A Novel Slurry-Based Biomass Reforming Process

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Project ID #

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Overview

Timeline

- 1 May 2005
- 31 Oct 2008
- 2%

Budget

- Total project funding
 - \$2.9 million, DoE
 - \$737k, cost share
- \$0, FY04
- \$300K, FY05

Barriers

- Barriers:
 - V. Feedstock Cost and Availability
 - W. Capital costs and efficiency of technology
- Barriers Addressed
 - Technology Energy Efficiency
 - Capital Cost
 - Feedstock Flexibility

Partners

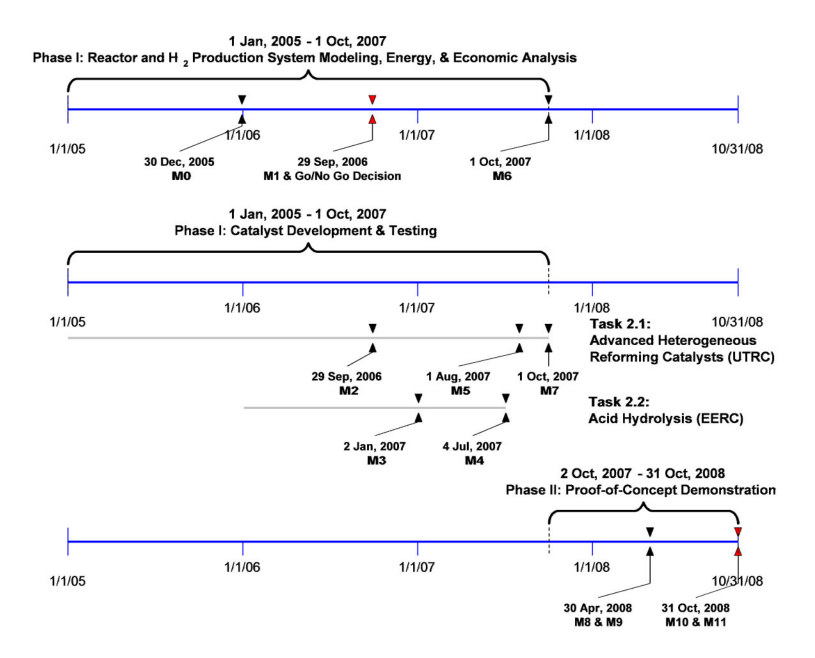
University of North
 Dakota Environment
 Energy Research
 Center

Biomass Slurry Reforming Objectives

DOE: \$1.75 kg 99.9+% H₂ with an LHV efficiency of 50%

- 1. Determine LHV Efficiency Using HYSYS
 - Major efficiency determinants and impact of catalyst efficiency/selectivity
 - Required hydrolysis rate per in unit input energy
 - Capital and energy cost of intermediate hydrogenation step
- 2. H₂ Cost via H2A Spreadsheet: Plant Cost, Rate of Return & Feedstock Costs
- 3. If DOE Cost and Efficiency Targets Can Be Met, Commence Next Phase
 - Optimum hydrolysis conditions: Energy and Capital Cost
 - Hydrolysis product chemical composition and physical properties
 - Sugar identification and concentrations
 - Identification and quantification of low molecular weight organics
 - Solubility, AMW and surfactant/foaming properties of lignin fraction
 - Catalysis discovery and testing
- 4. Micro-scale continuous operation of membrane reformer with batch hydrolysis
 - ~500 hr catalyst performance test
 - Collection of material and heat balance data important for plant design
- 5. Final Economic and Energy Analysis for Final Report

Project Schedule



Approach: Biomass Slurry to Hydrogen Concept

Slurry of ~ 10 % Ground
Biomass (Wood) in Dilute Acid
44% cellulose
19% hemicellulose
13% "other"
23% lignin
<1% "ash"
<1% protein

1 or more

Hydrolysis Steps Reformer Feed

~41% soluble C₆ and (C₆)_n "sugars"

~18% soluble "C₅" sugars

~10% "reformable others"

~31% lignin+cellulose fragments etc.

