2010 DOE Annual Hydrogen Program Review Meeting

Investigation of reaction networks and active sites in bio-ethanol steam reforming over Co-based catalysts

Acknowledgement US DOE

Umit S. Ozkan* (P.I.) I. Ilgaz Soykal Hua Song Xiaoguang Bao Burcu Mirkelamoglu

Heterogeneous Catalysis Research Group Department of Chemical and Biomolecular Engineering The Ohio State University Columbus, OH 43210



http://www.che.eng.ohio-state.edu/people/ozkan.html

**Ozkan.1@osu.edu* Project ID#: DE-FC36-05GO15033 June 8, 2010 PD001

This presentation does not contain any proprietary, confidential, or otherwise restricted information

Timeline

- Start Date May 1, 2005
 End Date April 31, 2010 (no-cost extension requested)
- ♦ 90% Complete

Budget

- Total project funding
 \$1,145,625 (DOE)
 \$299,715 (OSU cost share)
- ◆ Funding received in FY05
 >\$100,000(DOE)
 >\$10,458 (OSU Cost share)
- ◆ Funding received in FY06
 >\$185,000 (DOE)
 >\$147,987 (OSU cost share)

◆Funding received in FY07
 >\$290,473 (DOE)
 >\$67,316 (OSU)

◆Funding received in FY08
 >\$140,703 (DOE)
 >\$9,780 (OSU)

◆Funding received in FY09
 >\$429,449 (DOE)
 >\$54,163 (OSU)

Overview

Barriers

- A. Fuel Processor Capital Costs
- C. Operation and Maintenance
- D. Feedstock Issues

Partners

- Argonne National Laboratory
 - Advanced Photon Source Synchrotron techniques
- Chemistry Dept. at OSU (Prof. Hadad)
 Molecular simulation
- NexTech Materials, Ltd.
 - Catalyst manufacturing scale-up
- Directed Technologies Inc.
 - Economic analysis and feasibility considerations
- PNNL
 - Sharing findings



Relevance



Project overview

- Catalytic H₂ production from bioethanol
 - Renewable sources;
 - Plant matter (natural CO₂ recycling);
 - Lends itself well to distributed H₂ production strategy;
- Non-precious metal catalysts, lowtemperature operation
- A university program addressing many fundamental questions such as:
 - Catalytic active sites;
 - Reaction networks and mechanisms;
 - Surface species and intermediates;
 - Deactivation mechanisms;
 - Regeneration mechanisms.

Objectives-Relevance

- To acquire a fundamental understanding of the reaction networks and active sites in bioethanol steam reforming over Co-based catalysts that would lead to
 - Development of a precious metal-free catalytic system which would enable
 - □ Low operation temperature (350-550°C)
 - High EtOH conversion
 - □ High selectivity and yield of hydrogen
 - High catalyst stability
 - Minimal byproducts such as acetaldehyde, methane, ethylene, and acetone
 - Ease of operation
 - Addressing the barriers
 - □ Fuel Processor Capital Costs
 - Operation and Maintenance
 - Feedstock Issues



Approach Milestones and Progress



IVERSITY			
	Tasks	Progress	% comp
1	Setting up the experimental systems, establishing protocols, training	Built, installed a reactor system and established experimental protocols; Completed a HAZOP review for the ethanol reforming reactor system; Went through a safety inspection conducted by a DOE Safety Panel.	100%
2	Economic analysis	Completed an initial economic analysis based upon a 1,500 kg/day of hydrogen process using H2A model.	95%
3	Catalyst synthesis and optimization	Prepared over 100 batches of catalysts; Investigated the effect of support, promoters, precursors, active metal loading, synthesis techniques and parameters on catalytic performance.	95%
4	In-situ, pre- and post- reaction characterization	Completed characterization studies using CO, N ₂ O and H ₂ chemisorption, X-ray diffraction, temperature programmed reduction, oxidation, desorption, and thermogravimetric analysis, differential scanning calorimetry. Combined characterization studies (XPS, DRIFTS, LRS, TGA) with mechanistic investigations to understand the pre- and post-reaction nature of the catalyst, reaction mechanisms and deactivation characteristics; XANES and EXAFS studies using the Synchrotron Facilities at Argonne National Laboratories Advanced Photon Source.	90%
5	Activity tests, kinetic studies, deactivation, regeneration studies	Established correlations between synthesis parameters, structure and activity; Evaluated synthesized catalysts for ethanol steam reforming in the temperature range 200- 500°C and gas hourly space velocities from 5,000-300,000h ⁻¹ ; Conducted transient and steady-state experiments to understand the reaction network and mechanisms; Examined the role of pre-treatment procedures. Initiated studies with dimethyl ether (DME) steam reforming.	90%
6	Information dissemination	Published 12 articles in refereed journals, five articles in conference proceedings, and gave 27 presentations, one key-note lecture and several invited lectures.	90%
7	Data analysis, reproducibility tests, literature awareness	Built a literature data base; Data analysis and reproducibility tests are performed in conjunction with the experiments.	90%





