HCA AND TWC HYBRID SYSTEM FOR REDUCING COLD-START EMISSION

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ABSTRACT-In line with the Super Ultra Low Emission Vehicle (SULEV) regulation, the main idea in this study has been focused on the utilization of hydrocarbon adsorber (HCA) to adsorb the excess hydrocarbons emitted during a period of engine cold-start. As main recipes of HCA materials, many types of zeolite as well as the combination of alumina and precious metals were used. Representative physico-chemical factors of zeolite such as acidic and hydrophobic properties were characterized. The optimum recipe of HCA materials was also determined. Among the acid properties of zeolites, the Si/Al ratio was found to be the most important factor to get higher hydrocarbon adsorption capacity.

KEY WORDS: HCA, Zeolite, SULEV, Hybrid system

1. INTRODUCTION

One of the very important problems in automobile emission control is how to resolve the problem of cold start (Beck et al., 1997; Hirofumi and Tagomori, 1993). Hydrocarbons from cold start emission emit more than 85% of total hydrocarbon emission. Emission regulations are met today by using three-way catalysts (TWC), which reduce exhaust emissions through catalytic reaction (Ichikawa et al., 1999). However, these systems are inactive during cold start and warm-up of an engine, as they require a temperature level typically around 300°C for sufficient conversion. In order to satisfy the stringent Super Ultra Low Emission Vehicle (SULEV) legislation, the increase of catalytic activity at cold start condition is quite necessary. The research interest has been focused on the adsorption properties at cold-start conditions. The concept of using a hydrocarbon adsorber (HCA) to reduce the amount of cold-start hydrocarbons emitted from vehicles is well known (Sim and Chung, 2002; Engeler et al., 1993; Yamamoto et al., 2000).

Although the HCA technique is quite effective, tighter emission regulations may not be sufficiently met by simply using configuration with HCA. Therefore much research effort has utilized the HCA system in combination with TWC in order to reduce NOx and hydrocarbon emission at cold-start condition. To develop

the optimized operating condition of HCA and TWC, many of the HCA recipes have been tried, and their physico-chemical properties have been characterized. The adsorption characteristics of prepared HCA materials have also been examined. Furthermore, the catalytic activities of HCA and TWC combined system have been measured in a simulated flue gas and catalytic reactor system.

2. EXPERIMENTAL PROCEDURE

2.1. Preparation of Catalysts

Zeolite, as a major component of HCA material, was prepared by ion exchanging steps proposed by Iwamoto and Yahiro. The final HCA material was prepared through conventional wet impregnation method (Hong *et al.*, 1998), and the precious metals (3wt) Pd and Rh were added in the ratio of 9:1 (Paul and Todd, 1999). The catalysts were calcined in air for four hours at 550°C after drying for 2 hours at 150°C. In order to obtain extra function, base metal oxides such as Ba, Zr, Ce, and La were added to the samples in a similar manner (Long *et al.*, 1997).

2.2. Catalytic Activity Test

To simulate the emission gases emitted from gasoline engine, each reactant gas was passed through a mixing tank with enough mixing time. The mixing gases were then passed through the reactor having the hydrocarbon

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adsorber and three-way catalysts in a dual bed form. To determine the change of emission concentration, the activity test was performed before and after the gases were passed through the reactor. The simulated emission gas contents were: NO 450 ppm, CO 4200 ppm, C₃H₆ 1800 ppm, C₃H₈ 450 ppm, H₂ 1400 ppm, CO₂ 3.0%, H₂O 10%, O₂ 1.6%, and N₂ as a balance gas. The estimated Air/Fuel ratio (λ value) was approximately 1.0.

2.3. Characterization

Various temperature programmed desorption measurements were carried out on a conventional TPD system equipped with a thermal conductivity detector (TCD) cell. The catalysts were exposed to He gas at 350°C for 2 h in order to remove water and impurities on the surface. After pretreatment, the samples were exposed to adsorbate for 1 h. Finally, the programmed heating at a rate of 10°C/min was started and the samples were then heated to 650°C. The amount of the desorbed gas was continuously monitored with a TCD cell.

