

A Novel Slurry-Based Biomass Reforming Process

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Project ID #

PD 14

This presentation does not contain any proprietary or confidential information

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

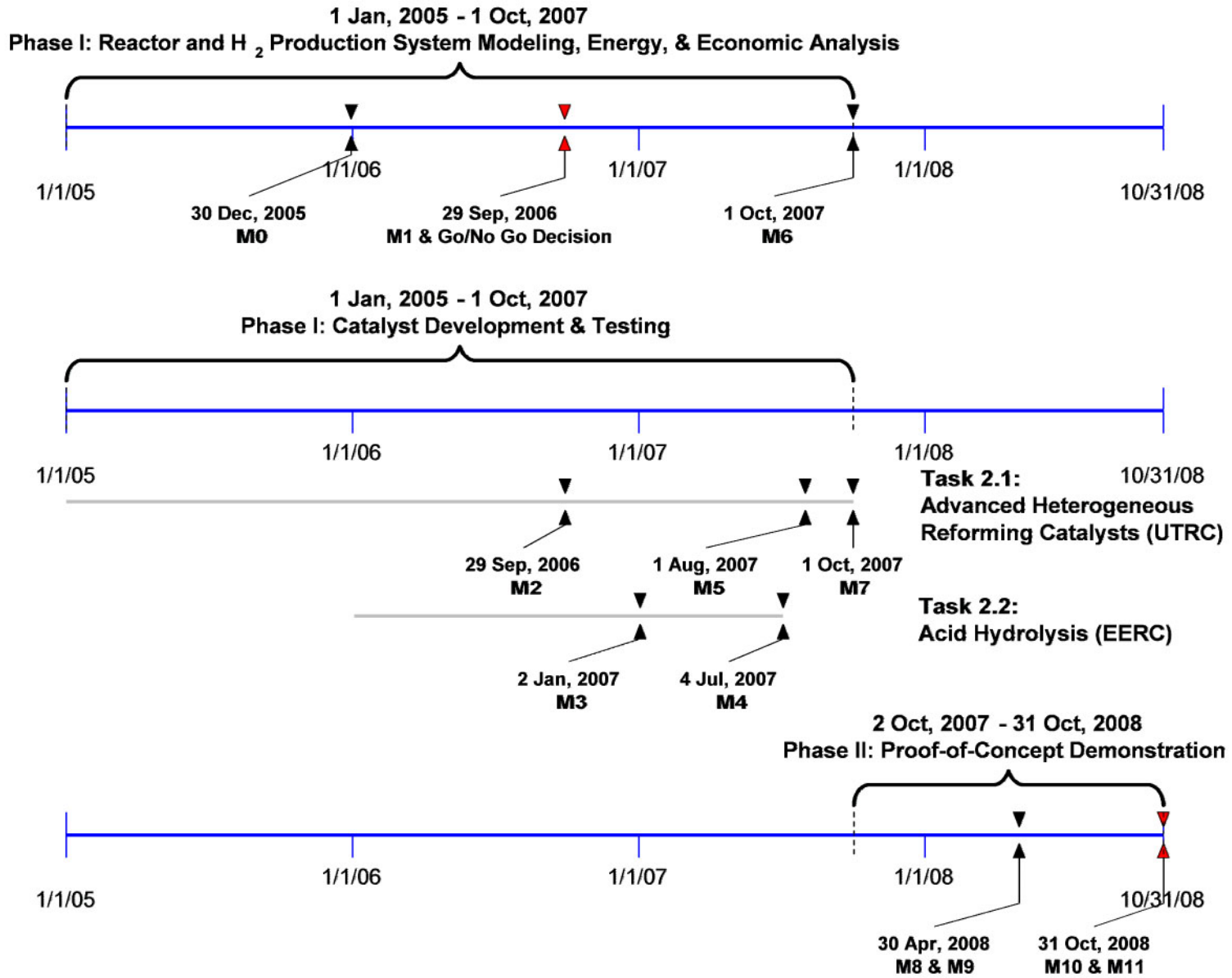
- University of North Dakota Environment Energy Research Center

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

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

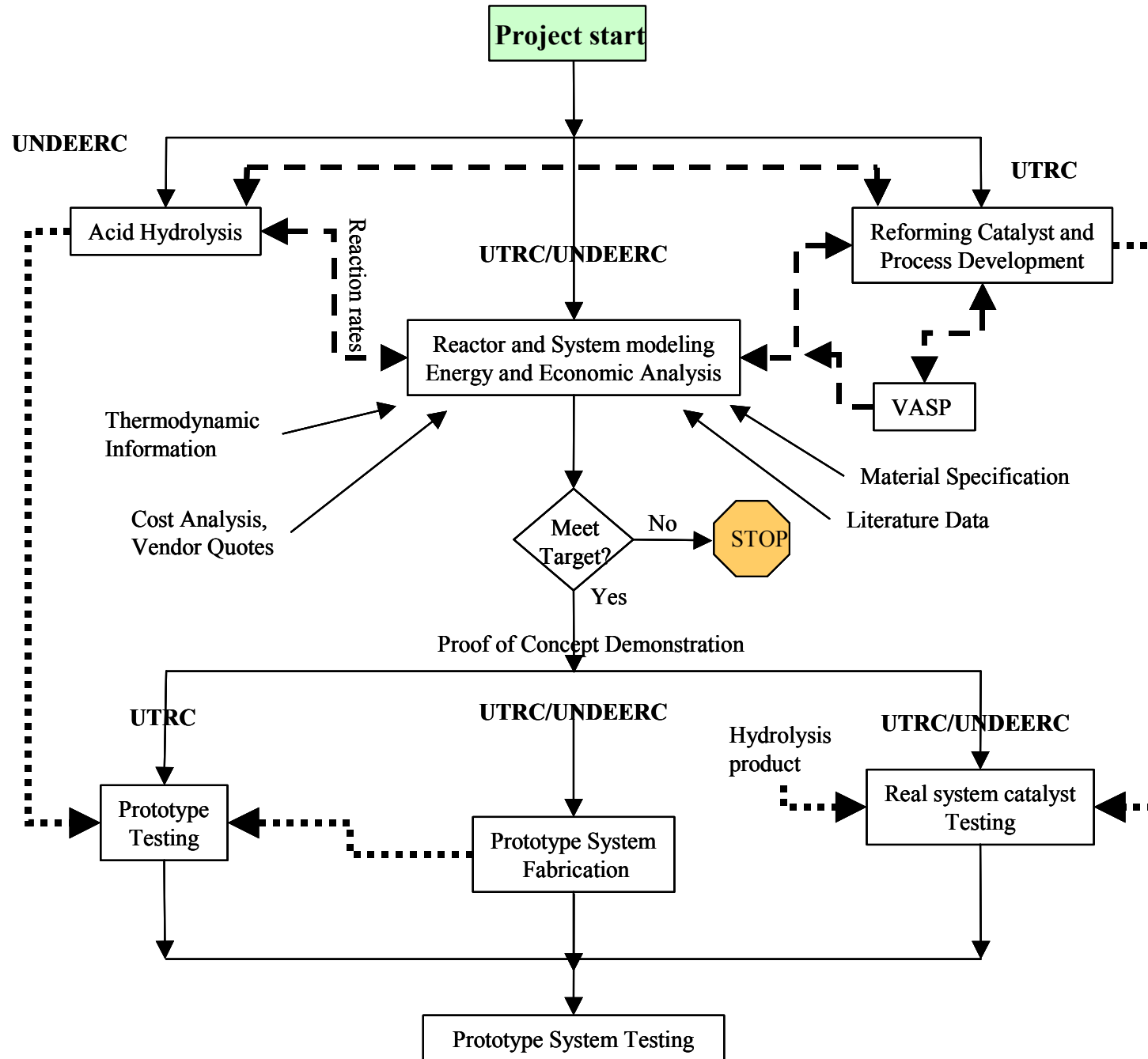
↓
High Selectivity Pt-MM rafts on engineered nano-structured oxide like Ti_[1-(x+y)]Dp_{1x}Dp_{2y}O₂