Hydrolysis targets

Preferential RCHO Hydrogenation Catalysts



High Selectivity Pt-MM rafts on engineered nano-structured oxide like Ti_[1-(x+y)]Dp1_xDp2_yO₂

~83 g 99.9+ H₂ / kg dry Feed Recovered Through Membrane

~9 g H₂ or Equivalent as fuel gas

~300 g Lignin and other fuel

~1 kg CO₂

Pt-Re/Ce $_{[1-(x+y)]}Zr_xDp_yO_2$ WGS Catalysts have high activity and very low CH_4 make

Optional Sugar Hydrogenation

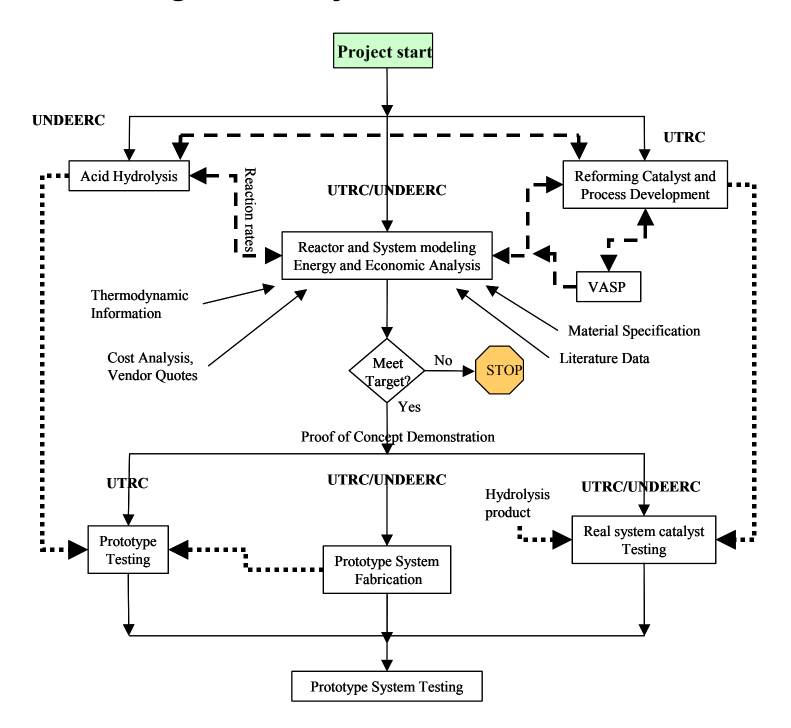
~59% sugar alcohols

~10% "reformable others"

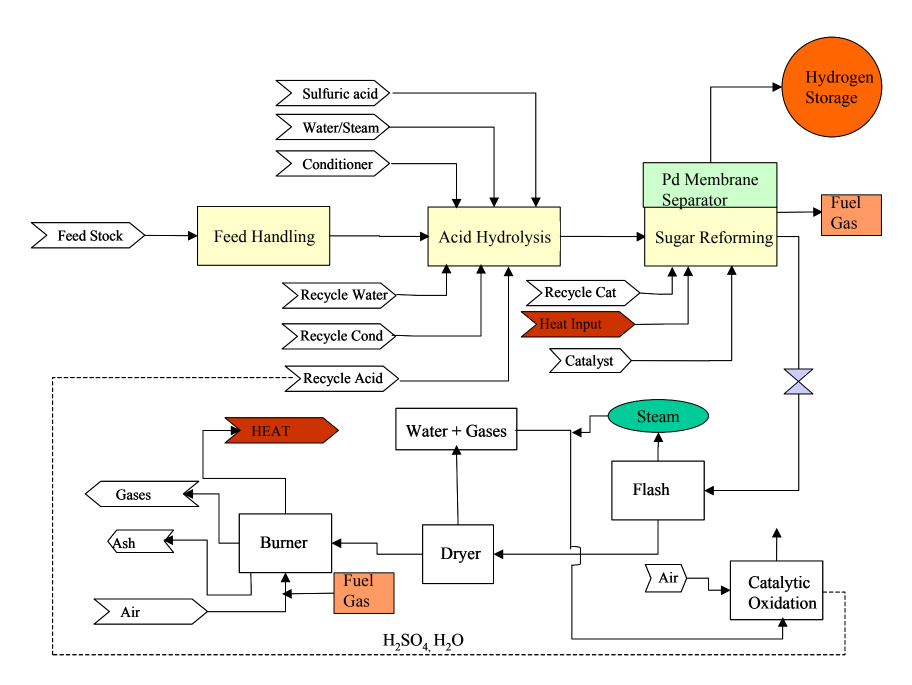
~31% lignin + cellulose fragments, etc.

