

# **Improved Processes to Remove Naphthenic Acids**

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## *Abstract*

In the first year of this project, we have established our experimental and theoretical methodologies for studies of the catalytic decarboxylation process. We have developed both glass and stainless steel micro batch type reactors for the fast screening of various catalysts with reaction substrates of model carboxylic acid compounds and crude oil samples. We also developed novel product analysis methods such as GC analyses for organic acids and gaseous products; and TAN measurements for crude oil. Our research revealed the effectiveness of several solid catalysts such as NA-Cat-1 and NA-Cat-2 for the catalytic decarboxylation of model compounds; and NA-Cat-5~NA-Cat-9 for the acid removal from crude oil. Our theoretical calculations propose a three-step concerted oxidative decarboxylation mechanism for the NA-Cat-1 catalyst.

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## ***Introduction***

Naphthenic Acid (NA), representing a group of organic acids in crude oils, causes serious corrosion problems in refinery units and transport pipelines. Due to its complex compositional heterogeneity, it is currently difficult to predict the severity of corrosion for an individual or a small group of NA compounds by any analytic measurements. A Total Acidity Number (TAN) or the neutralization number (Neut Number), defined by the number of milligrams of KOH required to neutralize the acidity in one gram of oil, and is therefore the commonly adopted criterion for predicting the corrosive potential of a crude oil. With this standard, high TAN oils (> 0.5 KOH/g) are less desirable than lower TAN oils and are labeled with lower price in market. Crude oils from California, Venezuela, North Sea, Western Africa, India, China and Russia have typically higher naphthenic acid contents.

Developing a naphthenic acid removal process and obtaining a fundamental understanding on the catalytic decarboxylation mechanism will significantly help the petroleum industry of the United State and worldwide in improving refining processing of heavy crude oils possessing high contents of the naphthenic acid. Being sponsored by DOE through this project (DE-FC26-02NT15383), we have assembled a team combining both experimental/theoretical researchers with extended academic/industrial expertise aiming at developing cost-effective methods to remove naphthenic acid from crude oils. Specifically we have two objectives. One objective is to develop a catalytic system to cleanly decarboxylate aliphatic and aromatic acids under low temperature conditions. In particular, we propose to develop a system that will be readily adaptable to removing naphthenic acids from crude oil. Our second objective is to develop an effective solid adsorbent that can remove naphthenic acid *via* solid liquid separation.

## ***Executive Summary***

Significant progress has been made since this project initiated. Following are the highlights of our achievements:

- Establishing the reaction systems for fast catalyst screenings. Various Batch-Type micro reactors, including sealed glass-tube reactors and stainless steel autoclaves, have been separately designed for the studies of model compounds and crude oils. All of them have been demonstrated to be working efficiently toward our specific requirements.
- Method of product analysis and reaction evaluation was established. For model compounds, primary gaseous products were quantified using GC analysis, while the unreacted acid was recovered using organic solvent through washing the reactor. The latter was subjected to another GC analysis and the acid conversion could be calculated herewith. For crude oil test, TAN measurements were employed to characterize the catalytic effect.

- Based on the computational studies which indicated that the acidities of naphthenic acids are generally structural insensitive, a set of model compounds has been selected for the purpose of the catalyst screening.
- Catalytic decarboxylation experiments of model compounds, and a mixed acid solution, were carried out using various catalysts, including commercially available chemicals and specifically developed catalysts. Among the early results, NA-Cat-1 exhibited the highest catalytic decarboxylation activity toward model NA compounds, which was recognized from the CO<sub>2</sub> formation.
- Detailed theoretical and experimental studies of the decarboxylation reaction catalyzed by NA-Cat-1 have been conducted.
- Another important progress is the observation of the decarboxylation effect of NA-Cat-2 toward model compounds. The CO<sub>2</sub> yield well matched the acid conversion; and the decarboxylation product, naphthalene, was also detected. This indicated a direct decarboxylation reaction occurred in the presence of NA-Cat-2.
- Mechanistic understandings of catalytic decarboxylation reactions based on the compounds modeling are continuing. Three major reaction pathways, namely concerted, radical and anionic, are being investigated. A series of stable, intermediate and transition state configurations are being studied.
- Directly applied the catalyst to the crude oil tests. Improvement of catalytic activity was achieved by doping an active metal on supports. As a result, the TAN could be significantly reduced.
- Two presentations regarding the naphthenic acid removal project were given in the 38<sup>th</sup> ACS Western Regional Meeting held in Long Beach, CA from October 15 to 18. The titles of our presentations are “Acidity and 1-Octanol/Water Distribution Coefficients of Naphthenic Acids” and “Decarboxylation of Naphthenic Acid Model Compounds”. The support of DOE was highly acknowledged.

