Biofuel by isomerizing metathesis of rapeseed oil esters with (bio)ethylene for use in contemporary diesel engines

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Rapeseed oil methyl ester (RME) and (bio)ethylene are converted into biofuel with an evenly rising boiling point curve, which fulfills the strict boiling specifications prescribed by the fuel standard EN 590 for modern (petro)diesel engines. Catalyzed by a Pd/Ru system, RME undergoes isomerizing metathesis in a stream of ethylene gas, leading to a defined olefin, monoester, and diester blend. This innovative refining concept requires negligible energy input (60°C) and no solvents and does not produce waste. It demonstrates that the pressing challenge of increasing the fraction of renewable fuels in engine fuel may be addressed purely chemically rather than by motor engineering.

Introduction

The increasing environmental awareness and upcoming shortage of fossil oil have triggered a growing interest in sustainable mobility concepts. A liquid biofuel with high energy density that requires no modification to the millions of existing car, truck, ship, and aircraft engines and that can be distributed within current infrastructure would ideally complement electrical solutions for the mobility of the future (1).

Around 80% [107 × 10^6 metric tons per annum (Mt/a)] of the worldwide plant oil production is required to cover food demands; the remaining 20% are converted into biofuels (11 × 10^6 Mt/a) or put to industrial use (15 × 10^6 Mt/a) (2, 3). Biodiesel is presently generated by methanolysis of vegetable oils or animal fats, which transforms the triacylglycerols to fatty acid methyl esters (FAMEs) andglycerol (4). In the European Union, a FAME content of 7% in diesel fuel is imposed by European Norm (EN) 590 (5), and 10% will be enforced by 2020 (6). Besides being a renewable and biodegradable material, biodiesel has beneficial properties over conventional diesel, such as its inherent lubricity, lower sulfur content, and higher flash point. Certain disadvantages, including its instability toward oxidation (7), high viscosity and pour point, and increased nitrogen oxide release (8), can be tackled with the help of additives (9).

The main obstacle to increasing the biodiesel content in motor fuels arises from its unfavorable boiling behavior (10). The standard EN 590 for commercial diesel fuel suitable for powering standard diesel engines calls for a smoothly rising boiling point curve within strict limits (Fig. 1) that ensure optimal fuel ignition and combustion. Petrodiesel, which fulfills these requirements, consists of a mixture of linear and branched hydrocarbons with various chain lengths. In contrast, a typical biodiesel based on rapeseed oil methyl ester (RME) mainly contains linear molecules with 19 carbon atoms, that is, 65% methyl oleate [18:1], 22% methyl linoleate [18:2], 8% methyl linolenate [18:3], 1% methyl stearate [18:0], and 4% methyl palmitate [16:0] (11). As a result, its boiling range starts at too high a temperature and is disadvantageously narrow (330° to 400°C), clearly outside the EN 590 specifications (Fig. 1). This adversely affects its ignition behavior and precludes late-stage injections as are required by modern diesel engines with particulate filters (12–14). In pure form, RME and related biodiesels can, thus, only be used in dedicated engines especially engineered to cope with these challenging physical and chemical properties (15).

The only known strategy to convert vegetable oils into biofuel suitable for use in standard diesel engines is their conversion into mixtures of saturated hydrocarbons by energy-intensive hydroprocessing (16, 17). Products obtained by conventional cross-metathesis of RME do not meet the specifications of EN 590 (17). An alternative, low-temperature refining concept that allows converting vegetable oils into biodegradable product blends with petrodiesel-like boiling ranges would be of tremendous interest. This key challenge is addressed by the isomerizing metathesis process disclosed herein, which enables generating an EN 590–compatible fuel (Fig. 1) from fatty acid esters and ethylene, which is abundantly available from bioethanol or shale gas (18).