Only if advanced catalysts seem unlikely reach g H₂ / kg feed goals

Original Project Plan Overview

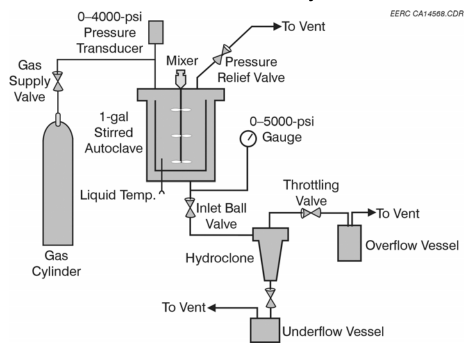


Approach: Initial Process Inputs and Outputs



Approach: Experimental Design to Optimize Hydrolysis

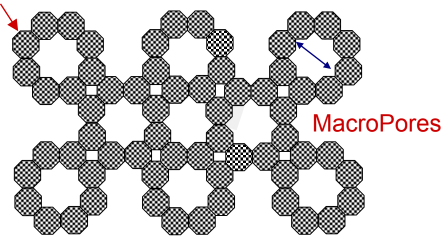
- Overall efficiency depends on optimizing hydrolysis energy / acid requirements
 - Lower acid concentration
 - + Less expensive alloys etc.
 - + Higher SA & activity reforming catalysts = smaller reforming reactors
 - + Less unnecessary chemical degradation = higher H₂ yield
 - Lower Temperature
 - + Increased residence time thus larger volumes and increased costs
 - + Lower autogenous steam pressures = lower capital costs
 - + Less expensive alloys etc.
 - + Less dehydrogenation etc. = higher H₂ yields
- Poplar assumed to be initial feed; grinding energy similar to mechanical pulping
- Input data for refined economic and efficiency model



Nano-Engineered Noble Metal / Doped Metal Oxide Catalyst

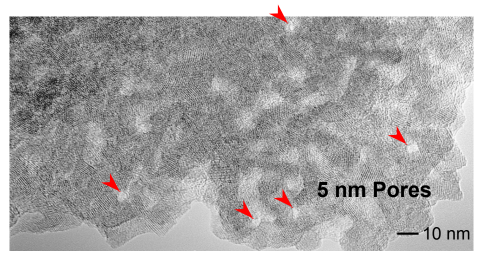
Design & synthesize active oxide structure to maximize accessible sites/vol.