## *Experimental*

### **1. Selection of Model Compounds and Oil Samples**

- a. *Model Compounds.* A set of carboxylic acids, such as naphthoic acid (C<sub>10</sub>H<sub>7</sub>COOH) and cyclohexane carboxylic acid (CHCA), has been selected as the model compounds to represent the aromatic and saturated naphthenic acids. A majority of catalytic tests with model compounds were performed with naphthoic acid.

- b. *Mixed Acid Solution.* To apply the catalysts to a reaction system that is close to the actual crude oil in composition, a mixed acid solution was prepared by dissolving five organic acids in dodecane. The structures and concentrations of these acids are listed in Table-1.

**Table 1: Composition of the mixed acid solution**

Carboxylic acids	Concentrations in dodecane (wt%)
Cyclopentane carboxylic acid (CPCA)	2.471
Cyclohexane carboxylic acid (CHCA)	1.927
Benzoic acid (BA)	0.871
Pentyl-cyclopentane carboxylic acid (C <sub>5</sub> H <sub>11</sub> -CHCA)	1.099
Heptyl-benzoic acid (C <sub>7</sub> H <sub>15</sub> -BA)	1.107

- c. *Oil Sample.* With several crude oil samples in hand, we measured their total acid number (TAN), which varied from 0.27 to 4.38. Among them, Texaco crude oil donated by ChevronTexaco with Total Acid Number (TAN) of 4.38 was selected for our oil test (Figure 1).

## 2. Catalyst Preparations

Commercially available chemicals could be directly used as catalysts or they can be subjected to the activation treatment. For instance NA-Cat-1 was calcined at 700°C for one-hour prior to use. For supported metal catalysts such as Cu/SiO<sub>2</sub>, metals were introduced by impregnating their salt solutions to supports followed by evaporation of water and hydrogen reduction at 400 °C, 50ml/min for 1 hr (Figure 1). For another type of supported metal catalysts, the active metal was doped by thermal decomposition of a metal-containing compound under controlled temperature and vacuum conditions.



**Figure 1: Catalysts and Crude oil Samples**

### 3. Experimental Setups and Operation Procedures

- a. *Sealed Glass Tube Batch Reactor (Figure 2)*. Acid sample, catalyst, and other additive (if any), in orders of milligram, were sealed in a glass tube under vacuum. The sealed glass tubes were placed in oven to start reaction under controlled reaction conditions. Typically, the reactions were carried out at the temperature range of 200 to 300 °C for 4 hrs. The reaction gas was collected and quantified in a vacuum line *via* a standard gas transfer method.