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

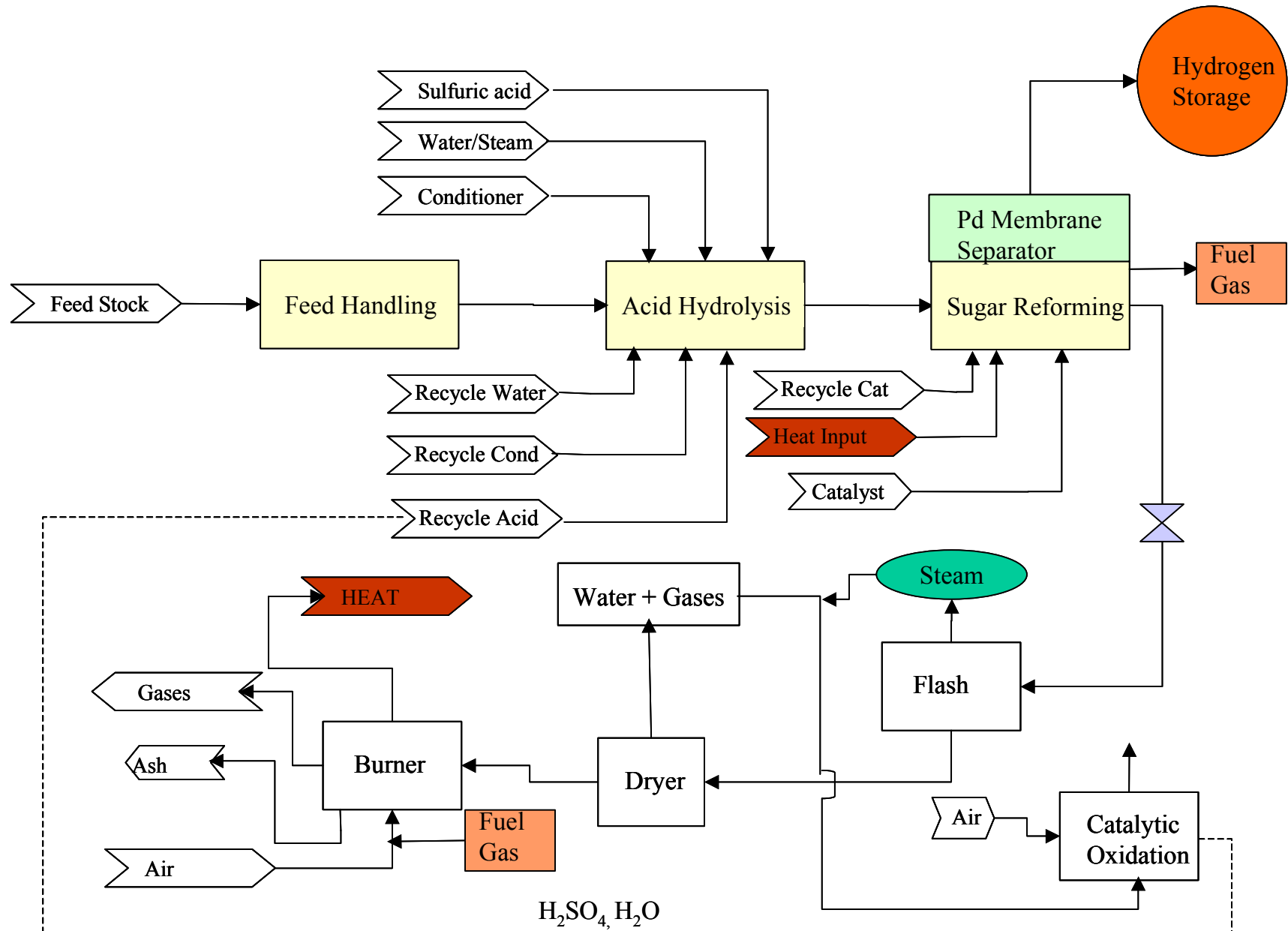
~9 g H₂ or Equivalent as fuel gas
~300 g Lignin and other fuel
~1 kg CO₂

Pt-Re/Ce_[1-(x+y)]Zr_xDp_yO₂ WGS Catalysts have high activity and very low CH₄ make