Nanoparticle (< 3.5 nm) Micropore (≥ 5 nm)



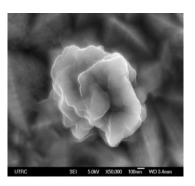
Conceptual Porous Metal-Oxide Framework
Shown in 2D

Self assembly used to create high surface area, large pore thermally stable active oxide support with 100% dispersed 2 wt% Pt based mixed metal clusters

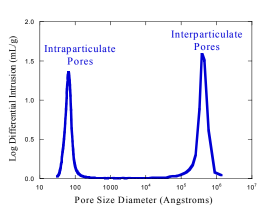


Conceptual Structure Realized

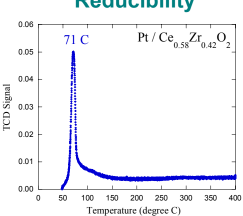
Fractal Morphology



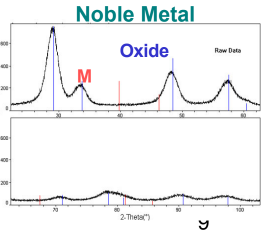
Large, Bimodal Pore Structure



Low Temperature Reducibility

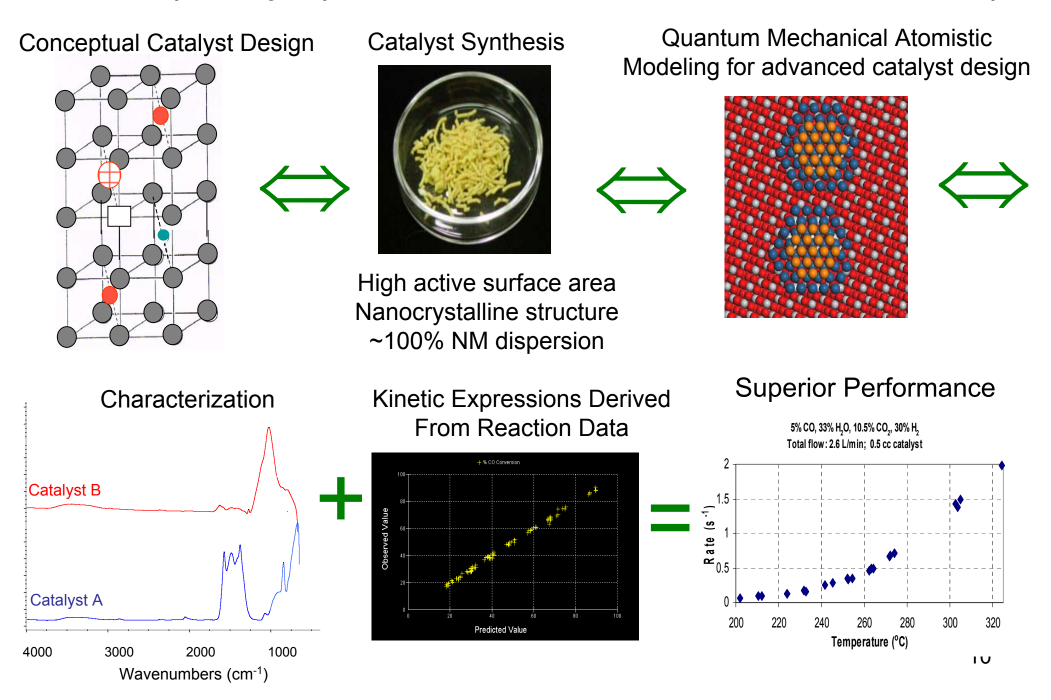


100% Dispersed
Noble Metal

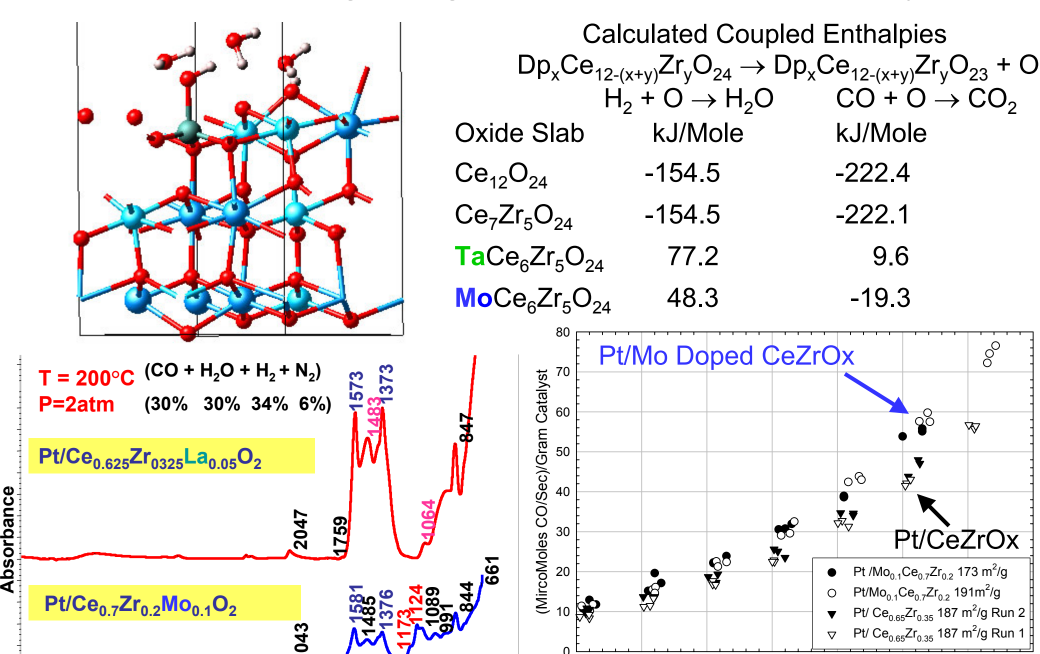


UTRC Catalyst Discovery Approach

Atomistic catalyst design, synthesis, characterization, reaction studies & kinetic analysis



VASP Modeling Insights Led To Better Catalysts



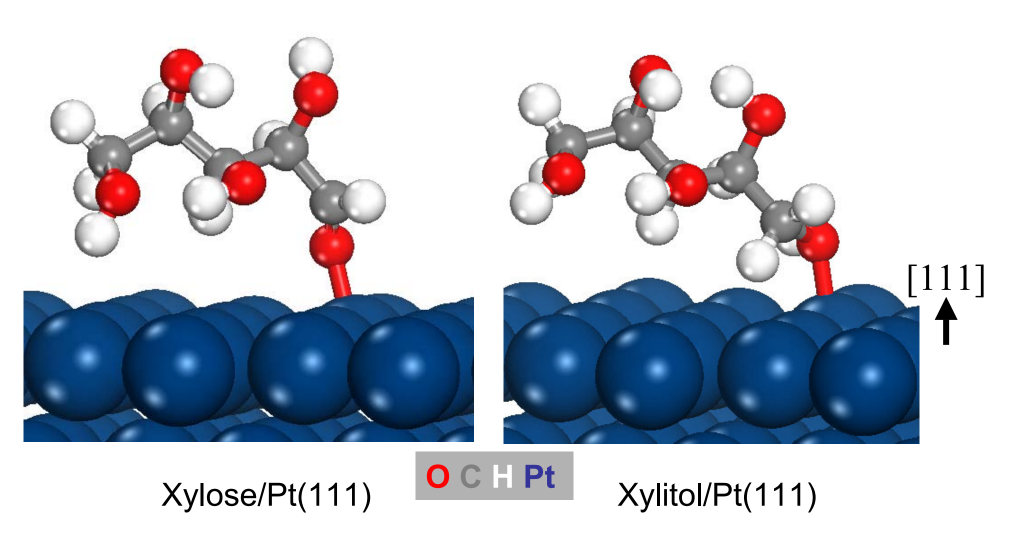
Wavenumbers (cm-1)

Higher Activity Catalyst w Similar Pt & SA

Temp C, Initial Down Ramp 4.9% CO, 10.5%CO₂, 33%H₂O, 30.3%H₃

Xylose Adsorbs More Strongly Than Xylitol on Pt(111)