Technical Accomplishments and Progress Understanding how to control the most important catalytic properties











Technical Accomplishments and Progress Understanding the Reaction Network





"Reaction Networks in Ethanol Steam Reforming over Supported Co Catalysts," Industrial and Engineering Chemistry Research, accepted



50 150 250 350 550 650 450 Desorption Temperature (°C)

 $CH_{3}CH_{2}OH \rightarrow CH_{3}CHO + H_{2}$ $CH_{2}CH_{2}OH \rightarrow CH_{4} + CO + H_{2}$ $CH_3CH_2OH \rightarrow C_2H_4 + H_2O$ $CH_{3}CHO \rightarrow CH_{4} + CO$ $2CH_3CHO \rightarrow CH_3COCH_3 + CO + H_2$ $CH_{2}CHO + 3H_{2}O \rightarrow 2CO_{2} + 5H_{2}$ $CH_{2}CH_{2}OH + H_{2}O \rightarrow 2CO + 4H_{2}$ $CH_{3}CH_{2}OH + 2H_{2}O \rightarrow 2CO_{2} + 6H_{2}$ $CO + 3H_2 \rightarrow CH_4 + H_2O$ $CH_4 + 2H_2O \rightarrow CO_2 + 4H_2$ $CO + H_2O \rightarrow CO_2 + H_2$ $CO_2 + H_2 \rightarrow CO + H_2O$ $C_2H_4 \rightarrow coke$ $CH_4 \rightarrow C + 2H_2$ $2CO \rightarrow CO_2 + C$





Technical Accomplishments and Progress Understanding the major deactivation mechanisms





Findings from this project that allow us to prevent deactivation

- Increasing oxygen mobility by creating anion vacancies in the support
- Neutralizing acidic sites
- Improving dispersion by using organic ligands in the synthesis
- Improving activity to reduce the operating temperature
- Eliminating the need for pre-reduction with H₂

Results published in

Song, H., Ozkan, U.S., "Ethanol Steam Reforming over Co-based Catalysts: Role of Oxygen Mobility," *J. Catalysis*, **261** 66-74 (2009). Song, H. Ozkan, U.S., :Changing the Oxygen Mobility in Co/Ceria Catalysts by Ca Incorporation: Implications for Ethanol Steam Reforming," Invited paper. *Journal of Physical Chemistry A*. **114**, 3796-3801 (2010).

Technical Accomplishments and ProgressA catalyst formulation with high activity, selectivity and stability







- Over 95% H₂ yield
- No liquid by-products above 300°C
- Stable activity in accelerated aging study

Results published in

Song, H., Tan, B. and Ozkan, U.S., "Novel Synthesis Techniques for Preparation of Co/CeO2 as Ethanol Steam Reforming Catalyts," *Catalysis Letters*, **132**, 422-429 (2009)





Technical Accomplishments and Progress Understanding the effect of catalyst pre-treatment



New capability New collaboration New insight

XAFS spectroscopy was utilized to provide insight on the oxidation state and the local coordination environment of Co during ethanol steam reforming with Co/CeO₂

Recent

rogres

X-ray absorption spectroscopy capability was recently added

XAFS (X-ray Absorption Fine Structure) spectroscopy provides information on the local atomic structure of a selected element, including:

- formal oxidation state
- coordination
- bond distances
- nature of immediate neighbors

Advanced Photon Source (APS) at the Argonne National Laboratories was utilized

The APS is a third generation synchrotron facility with the nation's most brilliant X-ray beams









Technical Accomplishments and Progress

Vibrational spectroscopy under actual reaction conditions (operando)

DRIFTS during Temperatureprogrammed desorption

 Thermal transformation of adsorbed species
 Provides limited information on the surface species during ethanol steam reforming





Acetaldehyde formation and decomposition to CH_4 and CO_2 can be directly observed.

Recen

Progres

DRIFTS during ESR Reaction

Spectra collected while reaction is progressing
 Provides information on the surface species and mechanistic steps under true reaction conditions



T · H · E OHIO STATE UNIVERSITY

Technical Accomplishments and Progress

Dimethyl ether steam reforming over Co-based catalysts



4000 3600 3200 2800 2400 2000 1600 1200 800 Wavenumbers (cm⁻¹)



 Multi-source clean fuel which can be produced from a wide spectrum of feeds including lignocellulosic biomass

Inert, non-carcinogenic, nonmutagenic, noncorrosive, and virtually non-toxic

Hydrogen content is equal to that of ethanol

In-situ DRIFTS data show that Co/CeO₂ catalysts can be used in steam reforming of DME