In isomerizing metatheses, an isomerization catalyst constantly moves double bonds up and down along alkyl chains within a molecule, while an olefin metathesis catalyst continually shuffles the alkyl residues attached to the double bonds between two molecules (19–22). This iterative, cooperative action of two orthogonal catalysts allows converting single olefins into olefin blends with carbon-chain lengths evenly distributed around the chain length of the starting material. As a technology, isomerizing olefin metathesis has recently made a substantial leap forward from the previous pioneering works in the 1990s (19, 23–27). However, it has never been envisaged as a tool for diesel refining. The abundance of potential catalyst poisons in technical-grade plant oils (for example, nitrogen compounds), the incompatibility of multiply unsaturated olefins and of ethylene with isomerization catalysts, the requirement of expensive solvents in most metatheses, and the difficulty to reproducibly tune product distributions are only some of the challenges that seemed to preclude the use of isomerizing metathesis in biodiesel refining. Consequently, no solvent-free isomerizing cross-metathesis of FAMEs and short-chain olefins has been reported to date. When using ethylene and pure methyl oleate in our previously developed isomerizing metathesis process, we were unable to obtain full conversion or homogeneous product distributions even in the presence of solvent (19). The corresponding process with unpurified RME or other natural FAMEs suitable for fuel production was, so far, entirely out of reach.

Results

Years of process design led to an effective catalyst system for a simplified model reaction: the isomerizing cross-metathesis of equimolar amounts...
of 1-hexene and unpurified RME from grocery-grade rapeseed oil. In the presence of 0.05 mole percent (mol %) [Pd(μ-Br)(‘Bu₃P)]₂ (IC-1) and 0.05 mol % of an N-heterocyclic carbene (NHC)–indenylidene ruthenium complex (Ru-1), an even distribution of olefins, monoesters, and diesters formed at 50°C without added solvent (Fig. 2).

Alternative olefin metathesis catalysts were also evaluated, including second-generation indenylidene-ruthenium complexes Ru-2 and Ru-3 and Hoveyda-type catalysts Ru-4, Ru-5, and Ru-6 (fig. S1). The results obtained are visualized in the superimposed olefin fraction histograms in Fig. 3. The non-isomerized metathesis products decene, tetradecene, and octadecene are overrepresented for all catalyst systems except for the IC-1/Ru-1 combination (black bars with superimposed black trend line). This indicates that all other metathesis catalysts adversely affect the isomerization activity of IC-1.

Incorporation of 1-hexene brought about an overall shortening of the chain lengths. As a result, the mixture displayed an evenly rising boiling point curve that closely matches that of petrodiesel in terms of initial and mean boiling points. However, the 1-hexene/RME product showed a recovery of only 93% at 360°C, which narrowly misses the EN 590 specification of 95% (Fig. 1). Toward the end of the distillation, it partially decomposed with smoke formation. This is a common problem for biodiesel caused by oxidation of sensitive polyunsaturated fatty acids.

**Fig. 1.** Boiling point curves of commercial diesel and biodiesel (RME) before and after isomerizing metathesis. The hashed areas represent the limits specified in EN 590. *, increasing decomposition; Recovery, percentage of the sample recovered in the receiving flask during distillation.

**Fig. 2.** Boiling point histogram of the product blend after isomerizing hexenolysis of RME. Conditions: 10.0 mmol of 1-hexene/RME (1:1), 0.05 mol % IC-1 and Ru-1, neat, 50°C, 20 hours. The mixture was hydrogenated for analysis. This histogram was overlaid with simulated curves for 1000 molecules per catalyst (± 0.05 mol % catalyst) of both 1-hexene and RME, with 30,000 metathesis and 7500 double-bond migration steps. ---, olefins; ---, monoesters; ---, diesters.
acid derivatives and is usually addressed by partial hydrogenation of the product fractions (28, 29). This was the best boiling behavior achieved for an isomerizing hexenolysis of RME.

An in-depth understanding of the influence of substrate and process parameters on the boiling behavior was derived from model reactions in combination with arithmetic considerations at equilibrium, and simulations of the reaction course.