Aldehyde O forms stronger bond than terminal alcohol O

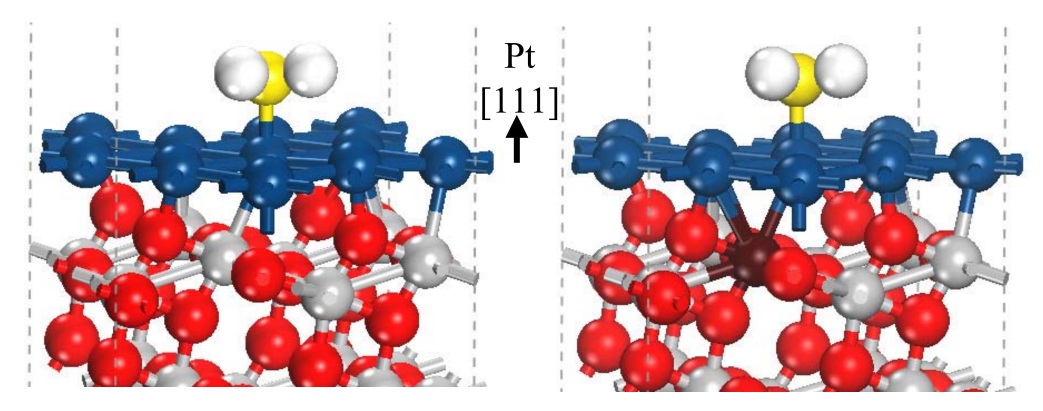


Binding Energy = -92 kJ/mole Binding Energy = -65 kJ/mole

Negative binding energy indicates exothermic process

Ce Dopant in TiO₂ Decreases H₂S-Pt Binding 16%

- Early results for Pt raft system, before full relaxation
- Anatase (101) TiO₂ with and without Ce



 $Pt(111)_{1ML}/AnataseTiO_2(101)$

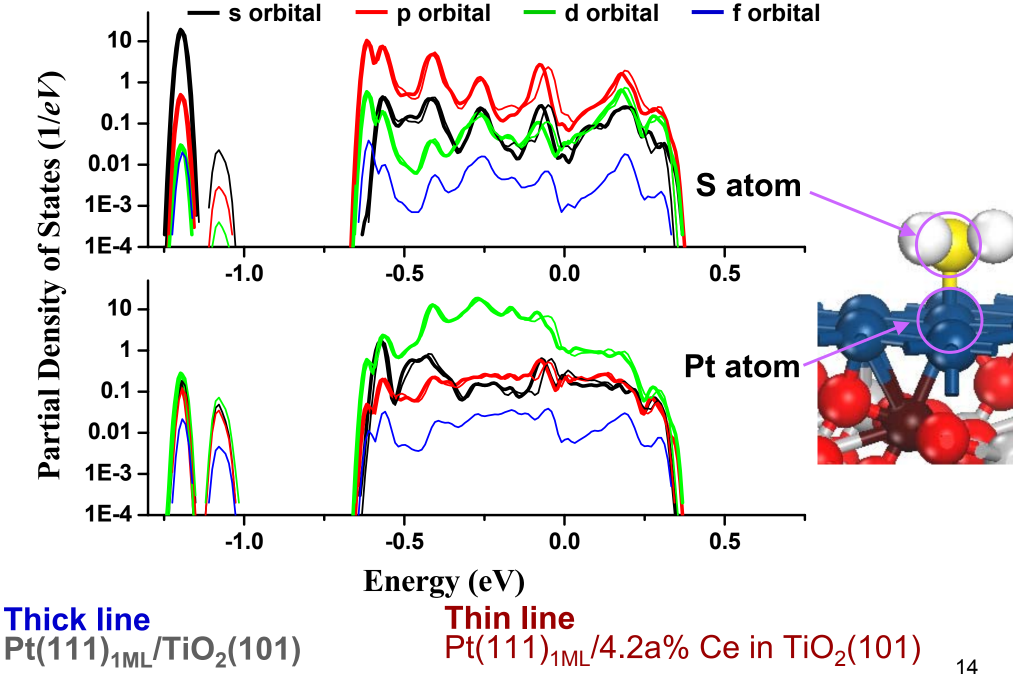
Pt(111)_{1ML}/4.2a% Ce_Anatase_TiO₂(101)