Formation of various adsorption and oxidation products observed on the surface

 M-OH 3650~3150cm⁻¹, O-H stretching
 CH₃-2962, 2927, 2865 cm⁻¹:C-H stretching 1385cm⁻¹: CH₃- bending
 Surface Carbonates and methoxy region 1176, 1166, 1101, 1079cm⁻¹ v(OC)

• CO₂

2361, 2336cm⁻¹: O=C=O stretching

 Molecularly adsorbed H₂O 1654cm⁻¹

Recen

rogres



Technical Accomplishments and Progress Active and selective catalysts for PROX



- Co/CeO₂ nanoparticles were found to have high activity as Preferential Oxidation (PROX) catalysts for removing CO from H₂ streams.
- In the 150-180°C, high CO conversion (~100%) and high O₂ selectivity to CO₂ (>90%).
- Examined the effect of O₂ concentration, H₂ concentration, space velocity, temperature on conversion and selectivity
- Examined possible side reactions (WGS, r-WGS, methanation)
- Examined time-on-stream activity
- Measured activation energies for CO oxidation and H₂ oxidation









Results published in

Woods, M. P. Gawade, P., Tan, B. Ozkan, U.S., "Preferential Oxidation of Carbon Monoxide on Co/CeO₂ Nano-particles," *Applied Catalysis B*: in press. doi:10.1016/j.apcatb.2010.03.015



Collaborations



- A new collaboration with Argonne National Lab (Dr. Jeffrey Miller and Dr. Christopher Marshall) has been initiated to incorporate XAFS studies in the project. Three consecutive proposals for "In-situ XAFS Characterization of cobalt-based catalysts for steam reforming of bio-derived liquids" have been submitted to Argonne National Laboratory to request beam time at the synchrotron facilities and they all have been accepted. Seven members of the P.I.'s team have registered as facility users at Advanced Photon Source of Argonne National Laboratory. Dr. Miller has been working closely with the team members in in-situ experimentation and data analysis.
- Collaboration with Chemistry Department (Prof. Hadad) on combining experimental work with molecular simulation has progressed. A paper, that resulted from this collaboration "Computational Study of the Ethanol Steam Reforming Reaction over Co/CeO₂(111): from Ethanol to Acetate" has already been presented at the ACS National Meeting in August 2009. A refereed journal article written jointly has been accepted for publication in *Journal of Physical Chemistry*. A second collaborative article is in preparation. A post-doctoral researcher and a graduate student are being jointly advised by the P.I. and Dr. Hadad.
- Collaboration with Nextech Materials, Inc. for scale-up of the catalyst manufacturing process is in place. A joint proposal (OSU and NexTech) to NSF has been accepted.
- Collaboration with PNNL. Sharing findings
- Directed Technologies, Inc. Economic analysis and feasibility considerations

Summary

Relevance:



A fundamental study aimed at developing precious-metal free catalysts for hydrogen production from bio-ethanol and other bio-derived liquids

Approach:

An integrative approach that combines catalyst synthesis, characterization, kinetic studies and molecular simulation in an iterative manner to develop a fundamental understanding of the nature of active sites, reaction networks and reaction mechanisms involved in ethanol steam reforming over Cobased catalysts

Technical accomplishments and progress:

Developed an understanding of the reaction networks and surface mechanisms that allow correlating specific sites with specific reaction steps and deactivation mechanisms

Developed techniques for tailoring the catalytic properties to meet the needs of the reaction

Developed promising catalyst formulations that can achieve high H_2 yields, high selectivities and high stability at low temperatures

Collaborations:

New partnership with Argonne National Lab for XAFS studies

Active partnership with Chemistry Department on molecular simulation

Partnership with NexTech for catalyst manufacturing

Partnership with PNNL and Directed Technologies, Inc. for sharing findings

Proposed Future work:

Long term stability tests, accelerated deactivation tests, kinetic measurements to obtain kinetic parameters and allow kinetic modeling, *in-situ* XAFS studies



Future Work

- Kinetic and mechanistic studies will be continued in conjunction with catalyst characterization under reaction conditions to obtain kinetic parameters. Rate expressions will be developed for kinetic modeling.
- Long term time-on-stream experiments will be continued.
- Accelerated deactivation and regeneration studies (e.g., higher C/S ratio and GHSV) will be continued.
- Economic analysis will be fine-tuned based on process modification with no pre-reduction.
- Molecular simulation work using DFT calculations will be continued and used to guide rational catalyst design. This is a collaborative activity.
- In-situ XAFS experiments will be performed at Argonne National Lab to elucidate the oxidation state and coordination environment of cobalt sites during reaction. Accelerated deactivation studies will be coupled with in-situ XAFS studies. This will be a collaborative activity.
- Catalyst synthesis for best performing catalysts will be scaled-up through industrial partnership. This is a collaborative activity.
- Several more publications are in preparation and will be submitted within the next 6 months.