Original Project Plan Overview

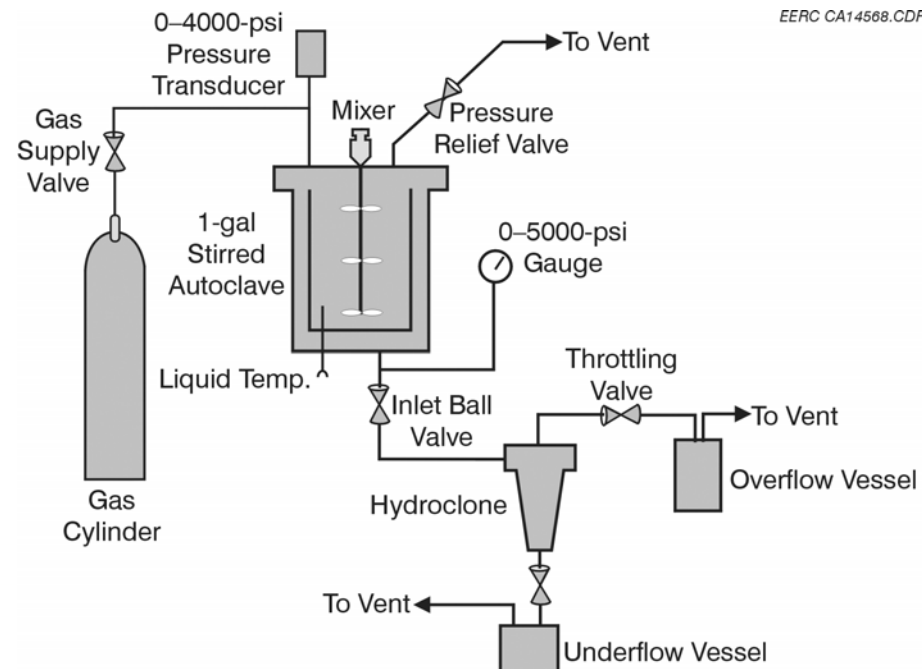


Approach: Initial Process Inputs and Outputs



Approach: Experimental Design to Optimize Hydrolysis

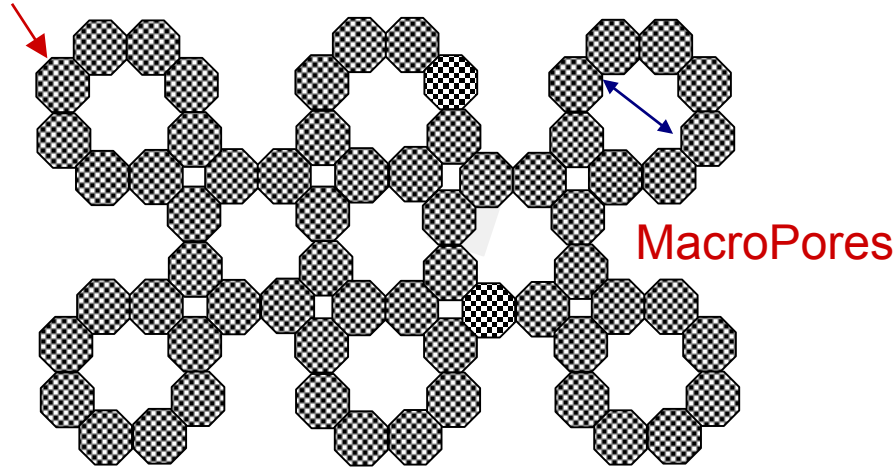
- Overall efficiency depends on optimizing hydrolysis energy / acid requirements
 - Lower acid concentration
 - + Less expensive alloys etc.
 - + Higher SA & activity reforming catalysts = smaller reforming reactors
 - + Less unnecessary chemical degradation = higher H₂ yield
 - Lower Temperature
 - + 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

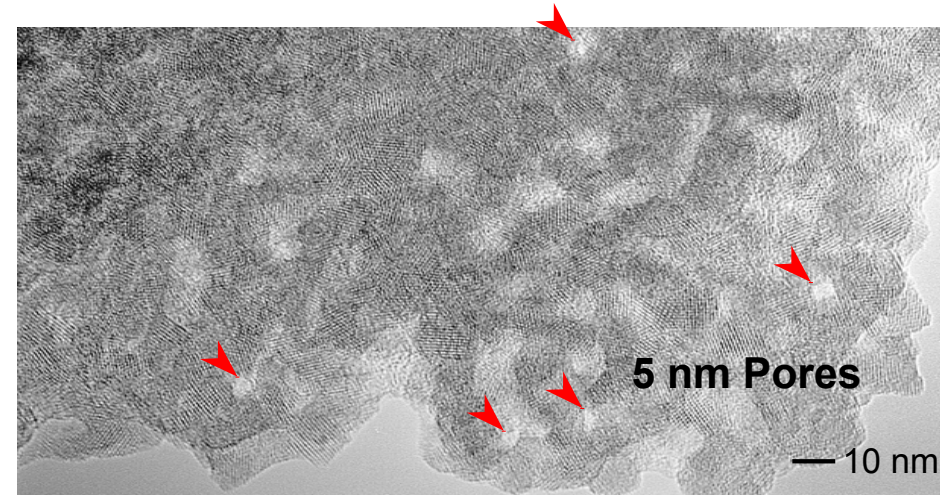
Design & synthesize active oxide structure to maximize accessible sites/vol.

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



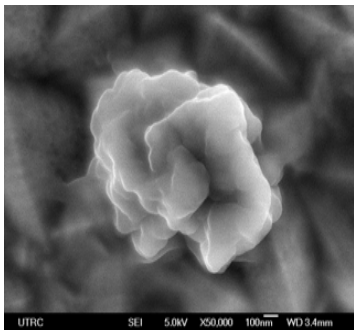
Conceptual Porous Metal-Oxide Framework Shown in 2D