3. RESULTS AND DISCUSSION

3.1. Effect of Si/Al Ratio

At first, the adsorption properties of ethylene on various kinds of ZSM-5 catalysts were examined depend-ing upon the Si/Al ratio (Figure 1). In automobile emissions, ethene is as important as propene among hydrocarbon emissions. Therefore, both gases were tested as probe hydrocarbon. Since both gases showed similar adsorption behavior and amounts depending upon Si/Al ratio, the data for both were cited inter-mixed.

Though the amounts of adsorption were similar in the case of 28 and 40 in Si/Al ratios, it was found that about 50% of adsorption amount increased when the Si/Al ratio

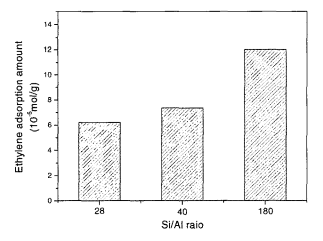


Figure 1. Ethylene adsorption amounts of HZSM-5 as a function of Si/Al ratio.

was 180. This could be interpreted that at higher range of Si/Al ratio, the hydrophobicity was jumped into higher value; therefore, the selective adsorption of non-polar material such as hydrocarbon was increased significantly. This result coincides with that of Engler *et al.*, which shows that at low Si/Al ratio of zeolite the polar materials like water, ammonia, and alcohols as well as non-polar materials such as hydrocarbon were well adsorbed regardless of polarity. But at high Si/Al ratio, by changing the surface property from hydrophilic to hydrophobic, the increased amount of hydrocarbon could be adsorbed selectively.

3.2. Effect of Hydrophobicity

In general, various kinds of experimental methods have been proposed for the quantitative determination of hydrophobicity. Among them, the method of hydrophobic index calculation proposed by Long was adopted for this study. In this method, the hydrophobic index could be expressed as the weight loss of X_{HC}/X_{H2O} in thermogravimetric analysis (TGA) curve. In this study, the hydrophobic index was defined as the weight loss from room temperature up to 150°C to the weight loss from 150°C up to 400°C in the thermogram curve (Figure 2) for each sample. The relationship between the hydrocarbon adsorption amounts and the measured values of hydrophobic index was examined. Surprisingly, as shown in Figure 3, a linear relationship between the hydrophobic index (h) and the amount of adsorption was observed for most zeolite samples. This result suggests that for the selection of

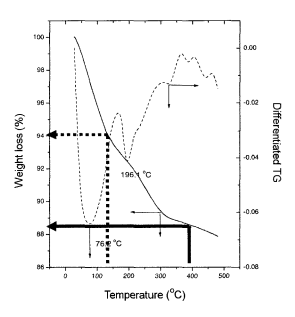


Figure 2. DTG and TG curve of H-Mordernite (Si/ Al=60, ←—: water weight loss, ←—: hydrocarbon weight loss).

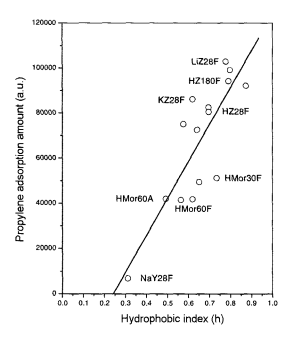


Figure 3. Correlation of hydrophobicity and adsorption amount of ion exchanged zeolites [F=fresh, A=aged, Mor=mordernite and Z=ZSM5].

zeolites to make an efficient HCA, the hydrophobic index is the most important factor to examine.

3.3. Effect of Surface Acidity

The ammonia temperature programmed desorption (TPD) method and amine titration method are known to be effective characterization techniques for the determination of total surface acidity and acid strength distribution on catalyst surface. As shown in Figure 4 and 5, the acid amount and acid strength distribution had been measured for ZSM-5 catalysts, depending upon the Si/Al ratio. As expected, upon increasing the value of Si/Al ratio, the total amount of acidity was reduced. From the ammonia TPD, it was found that strong acid sites at high temperature region were specially reduced as the Si/Al ratio increased. These phenomena could also be confirmed from the acid strength distribution diagram that large amount of acid sites stronger than pH 4 was reduced significantly with increasing the Si/Al ratio.

An important point is that, in spite of the reduction of

Table 1. Ethylene adsorption amount of ZSM-5.

