Development of Heterogeneous Basic Catalysts Supported on Silica for the Synthesis of High Quality Bio-Diesel from Waste Cooking Oil

Paweesuda NATEWONG, Yayoi MURAKAMI, Haruki TANI and Kenji ASAMI

(Received August 5, 2015)

Effect of addition of CaO and ZrO₂ to MgO/SiO₂ catalyst, which is effective for synthesis of High Quality Bio-Diesel (HiBD) was investigated using atmospheric agitated reactor at 430 °C and LHSV = 0.3 h⁻¹. Binary MgO-CaO/SiO₂ and MgO-ZrO₂/SiO₂ catalysts and ternary MgO-CaO-ZrO₂/SiO₂ catalyst were prepared by the incipient wetness impregnation method. Physical properties of these catalysts were characterized by XRD and BET methods. Both the fatty acids and the triglycerides in waste cooking oil were converted into hydrocarbon gases, CO, CO₂, water and hydrocarbon oil. CaO- and/or ZrO₂-added catalysts gave higher CO₂ yields than MgO/SiO₂ alone, and showed lower acid values. These observation indicates that the added oxides promote the decarboxylation reactions. Iodine values also decreased to some extent by the addition of these oxides.

Key Words
High quality bio-diesel, MgO/SiO₂ catalyst, CaO and ZrO₂ addition

1. Introduction
The production of bio-fuels as clean and renewable fuels has recently been paid a keen attention to because of a rapid price rise of petroleum and an increase in the greenhouse gas emission. Fatty acid methyl ester (FAME) and Bio Hydrofined Diesel (BHD) are known as current bio-fuels from triglycerides of vegetable oils and animal fats for diesel engine 1) 2). However, sub-raw materials other than oil are required in both FAME and BHD production processes; methanol and high pressure hydrogen gas are necessary for FAME and BHD, respectively.

We have developed a new biodiesel production process which gives mixed hydrocarbon liquids of diesel fraction with high selectivity without using any sub-

Faculty of Environmental Engineering, The University of Kitakyushu
1-1, Hibikino, Wakamatsu-ku, Kitakyushu-shi, Fukuoka 808-0135, Japan

meters 3) ~ 6), and named this fuel as High Quality Bio-Diesel (HiBD) 7). HiBD is expected as the next generation biodiesel because its properties are suitable for the latest common rail type engine. The production process of HiBD is quite simple, where triglycerides are converted to liquid hydrocarbons and light hydrocarbon gases through decarboxylation over solid catalysts as shown in Fig. 1. This process can be adapted to a lot of feed materials such as vegetable oil, waste cooking oil, and animal fats.

One of the most significant factors in the HiBD production is the catalyst which promotes the conversion of triglycerides. A magnesium oxide supported on silica (MgO/SiO₂) catalyst is found to be effective for this reaction, and its basic property promotes the decarboxylation 3) ~ 6). Since both CaO and ZrO₂ are known to have basic character and also ability to enlarge active surface area 6), and thus they are expected to enhance the activity. In the present study, therefore, addition of CaO and ZrO₂ to MgO/SiO₂ catalyst
was investigated.

2. Experimental

2.1 Preparation of catalyst

Silica-supported oxide catalysts were prepared by the incipient wetness impregnation method using aqueous solutions of Mg, Ca, and Zr nitrates. The silica support used was Fuji Silisia Q10 (particle size, 1.18-2.36 mm; SA 300 m²/g; PV, 0.95 cm³/g). After the impregnation, they were dried in an oven at 100 °C for 12 h, and then calcined in air at 500 °C for 3 h. Components of the catalysts are abbreviated by using loadings (wt%) and the initial letters of the oxides; for example, 5M5C5Z indicates that the catalyst contains 5 wt% of MgO, 5 wt% of CaO, and 5 wt% of ZrO₂, respectively. Physical properties of these catalysts were characterized by X-ray diffraction and N₂ adsorption.