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

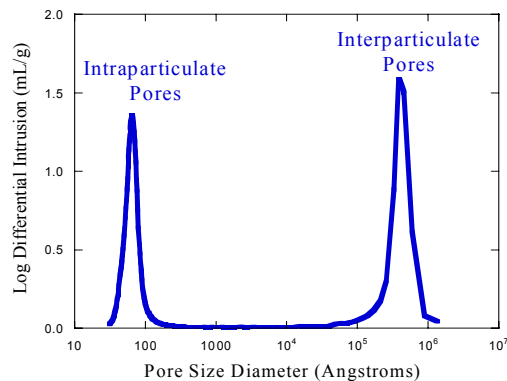


Conceptual Structure Realized

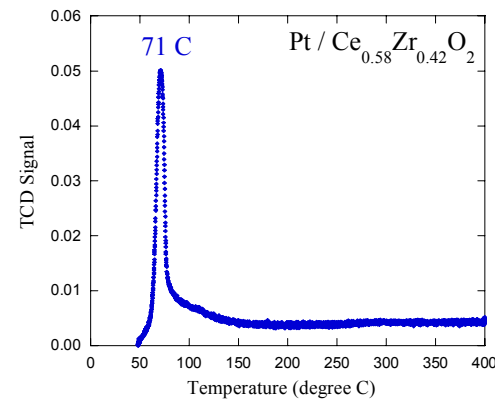
Fractal Morphology



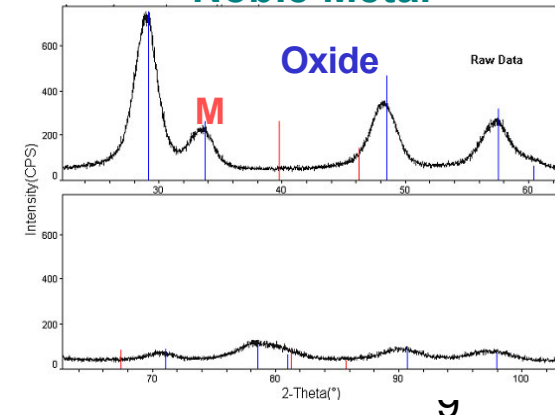
Large, Bimodal Pore Structure



Low Temperature Reducibility



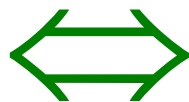
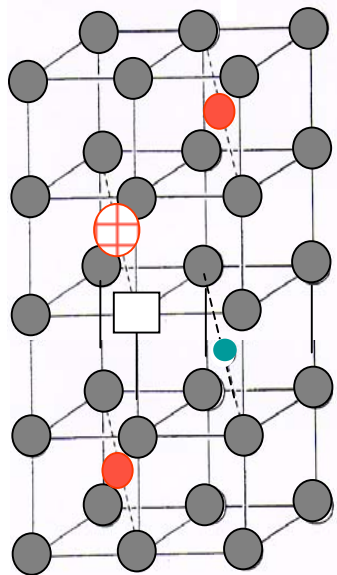
100% Dispersed Noble Metal



UTRC Catalyst Discovery Approach

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

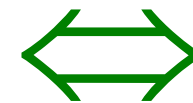
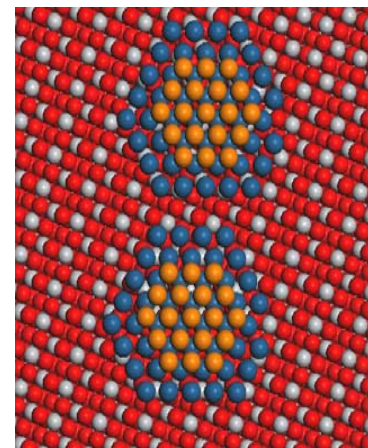
Conceptual Catalyst Design



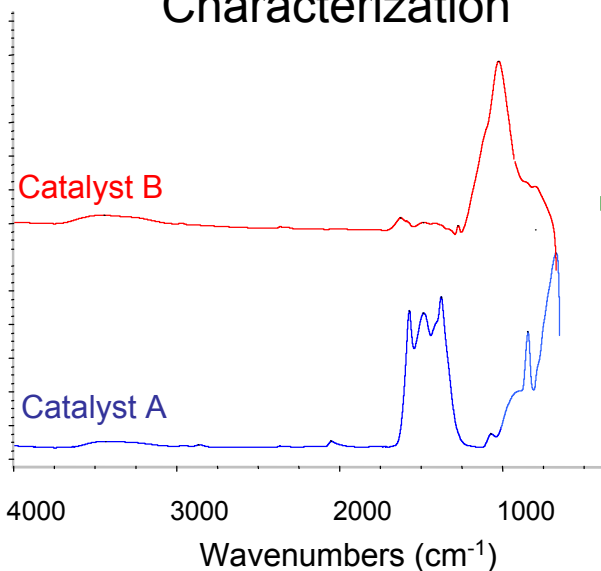
Catalyst Synthesis



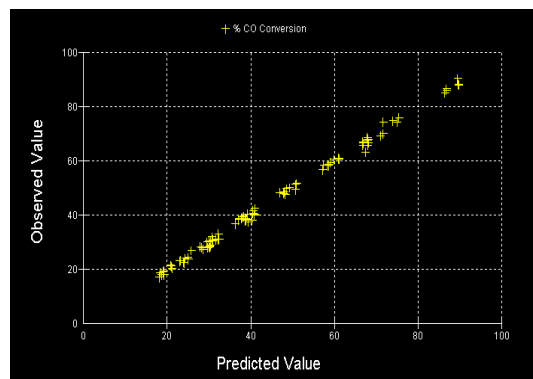
Quantum Mechanical Atomistic Modeling for advanced catalyst design



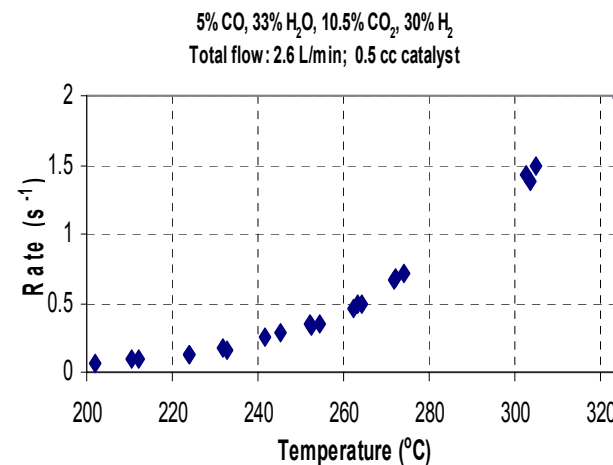
Characterization



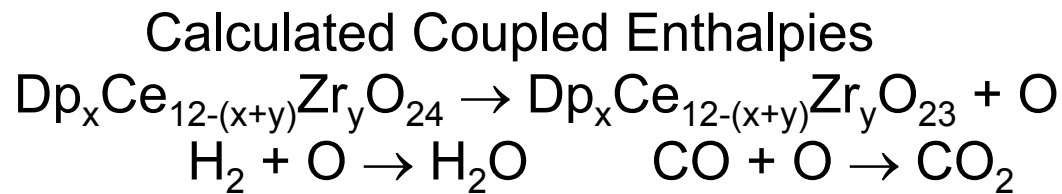
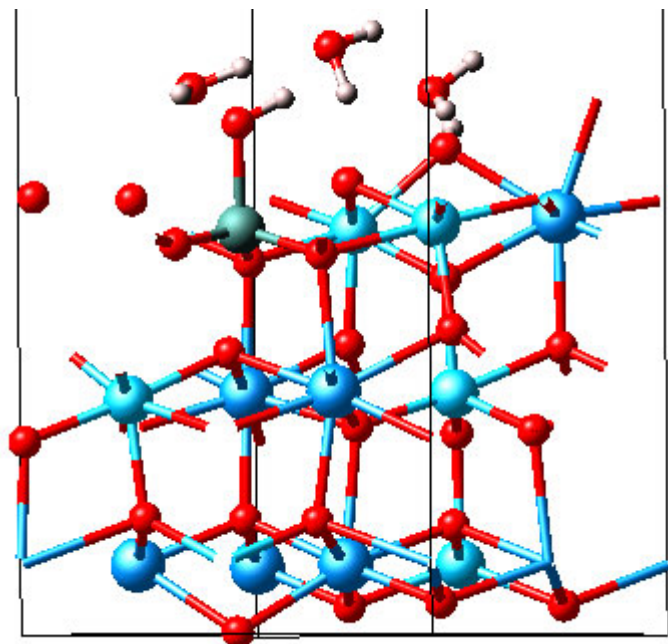
Kinetic Expressions Derived From Reaction Data