At equilibrium, the product distribution for the reaction of $X$ equivalents of 1-hexene and $Y$ equivalents of methyl oleate is governed purely by the relative abundance of C–C double bonds and functionalized and unfunctionalized chain termini. Thus, the ratio of olefins, monoesters, and diesters in the product mixture corresponds to $(2 \times X + Y)^2 : (2 \times X + Y) \times Y \times 2:Y^2$ (Fig. 2). For the reaction of pure methyl oleate ($Y = 1$) with one equivalent of 1-hexene ($X = 1$), the ratio of olefins, monoesters, and diesters following isomerizing hexenolysis should be 9:6:1. The boiling behavior of the product mixture is also influenced by the carbon-chain lengths of the olefin, monoesters, and diester fractions. The mean number of aliphatic carbon atoms per double bond was calculated using the formula $\bar{b} = (Y \times 15 + X \times a) / (X + Y)$, with $a$, $b$, $X$, and $Y$ as defined in Fig. 2. Two alkene carbons as well as the number of ester termini were added to obtain the mean chain lengths (MCLs). For the isomerizing hexenolysis with one equivalent of 1-hexene, the MCLs of the olefin, monoester, and diester fractions at equilibrium are 11.5, 12.5, and 13.5, respectively. Experimentally, mean values of $<12.9$ carbons for the olefins [uncorrected for volatiles not detectable by gas chromatography (GC) analysis], 14.4 for the monoesters, and 17.5 for

![Graph showing the carbon-chain length histograms for olefin blends obtained by isomerizing hexenolysis with different Ru catalysts.](image)

Fig. 3. Carbon-chain length histograms for olefin blends obtained by isomerizing hexenolysis with different Ru catalysts. Conditions: 10.0 mmol of 1-hexene/RME (1:1), 0.05 mol % IC-1 and Ru-cat., neat, 50°C, 20 hours. The samples were hydrogenated for GC analysis to simplify the chromatograms.

![Graphs showing simulated boiling point distribution of olefins (-----), monoesters (—), and diesters (–) and boiling point curve of the products after isomerizing ethenolysis.](image)

(A) Effect of the ethylene/RME ratio near equilibrium and (B) effect of the ratio of isomerization/metathesis steps per molecule in preequilibrium reactions. Recovery, percentage of the sample recovered in the receiving flask during distillation analysis.
the diesters were detected for RME. This indicates that the reaction did not reach equilibrium.

To rationalize the product distributions in preequilibrium mixtures, we developed a simulation program in which a given number of randomly chosen molecules from a mixture of 2000 molecules of methyl oleate/hexene at a given ratio (1:1 in the above case) undergo a single shift of their double bond. Then, another given number of randomly chosen molecules undergo metathesis. These two steps are iterated a given number of times so that different overall and relative turnover numbers for both catalyst systems can be modeled (see the “Simulation of the isomerizing hexenolysis reaction mixtures” in the Supplementary Materials).

The experimental product distribution for the isomerizing hexenolysis of RME (Fig. 2, histograms) fits best with the simulated outcome for an average of 15 metathesis and 3.75 isomerization steps per olefin molecule (Fig. 2, simulated curves). This translates to catalyst turnover numbers of 30,000 for Ru-1 and 7500 for IC-1, which are impressive values for solvent-free reactions. Slight deviations of the calculated product distributions from those experimentally observed were to be expected: The multiply unsaturated fatty acids in RME release short-chain fragments and the saturated components, namely, methyl palmitate (1%) and stearate (4%), cannot undergo isomerizing metathesis, which explains the protruding signals of C16 and C18 esters and the increased integral and MCL of the monoester fraction.

With all the necessary tools in hand, we investigated whether and how it is possible to modulate the boiling point curve of RME by isomerizing cross-metathesis with ethylene so that it would meet the EN 590 specifications. Figure 4A illustrates the predicted effect of the ethylene/RME ratio on the simulated boiling point distribution and net boiling point curve near equilibrium, at 150 isomerization and metathesis steps each per molecule. None of the boiling point curves meet the specifications at both middle and high % recovery. The median boiling temperature is too high at a 1:2 ethylene/RME ratio and too low at a 2:1 ratio. The curves for ratios around 1:1 are closest to the desired profile. The boiling behavior can be adjusted further by changing the number of isomerization and metathesis steps per molecule (Fig. 4B). When setting the metathesis steps to 15 per molecule and gradually increasing the number of isomerizations, the product distributions change from irregular patterns with several chain-length maxima per fraction (<10 isomerization steps) to increasingly broad, smooth product distributions around single maxima. At the same time, the amount of high-boiling long-chain compounds initially decreases, goes through a minimum of 10 isomerization steps per molecule, and then increases again. For an ethylene/RME ratio between 0.7:1 and 1.3:1, the boiling point curve is predicted to meet the specifications with a boiling temperature above the lower threshold at 65% recovery and below the upper threshold at 95% recovery if the number of isomerization steps is between 10 and 30.