Si/Al ratio	Ethylene adorption vol. (ml)
28	0.135
40	0.139
180	0.192

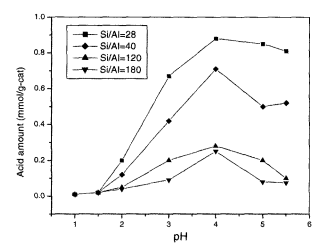


Figure 4. Acid amounts of ZSM-5 as a function of Si/Al ratio.

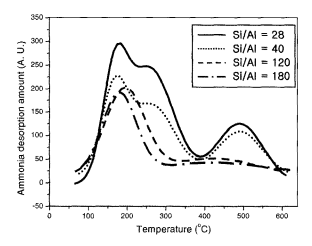


Figure 5. Ammonia TPD patterns for ZSM-5 with different Si/Al ratios.

total acidity at higher Si/Al ratio, the amount of hydrocarbon adsorption as described in Table 1 still shows a high value of adsorption. This means that between the two major factors governing the amount of hydrocarbon adsorption, which are the hydrophobic properties and acid amount, the effect of hydrophobicity is more dominant than acidity.

Depending upon alkali metals ion-exchanged in ZSM-5, minor changes in the amount of hydrocarbon adsorption could be monitored (Figure 6). This means that by changing some cations of zeolite to alkali-metals, a minor improvement in the adsorption amount could be obtained at a fixed Si/Al ratio.

To verify the source of acidity increase as well as adsorption amount for various ion-exchanged ZSM-5, the change of acid sites was examined through an ammonia

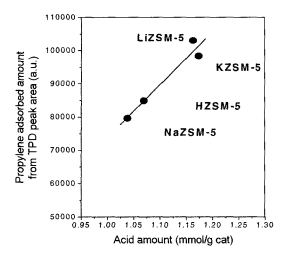


Figure 6. Correlation of acid amount and propylene adsorption amount on alkaline metal-exchanged ZSM-5 catalysts.

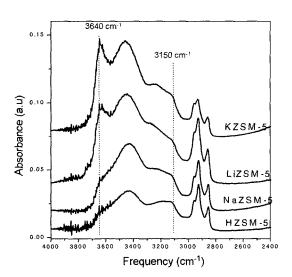


Figure 7. FT-IR result of alkaline metal-exchanged ZSM-5 catalysts.

adsorption experiment using in-situ IR. As a result, in the case of either Li⁺ or K⁺-exchanged ZSM-5, a new peak was observed at 364000 m⁻¹ (Figure 7), which can be interpreted as a Brönsted acid site. However, there was no peak growth on H⁺ and Na⁺ ion exchanged ZSM-5. This result suggests that for specially selected ion exchanged ZSM-5 catalysts, some Lewis acid sites could be transferred into Brönsted acid site so as to increase the amount of total acidity, which in turn results in the increase of hydrocarbon adsorption. This observation that the main site of hydrocarbon adsorption is Brönsted acid site rather than Lewis acid site through a detrium labeling experiment is supported by Eisuke *et al.* (2000).

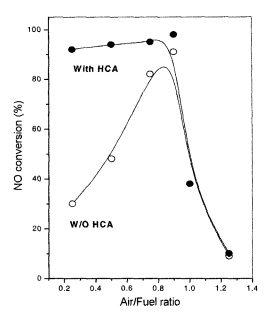


Figure 8. NO_x reduction over Pd-only catalyst with respect to Air/Fuel ratio on Pd-only catalyst.

Therefore, for a fixed Si/Al ratio, by changing the cations of zeolites to K⁺ and Li⁺ ions, more improved hydrocarbon adsorption amounts could be obtained. This principle could be applied to prepare HCA materials for the removal of excessibly emitted hydrocarbons during an engine cold-start period.

3.4. The Effects on Combination of HCA and TWC To observe the combination effect of HCA with TWC, the catalytic activity of TWC was measured in a simulated flue gas and reactor system. The combination effect was measured in the aspect of light-off temperature (LOT)₅₀, which is the temperature of the reaction exhibiting the 50% conversion of hydrocarbon, together with the final conversion of NO, CO, and hydrocarbon at reaction temperatures from 50°C to 400°C. Finally, in order to promote thermal durability as well as improved catalytic activity, some base metal oxides were added on the surface of HCA materials and the catalytic activity was examined.