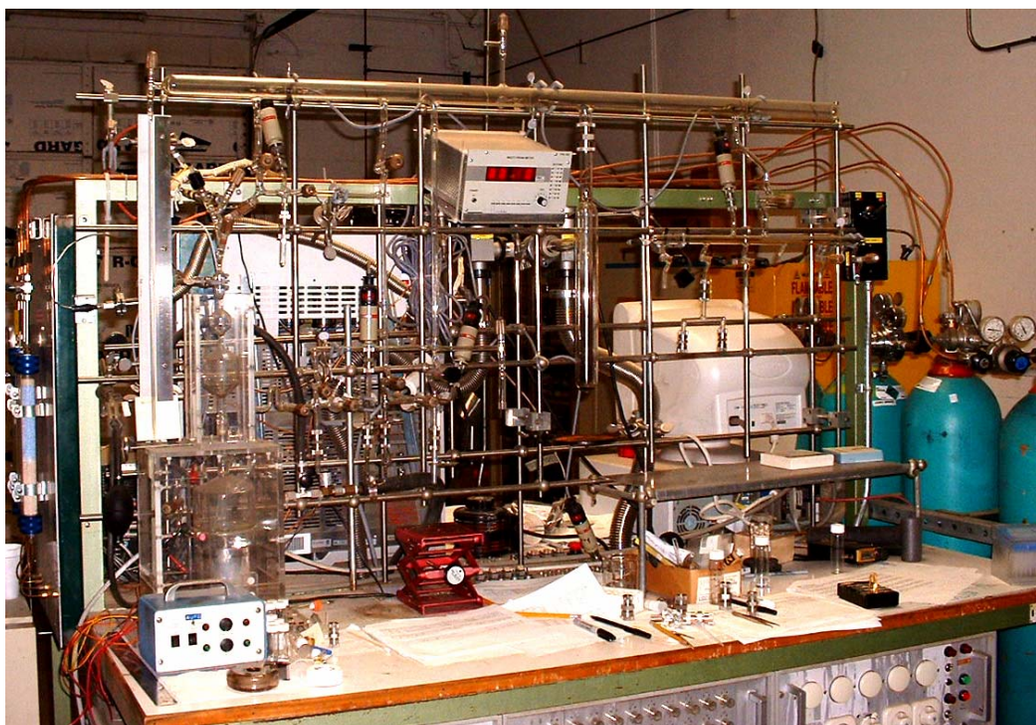


**Figure 2:** Various Micro Batch Type Reactors

- b. *Autoclave Reactor (Figure 2)*. For crude oil test experiments, as more sample amounts are needed for the TAN measurement, a different experimental procedure was established. An autoclave reactor with the volume of ca 40 ml was used to perform the reaction. The detail on the operation procedure can be described as the following:
  - i) loading ca. 12g oil and 0.24-0.60g catalyst (2~5 wt% of oil) to the reactor;
  - ii) premixing the two components by shaking the reactor for one hour;
  - iii) running the reaction at the temperature range of 250~300°C for 4 hr while keeping the reactor moving to achieve a good contact between the reactants and catalysts;
  - iv) after undergoing the reaction time, cooling the reactor and then recovering the treated oil through solvent extraction using dichloromethane as the solvent followed by vacuum filtration and evaporation of the solvent.

#### 4. Analysis Methods

- a. *GC Analyses.* The reaction gas was collected and quantified in a vacuum line via a standard gas transfer method (Figure 3). The gas was then analyzed with a GC, which is connected with the vacuum line. CO<sub>2</sub> as well as other gaseous products could be quantified herein, from which, the yields of CO<sub>2</sub> and other gaseous products could be calculated. For the reaction residue, using dichloromethane to recover the un-reacted acids and then subject to another GC analysis, from which acid conversion could be calculated.
- b. *Total Acid Number Measurement.* For crude oil tests, Total Acid Numbers (TAN), before and after reactions, were measured using a Standard Test Method for Acid Number of Petroleum Products by Potentiometer Titration, ASTM-D 664. Here TAN is defined as the KOH amount consumed in milligram to neutralize one gram of oil.



**Figure 3:** The Gas Transfer and Analysis Apparatus

#### 5. Theoretical Calculation Methods

The gas phase geometries of reactants, products, intermediates and transition states (TS) have been optimized using the B3LYP flavor of density functional theory. We have used the 6-31G (d) basis set for all of computations. All stationary points have positively identified for local minima (zero imaginary frequencies) and for TS (one imaginary frequency). Vibration frequencies are also calculated at all stationary points to obtain zero point energies (ZPE) and thermodynamic parameters.

## *Results and Discussion*

### 1. Catalytic Decarboxylation of Model Compounds

Table 2 lists the CO<sub>2</sub> generation during the catalytic reactions at 200 °C. Among the solid catalysts investigated, the amounts of the CO<sub>2</sub> generated from NA-Cat-1 catalyzed for both saturated and aromatics (30.38 and 33.20 ml/g, corresponding to the 17.4 % and 25.5 % mol conversion) are much higher than other solid catalysts. It should be noticed that no CO<sub>2</sub> formation does not necessarily mean no acid conversion, since there is high possibility of CO<sub>2</sub> being adsorbed by the metal oxides to form very stable carbonates. However, detection of CO<sub>2</sub> does demonstrate the conversion of acid compounds. In this sense, NA-Cat-1 has exhibited the highest reactivity towards the decarboxylation of the naphthenic acid compounds. Addition of the organic bases, such as pyridine, can slightly promote the catalytic reactivity. In the presence of pyridine, NA-Cat-1 catalyzed decarboxylation can even occur at 100 °C. Several transition metal oxides used here gave poor activities towards the catalytic decarboxylation at this condition.