Table 1. Optimization of the isomerizing ethenolysis of RME. Conditions for entries 3 to 12: 2.50 mmol of RME (based on methyl oleate), ethylene, Ru-cat., Ru-CAAC, IC-1, neat, 16 hours, 60°C.

<table>
<thead>
<tr>
<th>Entry</th>
<th>Ru-cat. (mol %)</th>
<th>Ru-CAAC (mol %)</th>
<th>IC-1 (mol %)</th>
<th>Ethylene (ml)</th>
<th>Reaction temperature (°C)</th>
<th>Average MCL</th>
<th>Evenness of the curve</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Isomerizing hexenolysis (see above)</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>50</td>
<td>15.0</td>
<td>Excellent</td>
</tr>
<tr>
<td>2</td>
<td>Sequential isomerizing ethenolysis</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>50</td>
<td>13.7</td>
<td>Good</td>
</tr>
<tr>
<td>3</td>
<td>Ru-1 (0.05)</td>
<td>—</td>
<td>0.05</td>
<td>300 (at 6 bar)</td>
<td>50</td>
<td>—</td>
<td>No conversion</td>
</tr>
<tr>
<td>4</td>
<td>Ru-1 (0.10)</td>
<td>—</td>
<td>0.10</td>
<td>20</td>
<td>60</td>
<td>18.0</td>
<td>Poor</td>
</tr>
<tr>
<td>5</td>
<td>Ru-5 (0.10)</td>
<td>—</td>
<td>0.10</td>
<td>20</td>
<td>60</td>
<td>15.3</td>
<td>Fair</td>
</tr>
<tr>
<td>6</td>
<td>Ru-7 (0.10)</td>
<td>—</td>
<td>0.10</td>
<td>20</td>
<td>60</td>
<td>15.2</td>
<td>Good</td>
</tr>
<tr>
<td>7</td>
<td>Ru-11 (0.10)</td>
<td>—</td>
<td>0.10</td>
<td>20</td>
<td>60</td>
<td>15.3</td>
<td>Fair</td>
</tr>
<tr>
<td>8</td>
<td>Ru-5 (0.10)</td>
<td>—</td>
<td>0.40</td>
<td>20</td>
<td>60</td>
<td>15.8</td>
<td>Fair</td>
</tr>
<tr>
<td>9</td>
<td>Ru-7 (0.10)</td>
<td>—</td>
<td>0.40</td>
<td>20</td>
<td>60</td>
<td>15.5</td>
<td>Excellent</td>
</tr>
<tr>
<td>10</td>
<td>Ru-11 (0.10)</td>
<td>—</td>
<td>0.40</td>
<td>20</td>
<td>60</td>
<td>15.4</td>
<td>Excellent</td>
</tr>
<tr>
<td>11</td>
<td>Ru-11 (0.10)</td>
<td>—</td>
<td>0.40</td>
<td>20</td>
<td>60</td>
<td>15.3</td>
<td>Excellent</td>
</tr>
<tr>
<td>12</td>
<td>Ru-11 (0.10)</td>
<td>—</td>
<td>0.40</td>
<td>300 (at 6 bar)</td>
<td>60</td>
<td>14.4</td>
<td>Excellent</td>
</tr>
<tr>
<td>13</td>
<td>Ru-11 (0.10)</td>
<td>0.10</td>
<td>0.40</td>
<td>Constant stream</td>
<td>60</td>
<td>12.9</td>
<td>Excellent</td>
</tr>
</tbody>
</table>
per molecule and if the number of metathesis steps is greater than 7 per molecule. The predicted MCLs for this mixture are 8.6, 11.4, and 14.4 for olefins, monoesters, and diesters, respectively, or 11.5 overall.

