl</i>						
Tallow ^j				17	12	
<i>Isopropyl</i>						
Soybean	52.6 ^k			-9 ^l	-12 ^l	
Tallow ^j				8	0	
<i>n-Butyl</i>						
Soybean ^f	51.7	40700	5.24 (40°)	-3	-7	185
Tallow ^j				13	9	
<i>2-Butyl</i>						
Soybean ^l				-12	-15	
Tallow ^j				9	0	

a) CN = cetane number; CP = cloud point, PP = pour point, FP = flash point. b) Some flash points are very low. These may be typographical errors in the references or the materials may have contained residual alcohols. c) Ref. 42. d) Ref. 55. e) Ref. 178. f) Ref. 17. g) Ref. 179. h) Ref. 177. i) Ref. 180. j) Ref. 95. k) Ref. 127. l) Ref. 123.

<i>Fuel Property</i>	<i>Unit</i>	<i>Test Method</i>	<i>Limit (min.)</i>	<i>Limit (max.)</i>
Density at 15°C	g / ml	ISO 3675	0.875	0.900
Kinematic Viscosity at 15°C	mm ² / s	ISO 3104	3.5	5.0
Flash Point (Pensky-Martens)	°C	ISO 2719	100	
CFPP April 15- September 30 October 1- November 15 November 16 - February 28 March 1 - April 14	°C	DIN EN 116		0 -10 -20 -10
Sulfur Content	wt.-%	ISO 4260		0.01
Carbon Residue - Conradson (10% distillation residue)	wt.-%	ISO 10370		0.30
Cetane Number		ISO 5165	49	
Ash	wt.-%	ISO 6245		0.01
Water	mg / kg	ASTM D 1744		300
Total Contamination	mg / kg	DIN 51419		20
Copper Strip Corrosion (3 h at 50°C)		ISO 2160		1
Acid Number	mg KOH / g	DIN 51558 Part 1		0.5
Methanol	wt.-%	tbs ^{b)}		0.3
Monoglycerides	wt.-%	tbs		0.8
Diglycerides	wt.-%	tbs		0.1
Triglycerides	wt.-%	tbs		0.1
Free Glycerine	wt.-%	tbs		0.02
Total Glycerine	wt.-%	tbs		0.23
Iodine Value	g Iodine / 100g	DIN 53241 Part 1		115
Phosphorus	mg / kg	tbs		10

a) CFPP = Cold-filter plugging point. b) tbs = to be standardized.

The iodine value (IV; see Table III) has been included in the European standards and is based on rapeseed oil as biodiesel feedstock. It is set at IV = 115, which would exclude soybean oil (neat vegetable oils and their methyl esters have nearly identical IVs) as biodiesel feedstock. The discussion in the previous section, however, shows that **Table VI. Suggested ASTM standard for pure (100%) biodiesel.^a**