**Table 2: Catalytic Decarboxylation of Model Compounds<sup>a</sup>**

Acid		Catalyst		Additive (mg)		CO <sub>2</sub> (ml/g)
Name	wt. (mg)	Name	Wt. (mg)	Name	wt. (mg)	
CHCA	49.3	NA-Cat-1	10.2			30.38
CHCA	50.9	NA-Cat-5	14.3			0.00
CHCA	51.9	BaO	11.5			0.02
CHCA	46.1	SrO	11.5			0.00
NA	51.5	None				0.00
NA	52.8	NA-Cat-1	19.7			33.20
NA	50.8	CuO	11.3			0.00
NA	52.4	Cu <sub>2</sub> O	11.7			0.00
NA	53.1	Ag <sub>2</sub> O	10			0.00
NA	50.5	Al <sub>2</sub> O <sub>3</sub>	10.1			0.00
NA	50.1	Ag <sub>2</sub> O	9.9	C <sub>5</sub> H <sub>5</sub> NO	52.8	4.40
NA	49.1	Cu <sub>2</sub> O	9.6	C <sub>5</sub> H <sub>5</sub> NO	48.4	5.60
NA	52.7	NA-Cat-1 <sup>b</sup>	21.3	Pyridine	56.2	20.80
NA	49.5	Ni/Al <sub>2</sub> O <sub>3</sub>	10.9			0.10
NA	49.2	Ni/SiO <sub>2</sub>	11.6			0.00

<sup>a</sup> Reaction temperature and time are 200 °C and 4 hrs,  
except for <sup>b</sup> reaction temperature and time for 100 °C and 4 hrs.

## 2. Catalytic Decarboxylation of the Acid Mixture

To test the effectiveness of the NA-Cat-1 series catalyst on decarboxylation of organic acids, a mixture of five acid compounds is prepared to partly simulate oil composition, but with higher concentrations of acids. As listed in Table 3, compared with no catalyst addition, high acid conversions were obtained from NA-Cat-1 for each acidic compound, which is similar with the case of single acid test. The acid conversions were further improved when small amounts of active metals were loaded on NA-Cat-1 (NA-Cat-3, NA-Cat-4), and the conversions even exceeded 90%.

**Table 3: Catalytic Decarboxylation of Acid Mixture<sup>a</sup>**

Mixture wt. (g)	Catalyst		Acid Conversions				
	Name	wt. (mg)	CPCA	CHCA	BA	C5H11-CHCA	C7H15-BA
2.4870	None		16.00	15.00	14.50	11.30	7.00
2.5261	Ni/Al <sub>2</sub> O <sub>3</sub>	25.80	5.00	5.70	15.00	10.70	4.80
2.5357	Ni/SiO <sub>2</sub>	25.50	21.50	18.90	15.40	16.00	8.70
2.5511	NA-Cat-3	25.50	70.80	70.50	91.20	92.40	97.50
2.3755	Cu/Al <sub>2</sub> O <sub>3</sub>	27.20	0.00	0.00	5.50	2.40	5.30
2.3806	Cu/SiO <sub>2</sub>	25.00	12.80	10.50	10.40	7.10	3.50
2.4960	NA-Cat-4	25.40	86.50	83.90	92.40	93.20	98.00
2.5432	NA-Cat-1	25.50	39.00	46.70	78.80	81.60	92.50