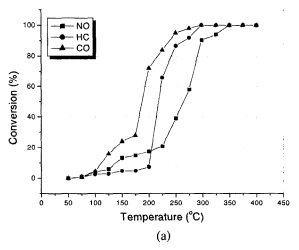
3.4.1. Catalytic activity of TWC only

Two types of TWC were used: Pd-only catalyst and Pt/Rh catalyst. From comparative studies, it was observed that the Pd-only catalyst is superior to the conventional Pt/Rh catalyst in the aspect of high temperature durability and durability in sulfur poisoning. However, the conventional Pt/Rh catalyst showed better catalytic activity in NO reduction specially at low temperature region. The major problem of Pd-only catalyst in the de-NO activity is that this catalyst shows very low activity at fuel rich

region mainly due to the hydrocarbon poisoning. Since the Air/Fuel ratio (λ =1 at stoichiometric ratio) was perturbed around stoichiometric ratio, it was important to monitor the removing efficiencies of NO, H/C, and CO from fuel rich to fuel lean region. Figure 8 shows the de-NO activity of Pd-only catalyst depending upon Air/Fuel ratio.

The de-NO activity at fuel rich region was very poor due to the poisoning of excess hydrocarbon. However, when the HCA was added, since the excess hydrocarbon was significantly adsorbed on HCA, the de-NOx activity at fuel rich region was much enhanced. This illustrates the advantage of HCA attachment on the TWC system.

The results of catalytic activity measurements are shown in Figure 9. Regardless of the type of catalyst, the behavior of CO conversion seemed quite similar for the whole temperature range. However, in the case of NO and HC conversion, two types of catalyst showed drastically different behaviors. In the LOT₅₀ of hydro-



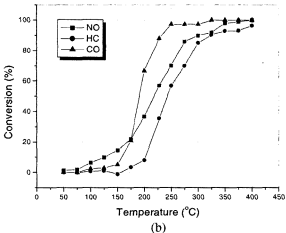


Figure 9. NO, THC and CO removal efficiency of commercial Pd-only(a) and Pt/Rh(b) catalysts.

carbon, while Pd-only catalyst was exhibited at 220°C, Pt/Rh catalyst showed at 243°C, which is higher by 23°C. Therefore, by considering the benefits of low LOT₅₀, Pd-only catalyst was selected for the combination study of TWC with HCA.

3.4.2. Combination effect of model HCAs and Pd-only TWC

Table 2 shows the various kinds of recipes for model HCAs which were prepared on monolith wash-coated by γ -Al₂O₃, precious metals, base metal oxide (BMO), and Li⁺ exchanged ZSM-5 (Si/Al=120, 180) based on the results of adsorption experiments.

Figure 10 and Figure 11 shows the activities of combined catalysts with Pd-only catalyst with prepared HCAs. In the case of HCA I, mixed with Pd-only catalyst, the LOT₅₀ of hydrocarbon was too high. HCA II was prepared by adding BMO. In this sample, La was specially added to prevent active site poisoning by excess hydrocarbon in rich condition together with promoted thermal stability. Ba addition was expected to increase NO adsorption amount, and Zr and Ce addition could promote thermal stability and oxygen storage capacity (OSC). When BMOs were added, the LOT50 of hydrocarbon was reduced by 10°C, and CO and NO were reduced by 20°C and 40°C, respectively, as compared with the results in Pd-only performance. In HCAII, zeolites, such as mordenite and Y-zeolite were added, and hydrocarbon LOT₅₀ and CO LOT₅₀ were both reduced by 10°C, and NO LOT₅₀ fell by 50°C in contrast to Pd-only catalyst. However, these samples were not good enough to reduce the LOT₅₀ of hydrocarbon significantly.

Kanazawa (Kanazawa *et al.*, 2001) reported a relationship between hydrocarbon adsorbtion amount and zeolite pore size with ZSM5, Mordenite, Y-zeolite, and 13X zeolite. ZSM5 was found to be inferior to the Y-zeolite in adsorb-ing C8 paraffin, C7 aromatics, and other higher hydro-carbons. The Y-zeolite, on the other hand, could not adsorb propylene. From these results, it was concluded that ZSM5 was effective for lower, C3 to C5 hydro-carbons, and Y-zeolite was effective for higher C6 and above. In case of adding Y-zeolite and Mordenite, the reducing ability of hydrocarbons could not be improved.