2.2 Catalytic test

Catalytic reactions were conducted in an agitated reactor system at 430 °C under atmospheric pressure. Fig. 2 shows the experimental setup for the HiBD production. 25 g of the granular catalyst was charged into the reactor, and it was heated up to the reaction temperature in an He flow (50 mL/min). Waste cooking oil with an acid value of 39.123 mg-KOH/g-oil and iodine value of 101 g-I/100 g-oil, which was supplied from the university restaurant, was introduced into the catalyst bed at 0.25 mL/min with a pump. The gaseous products which came out from the reactor were cooled and condensed at 0 °C, the uncondensed gaseous products were sampled and analyzed every 30 minutes during the reaction by a gas chromatograph with thermal conductivity detector (GC-TCD) and one with flame ionization detector (GC-FID) on line. Yields of the cracked oil and water were calculated on the basis of weight of the reactant oil fed. Carbon number distribution of the product oil was analyzed off line with another GC-FID. The amount of residue was determined by weight difference between catalysts before and after use. Total acid value, index of free acid contents, and iodine value, index of the amount of unsaturated bonding, of the oil were measured by potentiometric titration methods according to JIS 2501-2003 and JIS K0070-1992, respectively.

3. Results and Discussion

Table 1 shows the specific surface area (SA) and the pore volume (PV) of the fresh catalysts. When CaO and/or ZrO₂ was added to the MgO/SiO₂ catalyst, both SA and PV of the catalysts increased. This suggests that CaO and ZrO₂ have similar effect on surface area and pore volume. The highest SA and PV were obtained with 5M5C5Z/SiO₂ catalyst.

![Fig. 3 XRD patterns of A) 10M/SiO₂, B) 5M5C/SiO₂, C) 5M5Z/SiO₂ and D) 5M5C5Z/SiO₂ catalysts](image)
supported oxides of MgO, CaO, and ZrO₂ were detectable. Thus, these species would exist as highly dispersed ones on the support 9) 10).

**Fig. 4** shows product yield from waste cooking oil over the oxide catalysts by decarboxy-cracking. The major products recovered were cracked oil, dry gas (C₃-C₄ hydrocarbons), CO, CO₂, water and residue. The unreacted fatty acids was not detected, as the reacted fatty acids and the unreacted fatty acids may be condensed into liquid products. Therefore, liquid products not only contain the cracked oils but also comprise the unreacted fatty acids. Unreacted fatty acids would also remain as residue. CO₂ yields with CaO and ZrO₂ added catalysts were 1.1 ~ 1.4 times higher than that with MgO/SiO₂ catalyst. These facts show that the added oxides would assist MgO and promote the decarboxylation from intermediate free acids as well as triglycerides 11) 12). The increase in the specific surface area would be another effect as shown in Table 1. Formation of CO, even the yield was low, was observed over each catalyst. It may be formed by the decomposition of ketones as by-products 11) 12) or through the reverse water-gas shift reaction.

**Fig. 5** shows the carbon number distribution of the product oil. GC-MS analysis was done by area normalization method, which revealed that more than 99 wt% of the oil were hydrocarbons, and very small amount of free fatty acids and alcohols with different carbon numbers were contained. As a whole, the distribution profiles are similar to each other; a large amount of hydrocarbons with the diesel fraction (C₁₀-C₂₀) are the major products, while the compounds with carbon number higher than 21 were not observed. The main product obtained with the catalysts except for 5M5C/SiO₂ was C₁₇ hydrocarbons, which would be originated from oleic acid group (C₁₇H₃₃COO⁻) and linoleic acid group (C₁₇H₃₁COO⁻) in the waste cooking oil. The C₃ hydrocarbons could probably be produced from the triglycerides directly and after the decomposition to the corresponding free fatty acids, while the other hydrocarbons with shorter chain than C₁₇ would be formed by the cracking of the alkyl groups in the reactants and the products. The lower yield of C₁₇ with 5M5C/SiO₂ catalyst suggests the higher cracking ability than the other catalysts. C₁₈ compounds were found to be ketones as described above such as methyl ketone and ethyl ketone. Yield of these compounds increased by adding oxides, especially ZrO₂ 13) 14).