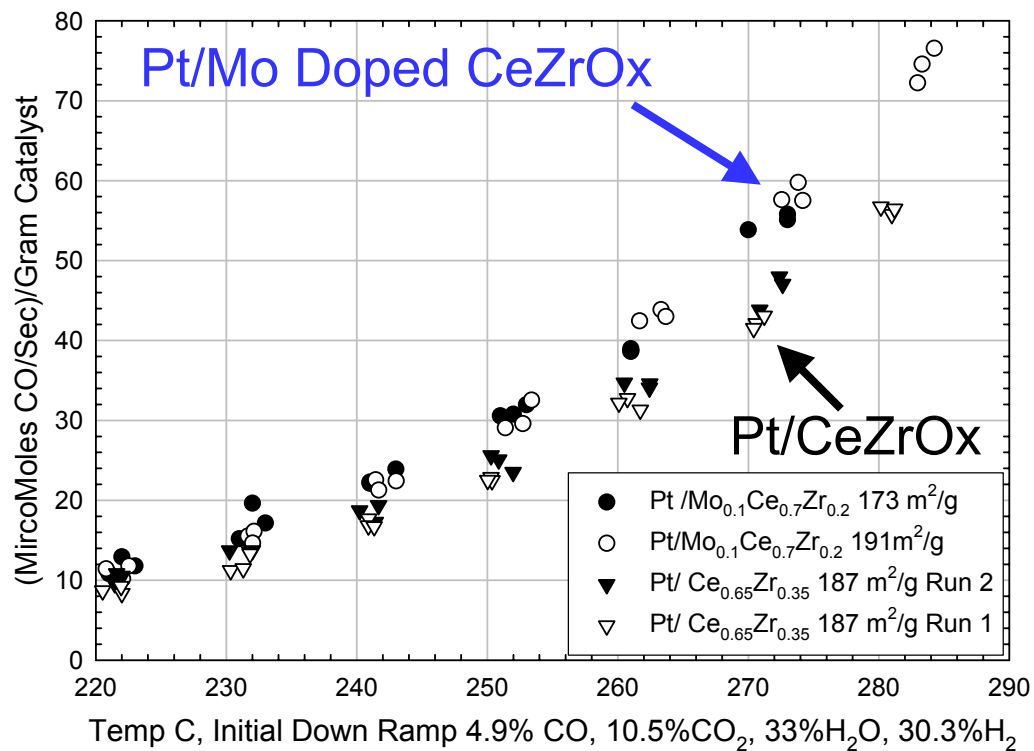
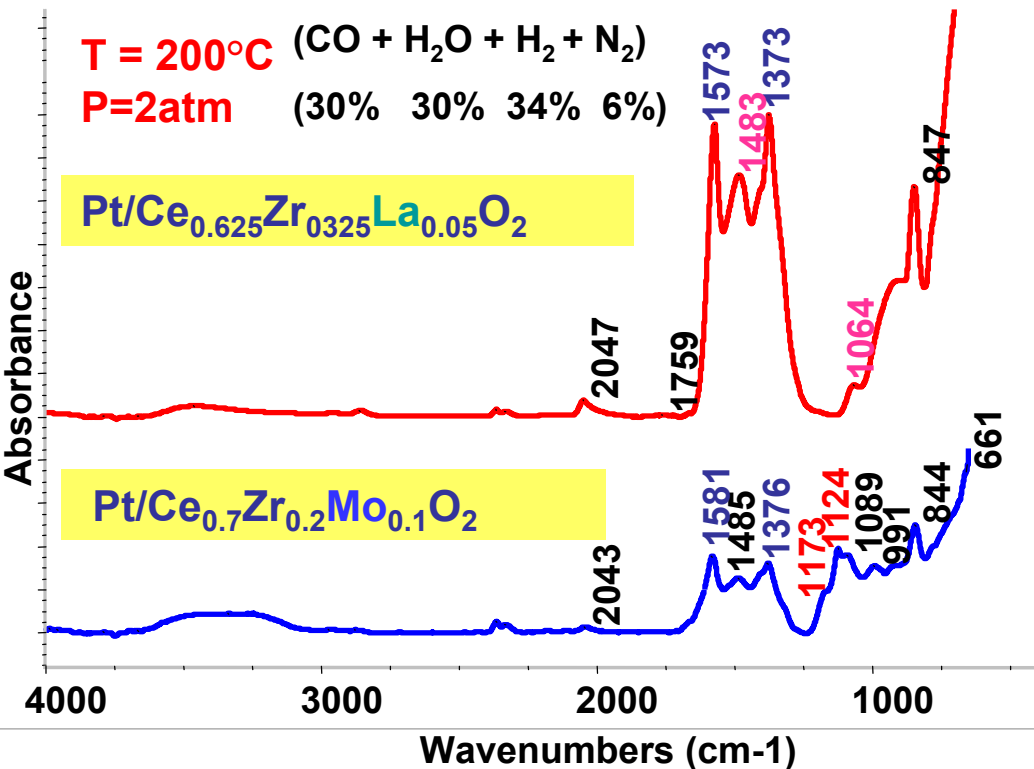
Superior Performance