Binding Energy -106.53 kJ/mole

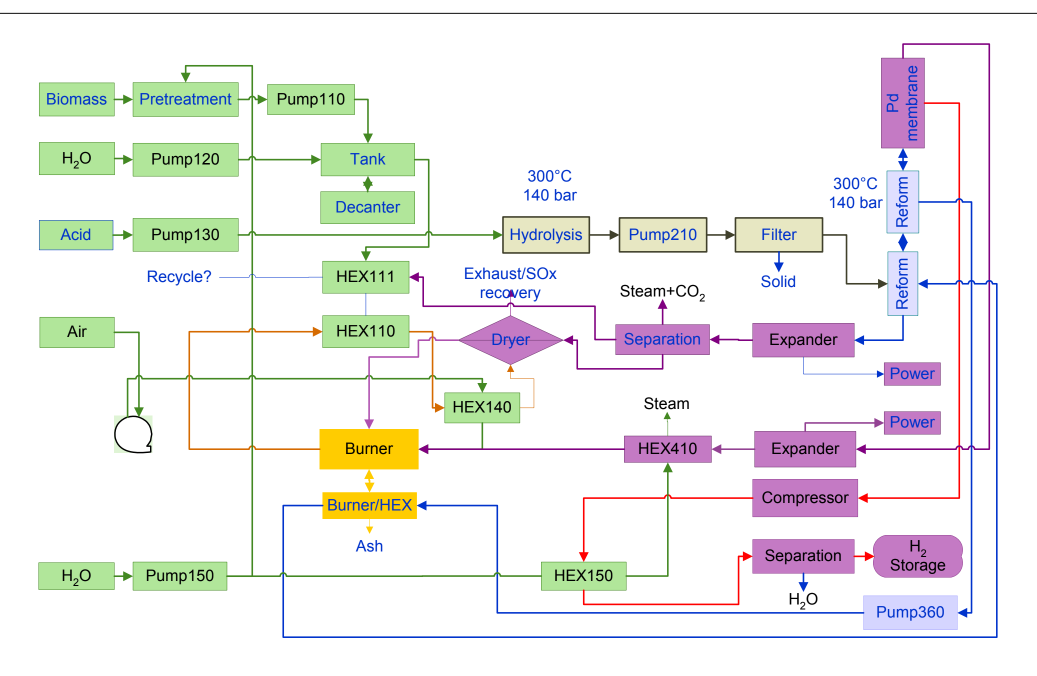
Binding Energy -89.50 kJ/mole



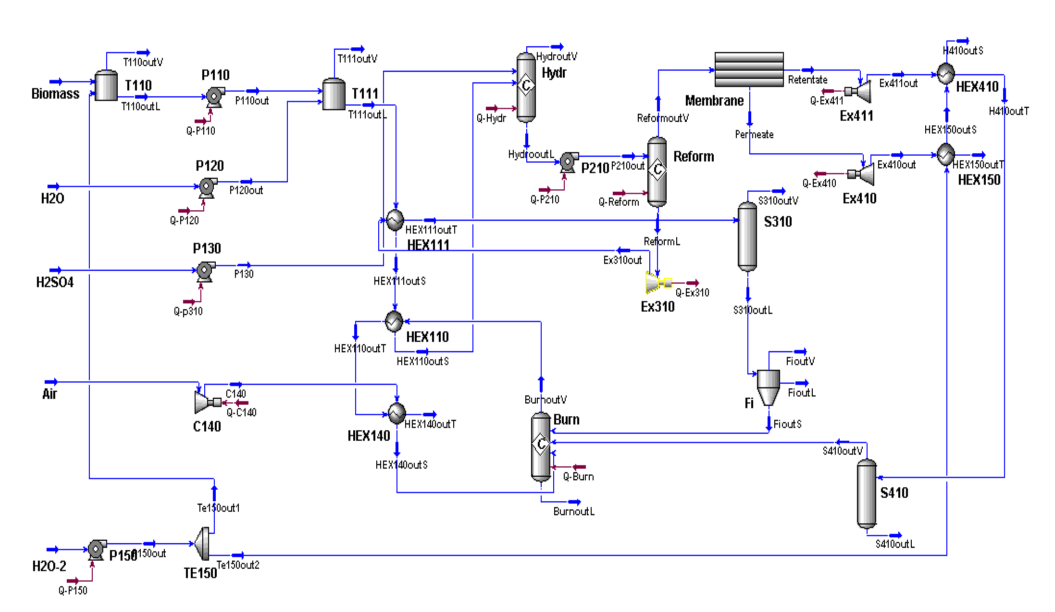
Oxide Dopant Shifts Pt & S DOS to Higher Energy