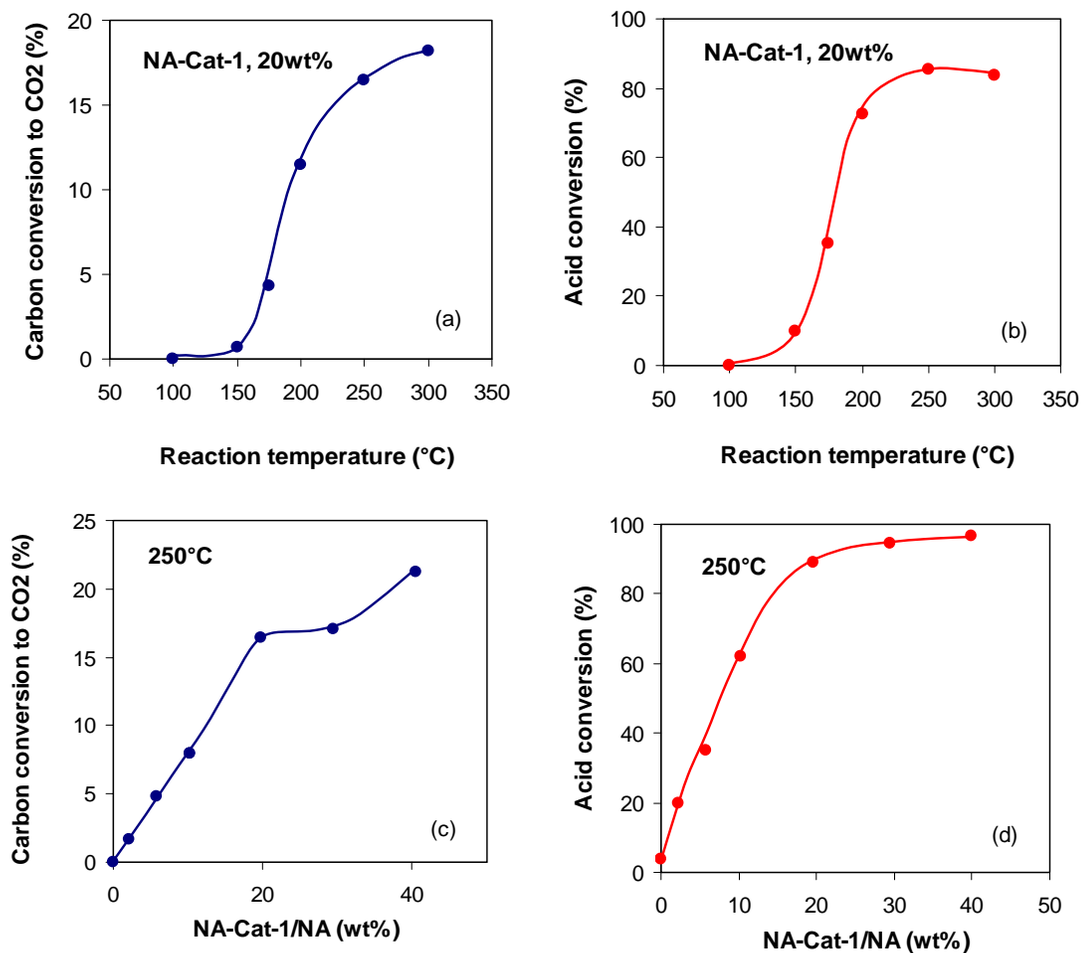
<sup>a</sup> CPCA, cyclopentane carboxylic acid; CHCA, cyclohexane carboxylic acid, BA, benzoic acid.

## 3. NA-Cat-1 Catalyzed Decarboxylation Reactions

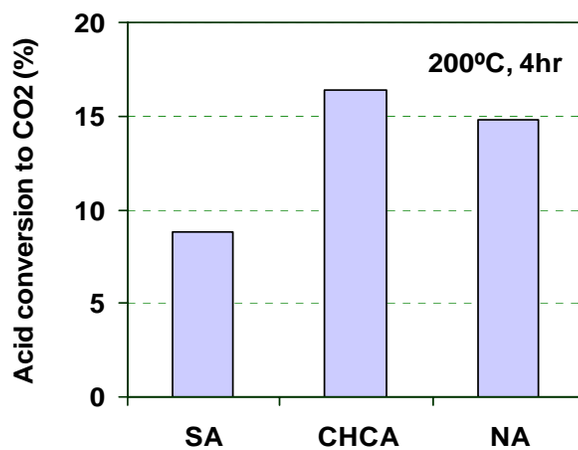
The effects of temperature and catalyst loading on naphthenic acid decomposition were investigated with NA-Cat-1. The reactions were separately run by changing reaction temperature in the range of 100 to 300 °C at a fixed NA-Cat-1 loading, 20 wt%, and changing the NA-Cat-1 loading from 0 to 40 wt% at 250 °C. Gaseous and remaining solid products were analyzed to obtain the data for CO<sub>2</sub> yield and acid conversion.

From the results shown in Figure 4, it can be seen that the reaction initialized at around 150 °C, followed by a sharp increase in range of 150-250 °C and then tended to level off at higher temperature. After 250 °C, more than 80% of acid has been converted, while the CO<sub>2</sub> yield was not as high. Increasing of the NA-Cat-1 loading in the range of 0-20 wt% linearly increases the CO<sub>2</sub> yield and the acid conversion, but further increasing of the NA-Cat-1 loading gave only limited enhancement.