Equipped with a link between the physical properties and the chemical composition of the product mixture, as well as an activity profile for the bimetallic catalyst system, we sought suitable catalysts and reaction conditions (see the “Development of the isomerizing ethenolysis of RME” in the Supplementary Materials). The best-known isomerization catalysts, among them the Chaudret, Cole-Hamilton, Behr, Grotjahn/Schrock, and Skrydstrup systems (20, 26, 30–33), were incompatible with ruthenium metathesis catalysts or deactivated by ethylene. IC-1 was the sole catalyst to display sufficient activity but only at ethylene pressures of ≤1 bar.

IC-1 was evaluated in combination with different metathesis catalysts for the isomerizing ethenolysis of RME (Table 1; see table S4 for full details). The average MCL of the three product fractions was used as a measure of ethylene intake and of the resulting shift toward lower boiling points (cf. figs. S12 to S15 for the effect with 1-hexene). In addition, we evaluated the shape of the product distributions obtained to ensure evenly rising boiling point curves.

Without ethylene, the average MCL is 18 (isomerizing self-metathesis of RME). The isomerizing hexenolysis, used as a further reference point, gave an average MCL of 15 (entry 1). To ensure a defined ratio of RME and ethylene, a two-step process consisting of non-isomerizing ethenolysis of RME followed by isomerizing metathesis of the ethenolyzed product was assessed as a third reference point. This resulted in an average MCL of 13.7 (entry 2). The deviation from the targeted value of 11.5 results from the incorporation of too little ethylene (0.83 rather than one equivalent), a low isomerization rate, difficulties detecting volatile short-chain olefins, and the presence of saturated fatty esters. The experimental boiling point curve measured for this reference material was marginally outside of the EN 590 specifications (see table S5), which confirms that the material has to meet all predicted parameters to reach the targeted behavior.

The system composed of Ru-1 and IC-1 was inactive under ethylene pressure, possibly because of decomposition of the ruthenium complex (entry 3). The catalyst loading was increased to 0.10 mol % to compensate for lower activity and stability in the presence of ethylene. Conversion was first observed after switching to atmospheric ethylene pressure (entry 4). Three Ru-NHC complexes (Ru-5, Ru-7, and Ru-11; figs. S1 and S25) mediated the isomerizing metathesis of RME in the presence of ethylene and resulted in almost even product distributions with single maxima (cf. table S4, entries 5 to 7). At an IC-1 loading further increased to 0.40 mol %, the distributions were more homogeneous, and protruding cross-metathesis products were no longer observed (entries 8 to 10). However, these catalysts incorporated too little ethylene into the product mixture, as visible from the medium chain lengths. We therefore needed to find a highly active ethenolysis catalyst that would retain its activity in the presence of IC-1. Fortunately, the complex Ru-CAAC described by Marx et al. (34) was found to be compatible with the isomerizing ethenolysis. However, Ru-CAAC did not promote the cross-metathesis of long-chain olefins, leading to the overproportional formation of volatile olefins.

The solution was to combine IC-1, Ru-CAAC, and Ru-11 into a mutually compatible, ternary system that is able to catalyze the isomerization, ethenolysis, and long-chain olefin cross-metathesis, respectively. At 60°C, the MCLs reached 14.3, 14.9, and 16.8, respectively (entry 11). Increasing ethylene pressure did not allow the average MCL (entry 12) to sufficiently reduce. However, by running the reaction under a constant stream of ethylene at atmospheric pressure in a specially constructed reactor, the MCLs were further reduced to <12.3, 12.2, and 14.0 carbons for olefins, monoesters, and diesters, respectively (entry 13). These values are close to the MCLs of 8.6, 11.4, and 14.4 predicted by simulation of the optimum product mixture. The gas chromatogram of the experimentally obtained mixture is comparable to that simulated for a 1.3:1 ratio of ethylene/RME, with an average of 15 metathesis and 12 isomerization steps per substrate molecule. This translates to turnover numbers of 15,000 for the metathesis and 3000 for the isomerization catalyst, which is well within the targeted range. Notably, the purity of the ethylene source does not have a significant effect on the catalyst system. When using industrial-grade ethylene (N3.5) in the process, the distributions and MCLs were identical.