VASP Modeling Insights Led To Better Catalysts



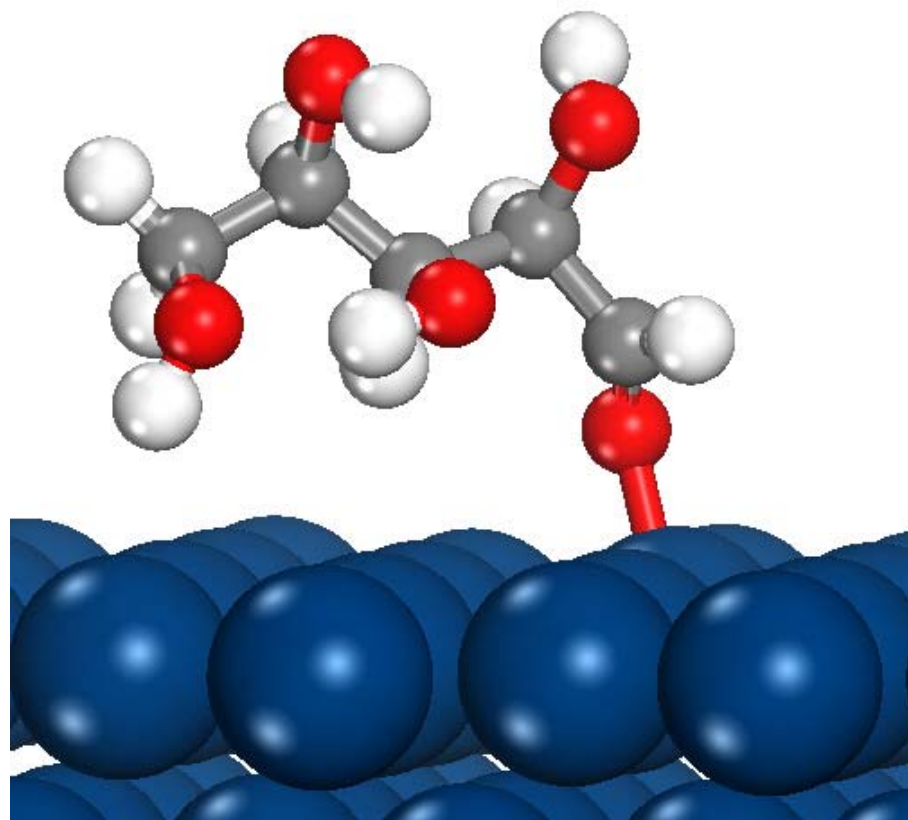
| Oxide Slab | $H_2 + O \rightarrow H_2O$ kJ/Mole | $CO + O \rightarrow CO_2$ kJ/Mole |
|----------------------------|------------------------------------|-----------------------------------|
| $Ce_{12}O_{24}$ | -154.5 | -222.4 |
| $Ce_7Zr_5O_{24}$ | -154.5 | -222.1 |
| Ta $Ce_6Zr_5O_{24}$ | 77.2 | 9.6 |
| Mo $Ce_6Zr_5O_{24}$ | 48.3 | -19.3 |



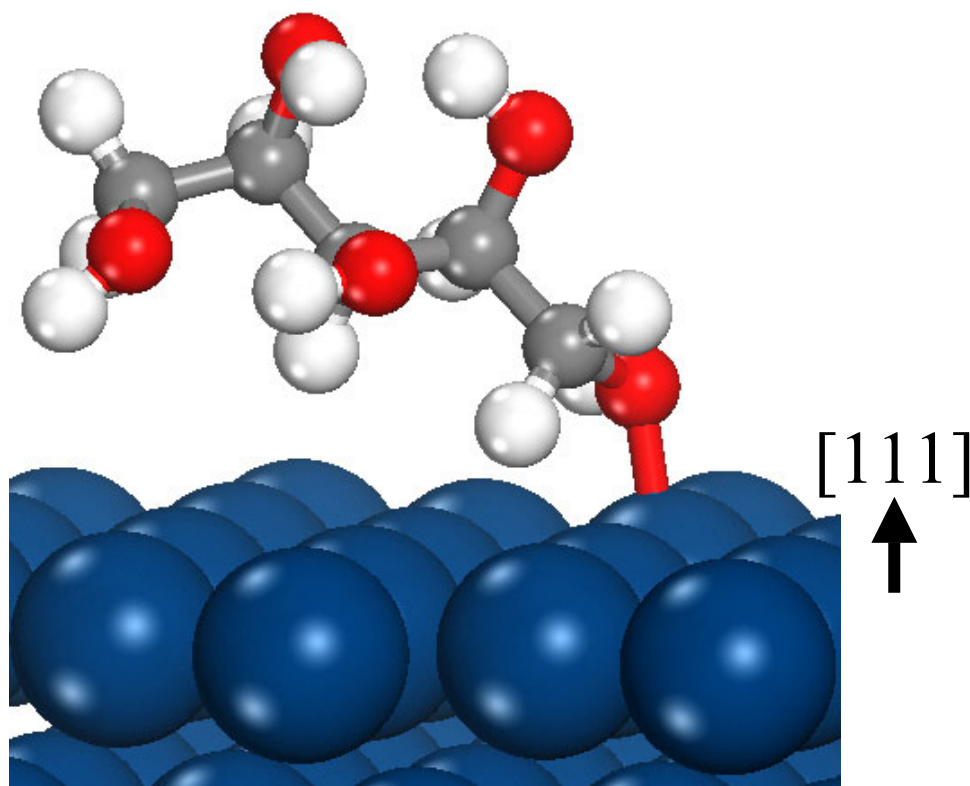
Higher Activity Catalyst w Similar Pt & SA

Xylose Adsorbs More Strongly Than Xylitol on Pt(111)

Aldehyde O forms stronger bond than terminal alcohol O



Xylose/Pt(111)



Xylitol/Pt(111)