On the role of NA-Cat-1, we currently consider its dual functionalities, namely catalytic decarboxylation and neutralization. Due to these two important features, it can be enrolled as a hopeful candidate to realize our two objectives.



**Figure 4:** NA-Cat-1 Catalyzed Decarboxylation Reactions  
(a, b) temperature effects; and (c, d) catalyst-loading effects



**Figure 5:** NA-Cat-1 Catalyzed Decarboxylation Reactions on Different Carboxylic Acids  
SA, Succinic acid; CHCA, Cyclohexane carboxylic acid; NA, naphthoic acid

To test the applicability of NA-Cat-1, we applied it to the model compound reaction with diverse reaction substrates (Figure 5). The three reaction substrates represent different acid structures, which are aliphatic, alicyclic and aromatic respectively. In each reaction, CO<sub>2</sub> formation was detected. This result indicates that NA-Cat-1 is able to promote catalytic decarboxylation for the acid substrates with different structures; in another word, this reaction is less acid-structure dependent.

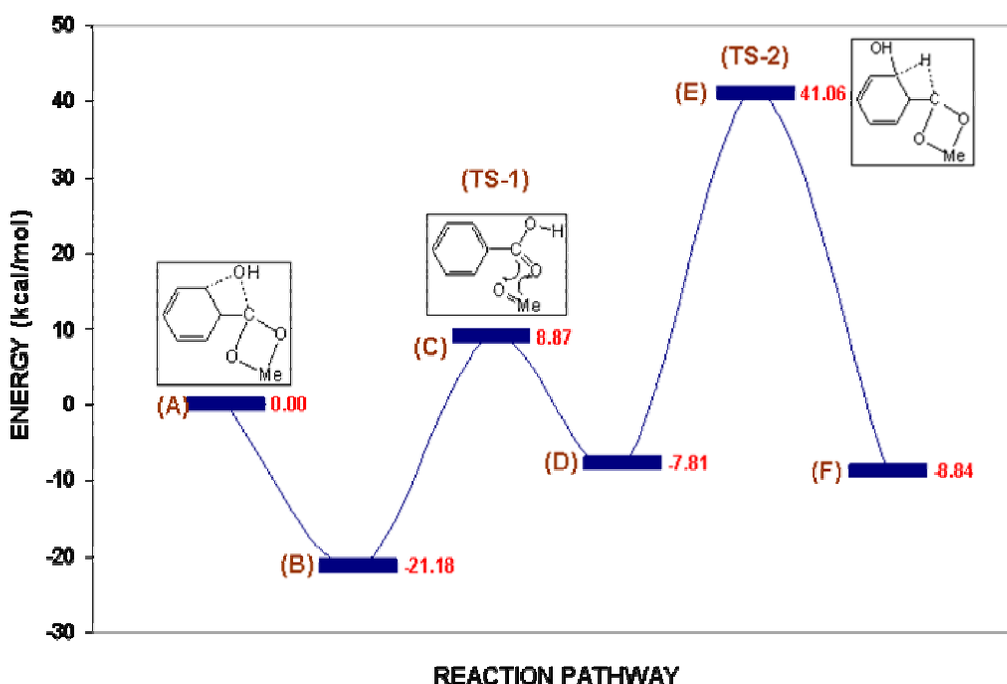
#### 4. Mechanistic Studies of the NA-Cat-1 Catalyzed Decarboxylation Reaction

A plausible concerted oxidative decarboxylation pathway in the presence of NA-Cat-1 has been theoretically studied in gas-phase with the energy diagram of all of stable, intermediate and transition states computed. The reaction path can be summarized as a three-step mechanism that converts benzoic acid to phenol (Figure 6):

Step 1: Nucleophilic attack at the C atom of the carboxyl group, from (A) to (B)

Step 2: Transfer of the hydroxy group *via* a 4-member ring transition state, from (B) to (D) through TS-1 (C)

Step 3: Proton transfer accomplishing by decarboxylation, from (D) to (F) through TS-2 (E).