**Fig. 6** shows the acid value (AV) and iodine value (IV) of the cracked oils obtained over the four oxide catalysts. The acid values obtained with the binary MgO-CaO and MgO-ZrO₂ catalysts and the ternary MgO-CaO-ZrO₂ catalyst were about 12.0 mg-KOH/g-oil, and lower than that with MgO alone (14 mg-KOH/g-oil). This would mean the promotion of decarboxylation from the free fatty acid which

---

**Fig. 4** Product yield from waste cooking oil over the oxide catalysts

**Fig. 5** Carbon number distribution of the product oil

**Fig. 6** The acid value and iodine value of the cracked oils
are the intermediates from triglycerides to hydrocarbons. Although these values were slightly higher than the level of neat regulation (0.5 mg-KOH/g-oil) for the biodiesel fuel, we have already developed an adsorption technique to clear the regulation for the oils with such acid values. Iodine values of all these oils were about 90 g-I/100 g-oil, which were lower than the neat regulation (120 g-I/100 g-oil). The IV's obtained with the binary and the ternary oxide catalysts were slightly lower than that with MgO alone. The added CaO and ZrO$_2$ might form new active sites and promote protonation and cracking of C=C double bonds.

4. Conclusion

Effects of addition of CaO and ZrO$_2$ to MgO/SiO$_2$ catalyst on the HiBD production from waste cooking oil have been investigated and the following conclusions were obtained. The binary MgO-CaO/SiO$_2$ and MgO-ZrO$_2$/SiO$_2$ catalysts and the ternary MgO-CaO-ZrO$_2$/SiO$_2$ catalyst were simply synthesized by incipient wetness impregnation method and could be directly used for the HiBD production. The catalyst exhibited good catalytic activities and promoted decarboxylation reaction, which gave higher CO$_2$ yields than MgO/SiO$_2$, and showed lower acid values mainly due to their high surface area and pore volume.

Acknowledgment

This work was supported by JST-JICA SATREPS program and NEDO, Japan. Natewong P. is grateful to the support from the Japanese Government (Monbukagakusho) Scholarship

References

1) Saka, S., All About Biodiesel, IPC Inc, (2006)
2) Maeda, S., Sci. Tech. Trends, 11-27 (2007)
3) Tani, H.; Shimouchi, M.; Haga, H.; Fujimoto, K., J. Jpn. Inst. Energy, 90, 466-470 (2011)
4) Tani, H.; Hasegawa, M.; Asami, K.; Fujimoto, K., Catal. Today, 164, 410-414 (2011)
5) Asami, K.; Komatsu, Y.; Ono, K.; Murakami, Y.; Tani, H.; Fujimoto, K., J. Jpn. Petrol. Inst., 58, 293-301 (2015)
6) Murakami, Y.; Tani, H.; Asami, K.; Fujimoto, K., J. Jpn. Inst. Energy, 94, 1074-1078 (2015)
7) Jpn. Reg. Trade Mark, 5468299
8) Jiao, X.; Li, N.; Xiao, F.; Wei, W., Energy Fuels, 27, 5407-5415 (2013)
9) Garcia, J.; Lopez, T.; Alvarez, M.; Aguilar, D. H.; Quintana, P., J. Non-Cryst. Solids, 354, 729-732 (2008)
10) Xu, H.; Chu, W.; Luo, J.; Zhang, T., Chem. Eng. J., 170, 419-423 (2011)
11) Chang, C. C.; Wang, S. W., Ind. Eng. Chem, 39, 1543-1548 (1947)
12) Watanabe, M.; Inomata, H.; Smith Jr, R. L.; Arai, K., Appl. Catal A, 219, 146-156 (2001)
13) Pestman, R.; Koster, R. M.; van Duijne, A.; Pieterse, J. A. Z.; Ponec, V., J. Catal, 168, (1997)
14) Watanabe, M.; Tida, T.; Inomata, H., Energy. Convers. Manage, 47, 3344-3350 (2006)
15) Fujimoto, K.; Murakami, Y.; Tani, H.; Asami, K., Jpn. Pat. Appl, 19, 2014-2075