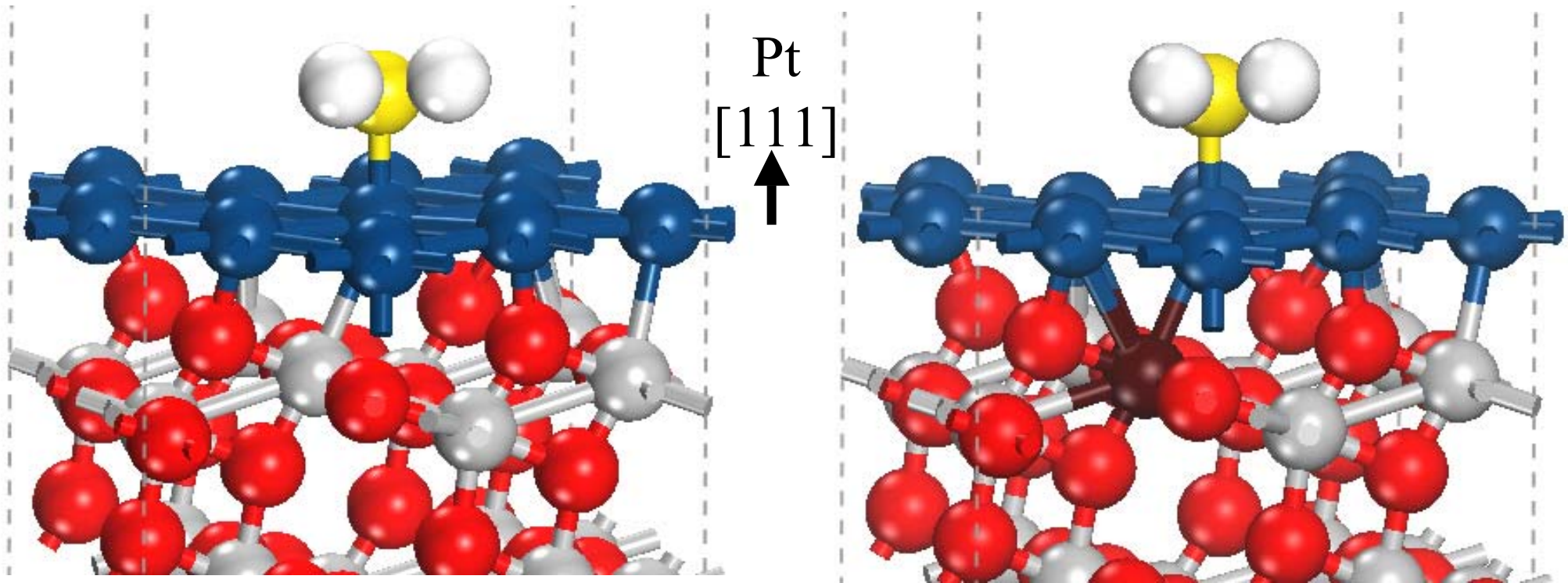
O C H Pt

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}/AnataseTiO₂(101)