The physical properties were experimentally quantified using a product mixture obtained on a larger scale. Thus, 135 ml of RME containing IC-1 (1.24 g), Ru-CAAC (243 mg), and Ru-11 (330 mg) was stirred under a stream of ethylene at 60°C for 16 hours (Fig. 5). Distillation analysis in an EN ISO (International Organization for Standardization) 3405 apparatus furnished an experimental boiling point curve that corresponded well with the prediction derived from the simulation (Figs. 1 and 4). The product blend passed the ASTM (American Society of Testing and Materials) D 6751 boiling specification for biodiesel (36), because 90% distilled below 360°C. It also fulfilled all three specifications for petrodiesel fuels laid out in EN 590, namely, a recovery of (i) <65% at 250°C, (ii) >85% at 350°C, and (iii) 95% at ≤360°C. As predicted, the boiling point curve of the ethylene/RME product had a terminal boiling temperature and an overall shape similar to petrodiesel.
In contrast to the 1-hexene/RME product, no decomposition was discernible during distillation of the ethylene/RME product, likely due to the lower content of polyunsaturated fatty acid esters following ethenolysis. Further analysis using standard methods for fuel testing revealed that the material had a sulfur content of <5 mg/kg, a viscosity of 2.12 mm²/s, and a lubricity of 232 μm, all values well within the EN 590 specifications. The acid value of 0.360 mg of KOH per gram was below the threshold for pure biodiesel. The cloud and pour points, which are not explicitly specified for temperate climatic zones, are significantly below 0°C and, thus, in a good range for unmodified fuel. To demonstrate the new biofuel’s suitability as motor fuel, we used it to propel a model diesel car (see movie S1).

DISCUSSION

In conclusion, isomerizing olefin metathesis with (bio)ethylene allows converting RME into olefin mixtures that match the boiling behavior of diesel fuel as specified in EN 590. This technology may turn out to be a decisive breakthrough toward increasing the content of renewables in diesel fuel. The activity, compatibility, and stability of the catalysts are decisive breakthrough toward increasing the content of renewables in diesel fuel as specified in EN 590. This technology may turn out to be a suitable concept but do not fulfill the industrial manufacturing standards. Many steps are still required to establish an industrial production of an EN 590–compatible biofuel. At the end of this development, we envision a fully continuous flow process in which an RME/ethylene feed passes through alternating sections of immobilized isomerization and metathesis catalysts dimensioned to provide the ideal ratio of turnover frequencies. The limiting challenge at this stage is the discovery of long-lived, economically viable, heterogeneous or heterogenizable catalysts.

MATERIALS AND METHODS

Procedure for the single-stage isomerizing ethenolysis

In a glovebox under nitrogen atmosphere, a 1-liter Parr autoclave was charged with [1-[(R)-2-(1-methylethyl)phenyl]-2-pyrrrolidinylidene]dichloro[2-[(1-methylethoxy-κO)phenyl]methylene-κC]-ruthenium(II) (CAS no. 959712-80-2; Ru-CAAC, 243 mg, 0.40 mmol), bromo(trimethylsilyl)dimethylsilane (DMDS) (1.0 g, 1.60 mmol), UniCure M73 SIPr (Ru-11, 330 mg, 0.40 mmol), and RME (135 ml, 400 mmol based on methyl oleate). The autoclave was closed and removed from the glovebox. The resulting reaction mixture was stirred under a stream of ethylene (99.995% purity) at atmospheric pressure for 16 hours at 60°C. The reactor was cooled to ambient temperature, and a 30% solution of H2O2 (40.9 ml, 400 mmol) was slowly added at 0°C under vigorous stirring. The organic phase was separated, dried over MgSO4, yielding 75 ml of a brown oil (55% based on volume). After high-temperature vacuum distillation (1 × 10-3 mbar, >350°C), the product mixture was obtained as a light yellow liquid (73 ml, >98 weight % recovery after distillation).

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