**Figure 6:** A Plausible Concerted NA-Cat-1 Catalyzed Decarboxylation Pathway

The computed transition barrier for TS-1, featuring the attack of the hydroxy group on the *ortho*-position of the aromatic ring (~30 kcal/mol) is consistent with our experimental conditions (200 °C, 4 hrs). The barrier for TS-2, featuring proton transferring from *ortho*- to *ipso*- (49 kcal/mol), however, is higher than expected. This could be partially due to the fact that

only gas-phase single molecule calculations were performed. Further calculations with larger metal oxide clusters, and/or water assisting are expected to lower this barrier.

## 5. The CO<sub>2</sub> Yield vs. the Acid Conversion

Even though many of the metal oxides tested did not show activities towards the CO<sub>2</sub> generation, it does not necessarily mean that no acids have been removed. In fact, by comparing the CO<sub>2</sub> yield and the acid conversion in the presence of several metal oxides, as shown in Table 4, it can be seen that the acid conversions are generally higher than the CO<sub>2</sub> yields. This large difference could be due to either the formation of carboxylic acid salts through acid-base neutralization reaction, or the formation of alkaline earth metal carbonates through the adsorption of CO<sub>2</sub> on oxides. The latter should also be a signature of catalytic decarboxylation process. In a sense, the case of NA-Cat-2 is an ideal decarboxylation catalyst as the acid conversion is almost equal with its CO<sub>2</sub> yield and the decarboxylated product, naphthalene, was also clearly detected, whose carbon yield reached ca.20%. This result clearly identified the occurrence of a catalytic decarboxylation process.

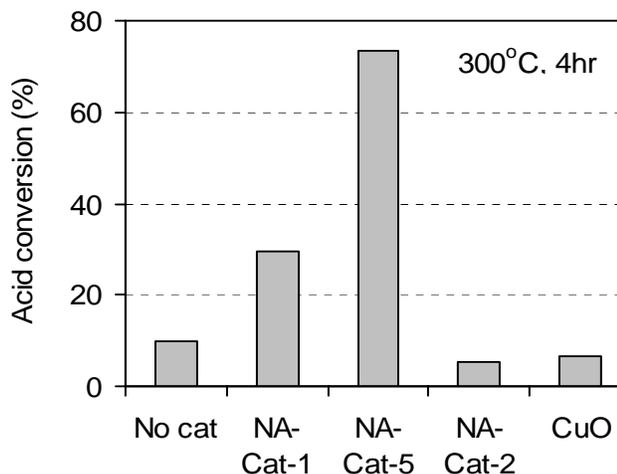
**Table 4: The CO<sub>2</sub> Yield and the Acid Conversion in the Presence of Several Metal Oxides**

Catalyst	Acid Conversion (%)	CO <sub>2</sub> Yield (%)
None	3.6	0.05
NA-Cat-1	81.6	17.1
NA-Cat-5	96.9	0
SrO	69.9	0
BaO	53.8	0.15
NA-Cat-2	53.7	53.1
CuO	63.3	17.2

Reaction temperature: 250°C; reaction time: 4 hr

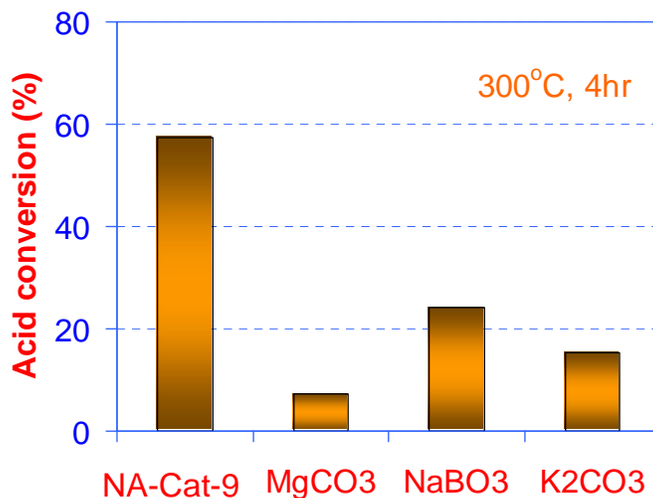
## 6. Test Runs with Crude Oil

The results of initial test runs of a group of metal oxide catalysts as reagents towards the decarboxylation reaction of crude oils are listed illustrated in Figure 7. Here the acid conversions were calculated based on the TAN reduction. NA-Cat-5 shows acid conversion as high as ~ 70%, while NA-Cat-1 did not show significant reactivity towards the acid removal as expected, and NA-Cat-2 and CuO almost had no positive influence to this process. It seems that there are many others