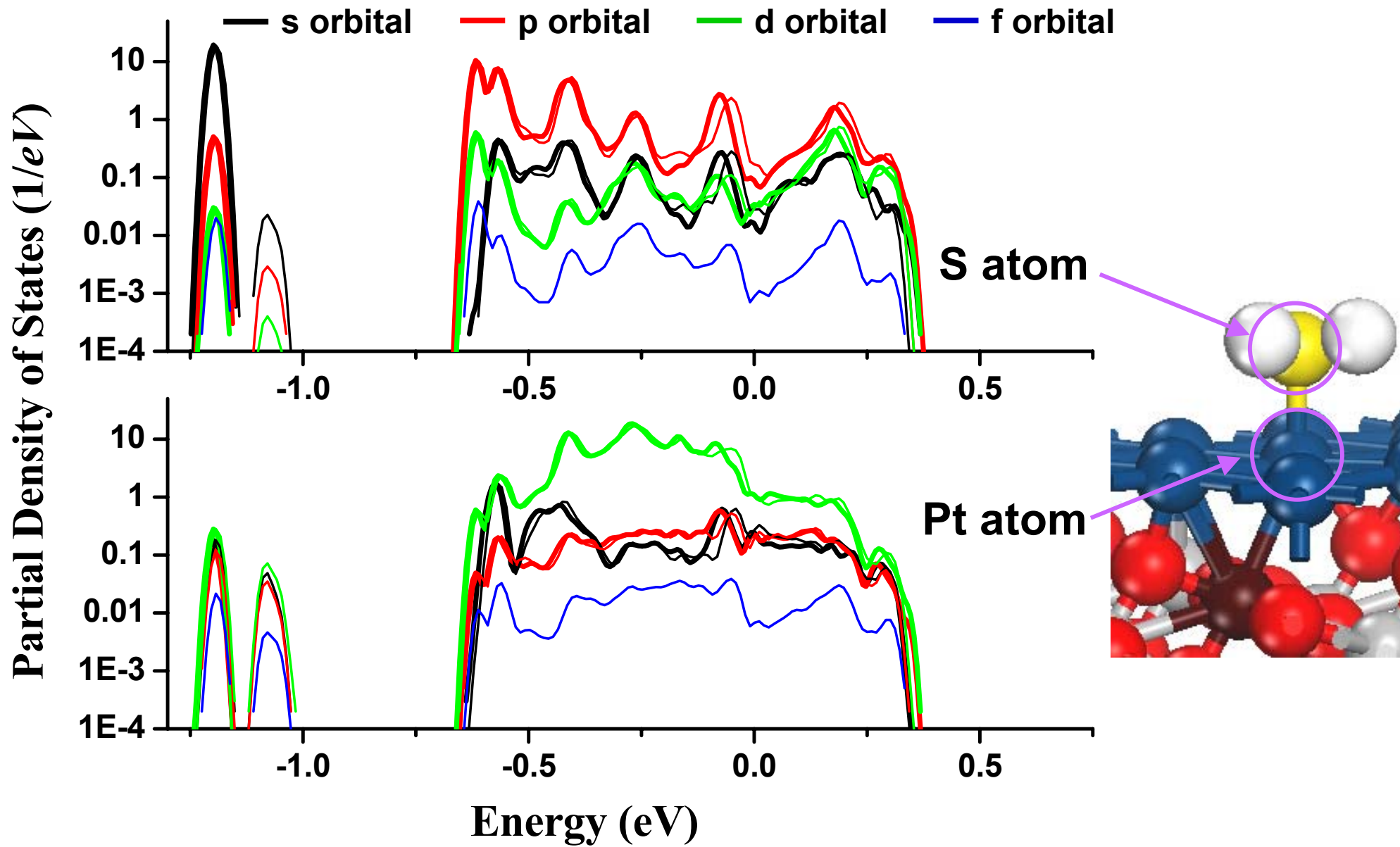
Binding Energy -106.53 kJ/mole

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

Binding Energy -89.50 kJ/mole

○ Ti H S Pt Ce

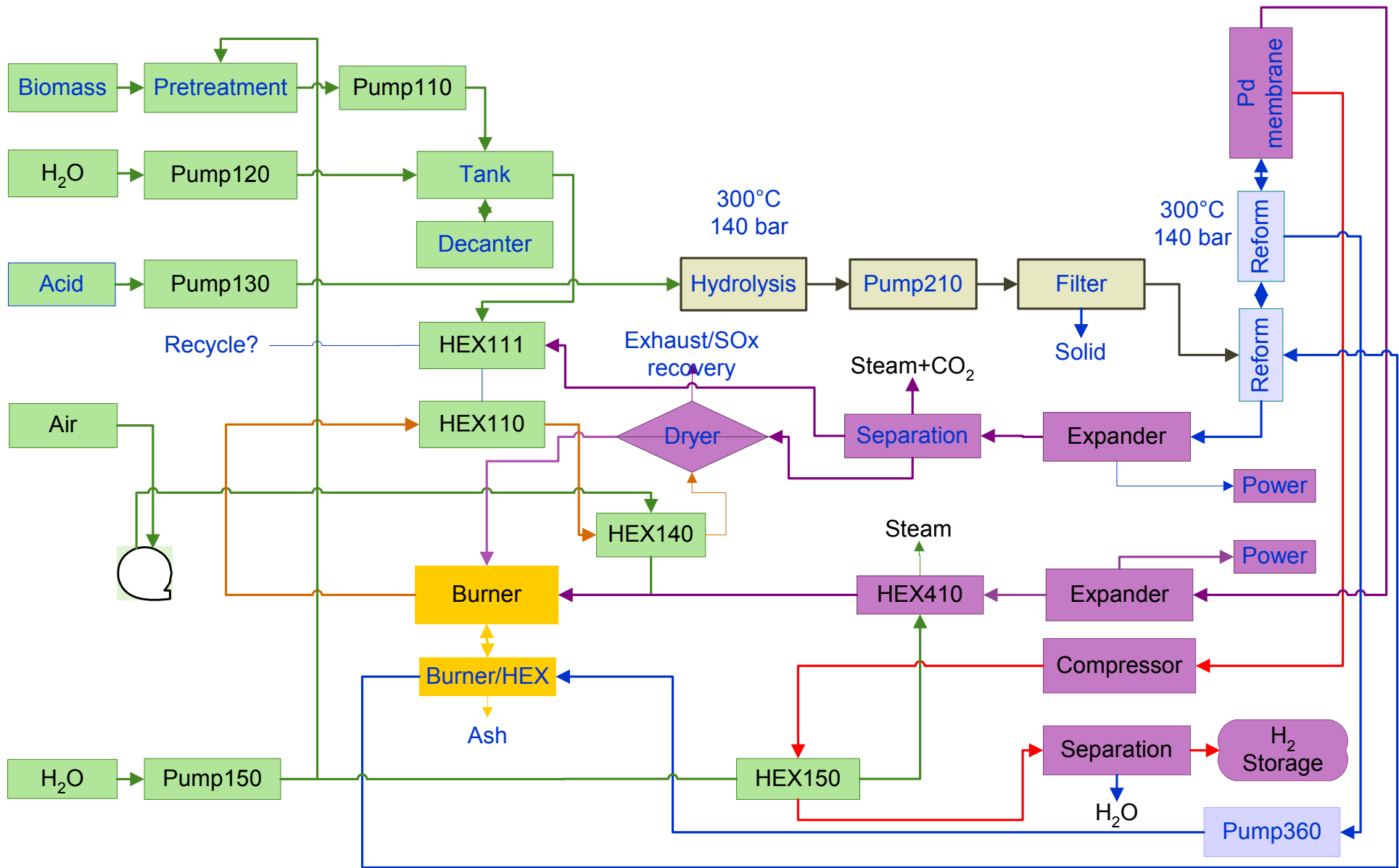
Oxide Dopant Shifts Pt & S DOS to Higher Energy



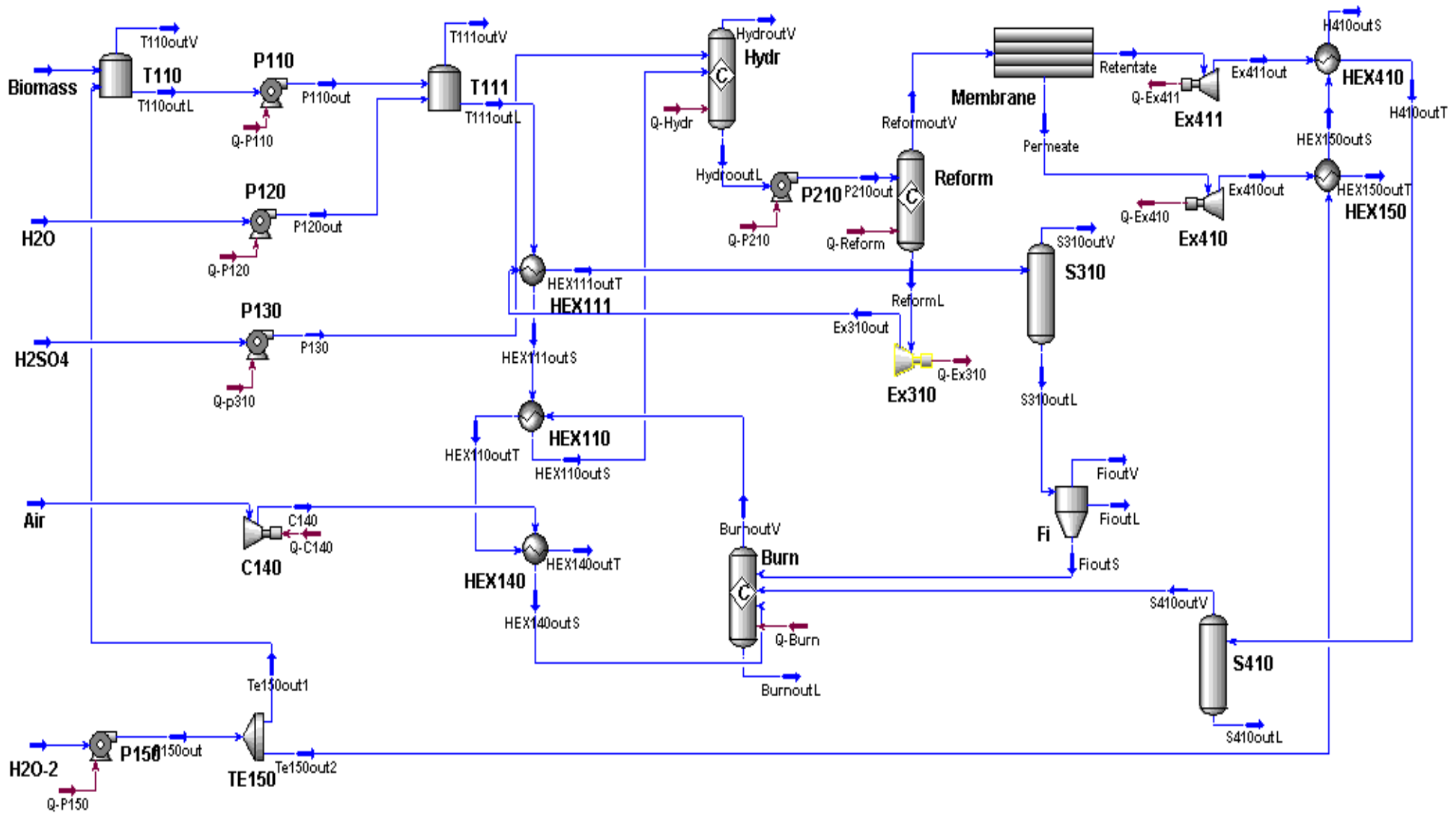
Thick line
Pt(111)_{1ML}/TiO₂(101)

Thin line
Pt(111)_{1ML}/4.2a% Ce in TiO₂(101)

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 H_2 /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 H_2
 - 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 H_2 /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