Progress: Conceptual Process Flow Diagram



Progress: Current HYSYS Process Flow Diagram



Future Work

• FY 2005:

- Initial feasibility analysis of a 2000 ton/day (dry) plant design showing a viable path towards the DOE's 2010 efficiency (50% LHV) and cost (\$1.75/kg H₂) targets.
- Low-level construction of catalyst synthesis & testing infrastructure

FY 2006:

- Is there a preliminary 2000 ton/day (dry) biomass plant design that could reach the DOE's 2010 efficiency (50% LHV) and cost (\$1.75/kg H₂) targets?
- GO/NO GO decision.
- Demonstrate an acid tolerant, model sugar solution reforming catalyst
 - + Promising kinetics and selectivity
 - + Path for cost-effective scale up (mass production) exists
- Identify preliminary hydrolysis conditions at UND-EERC and hydrolyzed product chemical composition and physical properties

Future Work

• FY 2007:

- Demonstrate effective hydrolysis conditions for actual biomass system and a path to scale-up for a viable plant design
- Demonstrate in the lab a potentially long lived, cost effective liquid phase biomass slurry reforming catalyst giving ~0.1 moles H₂/Total Pt equivalent-second
- Demonstrate that a plant designed with experimentally determined hydrolysis and reforming rates and conditions meets 50% LHV efficiency and \$1.75 /kg H₂
- Demonstrate wash coating of active catalyst on to selected support

FY 2008:

- Identify optimum hydrolysis conditions
- Demonstrate wash-coated reforming catalyst with actual hydrolyzed biomass
- Design, build, test and deliver proto-type continuous micro-scale reforming reactor to UND-EERC
- Complete 500 hrs of reformer operation and collect data important to full scale pilot unit design
- Estimate H₂/kg cost and LHV efficiency using 2000 T/day plant design finalized with actual batch hydrolysis and continuous micro-scale reforming reactor data.

Hydrogen Safety

The most significant hydrogen hazard associated with this concept is the 10% H₂ content of the up to 2000 psig process gas.

Hydrogen Safety

Our Approach to deal with the hazard in the laboratory is:

- H₂/Flammable gas detectors and ventilation interlock
 - System alarms if > 10% LFL (0.4% H₂) detected
 - All heater power and flammable gas flows shut off if either >25% of lower flammable limit (1% H₂) detected, or drop in ventilation rate
 - System design limits flammable gas flows to <10% of lower flammable limit based on measured ventilation rate

Hydrogen Safety

Our Approach to deal with the hazard in the proposed micro-scale demonstration unit is:

- Multiple H₂/Flammable gas detectors
- System alarms if >10% LFL (0.4% H₂) detected
- All heater power and flammable gas flows shut off if
 >25% of lower flammable limit (1.0% H₂) detected at unit.
- N₂ purging of all potential sources of ignition