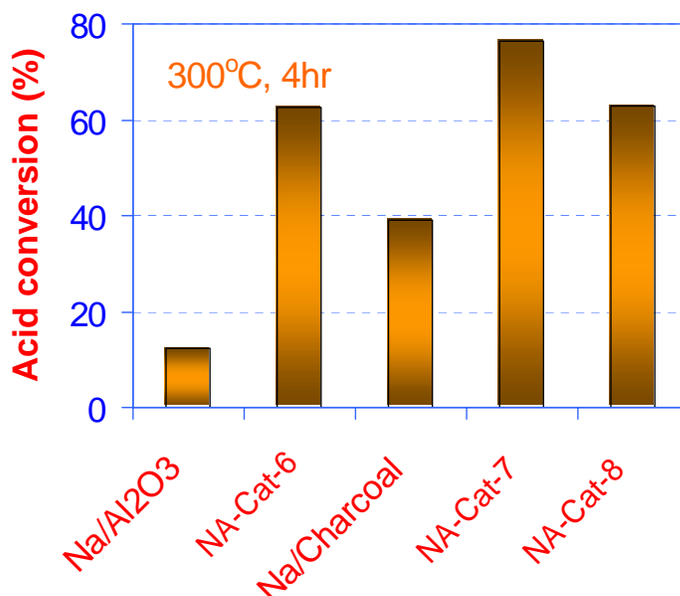
**Figure 7: Acid Conversion of the Texaco Oil Sample on Various Metal Oxide Catalysts**

factors that affect the performance of the catalysts in crude oil reaction. In particular, catalytic deactivation that resulted from the impurities in crude oils is an important issue.



**Figure 8:** Acid Conversion of the Texaco Oil Sample on Various Carbonate Catalysts

To improve the behavior of catalyst and promote the acid removal of crude oil, more autoclave tests were performed with a group of carbonates used as catalysts (Figure 8) and several active metal supported catalysts (Figure 9), which were specifically developed for this purpose. Among the carbonates tested, NA-Cat-9 gave relatively high acid conversion compared with others; NA-Cat-6, NA-Cat-7 and NA-Cat-8 were identified to be very effective in reducing the acidity of crude oil. These findings gave us new insight on the process that we are currently working with.



**Figure 9:** Acid Conversion of the Texaco Oil Sample on Various Active Meta-Supported Catalysts

## ***Conclusion***

In the first year of this project, we have established our experimental and theoretical methodology for the study of catalytic decarboxylation and acid removal from crude oil and obtained promising results.

- Our research revealed the effectiveness of several solid catalysts such as NA-Cat-1 and NA-Cat-2 for the catalytic decarboxylation of model compounds and mixed acid solution.
- Several improved catalysts were found to be very effective in acid removal from crude oil, and significant TAN reduction was observed.
- We have obtained preliminary understanding on the reaction mechanism and other aspects related to the projects.

## ***Work plan for the 2<sup>nd</sup> and 3<sup>rd</sup> years (2004 and 2005)***

Our efforts for the 2<sup>nd</sup> and 3<sup>rd</sup> years of this project will follow the outline as proposed in our proposal to i) develop an advanced catalytic system that can effectively remove naphthenic acid from model compounds and crude oils; ii) improve the fundamental understanding of the decarboxylation reaction mechanism; and iii) develop a readily adaptable naphthenic acid removal *via* solid liquid separation.

To achieve these goals, several action plans will be taken:

- More experimental tests of our developed catalytic systems in the crude oils,
- Design and synthesis of new solid catalysts that can significantly lower the acidity of crude oil and keep long catalytic life,
- More dynamic measurements with the developed catalysts for crude oil using time resolved multiple cold trap
- Detailed surface characterization of pure solid catalysts using XRD and XPS analysis.
- A detailed mechanistic understanding of decarboxylation reaction combining the theoretical and experimental efforts,
- Computational modeling of the adsorption of naphthenic acid on the solid surfaces,
- Establishing of experimental procedure to develop solid adsorbent approach,
- Develop and optimize a process for efficiently removing naphthenic acids from crude oil based on the experimental and the computational results.

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