Table 2. Hydrocarbon adsorber catalysts compositions.

Model	Catalyst composition
HCA I	PM 3 wt% + γ -Al ₂ O ₃ + Me-ZSM-5(120)
HCA II	PM 3 wt% + γ -Al ₂ O ₃ + Me-ZSM-5(120)
	+ BMO
HCA III	PM 3 wt% + γ -Al ₂ O ₃ + Zeolite Y +
	MOR(120) + Me-ZSM-5(120)+ BMO
HCA IV	PM 3 wt% + γ -Al ₂ O ₃ + Me-ZSM-5(180)

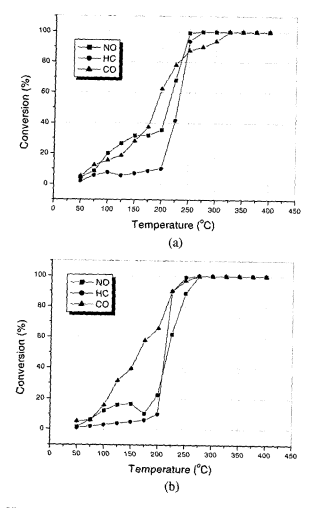
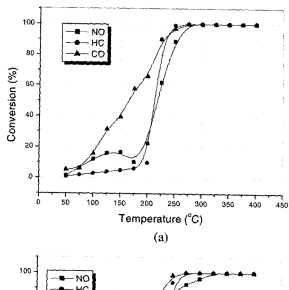


Figure 10. Combination effects of HCA I(a), HCA II(b) with Pd-only TWC.

In the case of HCA IV system, the combination effect of HCA prepared by ZSM-5 (Si/Al=180) was investigated. As shown in Figure 11, NO LOTso was less by 80°C compared to the Pd-only catalyst, CO LOT50 by 15°C, and hydrocarbon LOT50 by 20°C. It was surprising that the reduction of LOT_{50} in NO was higher than LOT_{50} in hydrocarbon. This effect probably comes from the prevention of hydrocarbon poisoning by uptaking the excess hydrocarbon through HCA. This could be interpreted that at higher range of Si/Al ratio, the hydrophobicity has been jumped into higher value, and therefore, the selective adsorption of non-polar material has been increased significantly. Therefore, hydrocarbon adsorption amount is increased, and hydrocarbon and NO is easily reduced. From these results, it could be seen that Si/Al ratio or hydrophobicity is a very important factor in selecting HCA. HCA materials using zeolite that have high Si/Al ratio and prepared by ion-exchanging method showed excellent reduction in LOT50 in NO by 80°C. Its



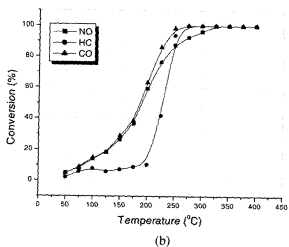


Figure 11. Combination effects of HCA III(a), HCA IV(b) with Pd-only TWC.

main reason could be explained by its acid characteristics and hydrophobicity of zeolite.

4. CONCLUSION

- (1) At higher range of Si/AI ratio, the hydrophobicity has been jumped into higher value. Therefore, the selective adsorption of non-polar material such as hydrocarbon has been increased significantly.
- (2) For specially selected ion exchanged ZSM-5 catalysts, some Lewis acid sites could be transferred into Brönsted acid site so as to increase the amount of total acidity, resulting in the increase of hydrocarbon adsorption.
- (3) If hydrocarbon adsorption catalyst is used with threeway catalyst, it could prevent Pd from poisoning by hydrocarbon. In addition, it could make better NO and CO conversion catalytic activity as well as lowering

- LOT₅₀ about 20-30°C. It is beneficial to the increase of catalytic activity at cold start condition.
- (4) BMO were added to promote temperature durability. This was found to be very helpful in promoting catalytic activity at low temperature region.
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