

# Solar High-Temperature Water Splitting Cycle with Quantum Boost

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May 16, 2012



PD027

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# Overview



- Start Date: 09/01/2007
- End Date: 8/31/2014
- Percent Complete: 45%



- Total project funding
  - DOE share: \$5.4M
  - Contractor share: \$1.4M
- Funding recvd. in FY11: \$750k
- Planned Funding for FY12: \$700k



- U. High-Temperature Thermochemical Technology
- V. High-Temperature Robust Materials
- W. Concentrated Solar Energy Capital Cost
- X. Coupling Concentrated Solar Energy & Thermochemical Cycles
- 2017 H<sub>2</sub> Production Target: \$3.00/kg
- 2017 Cycle Efficiency Target: >35%

### **Partners**

- SAIC (Lead)
   Solar System/Receiver
- UCSD/TChemE
   Thermochemical
   Process
- Electrosynthesis
   Salt Electrolysis







# **Overall Project Objectives** Relevance

- RD&D of the Viability of a <u>New</u> & Improved Sulfur Family Thermochemical Water-Splitting Cycle for large-scale hydrogen production using <u>solar</u> energy
- More Specifically, Overall Project Goals are to:

Evaluate SA water-splitting cycles that employ photocatalytic or electrolytic hydrogen evolution steps and perform lab testing to demonstrate feasibility of the chemistry

- Perform economic analyses of SA cycles as they evolve
- Select a cycle that has high potential for meeting the DOE 2017 cost target of \$3/kg hydrogen and efficiency goal of >35%
- Demonstrate technical feasibility of the selected SA cycle in bench-scale, closed-loop tests
- Demonstrate pre-commercial feasibility by testing and evaluation of a fully-integrated pilot-scale closed-cycle solar hydrogen production plant

## **Past Year Project Objectives**

- Complete Optimization of Electrolytic Oxidation Process
  - Test at higher temp (130°C) & higher pressure regimes
  - Incorporate better catalysts that lower the over-potential at the anode & allow operation over a range of current densities
  - Maintain pH at anode and cathode to avoid formation of reduced sulfur species
  - Perform 500 hour durability test & cell design improvement
- Complete Evaluation of High-Temperature K<sub>2</sub>SO<sub>4</sub> Sub-Cycle
  - Study phase equilibria of salt mixtures and identify favorable conditions for the process
  - evaluate salt mixtures to expand operational range
  - Optimization of process parameters and identification of side reactions and by-products
- Economic and Solar System Analysis
  - Continued Refinement of the Solar Field, Aspen Plus, and H2A Analyses to Incorporate Chemical Process Modifications
  - Evaluation of receiver materials, storage and control strategies



# **Impact on Program Barriers**

- <u>U. High-Temperature Thermochemical Technology</u> This project is developing a new thermochemical cycle intended to operate at lower temperatures and be more cost effective than other cycles.
- <u>V. High-Temperature Robust Materials</u> Materials able to operate in the high temperatures and caustic environment of the receivers and reactors are being developed.
- <u>W. Concentrated Solar Energy Capital Cost</u> –System costs are being reduced by optimizing the solar-chemical plant interface.
- X. Coupling Concentrated Solar Energy & Thermochemical Cycles

   The solar field and receivers are being evaluated and designed to
   provide appropriate amounts of energy at necessary temperature levels
   and with appropriate controls.

## Approach

# Approach

- Develop an Improved Sulfur-family Thermochemical Water-Splitting Cycle with Fewer Phenomenological Hurdles
- Conduct Laboratory Evaluation of all Processes within the Cycle
- Develop Comprehensive Aspen Plus<sup>®</sup> Flowsheet(s) of the Cycle and Modifications
- Perform H2A Analysis of the Flowsheeted Cycle(s) to Compare Design and Operational Approaches
- Design a Solar Collector Field and Receivers to Complement Chemical Process and Maximize Solar/System Efficiency
- Refine Process with Improvements Suggested by Lab Testing, Solar Configuration Studies, and H2A/Aspen Plus Analyses
- Perform Bench-Scale On-Sun Testing of Individual Chemical Processes, then Integrated Pilot-Scale Testing

# Milestones



#### 2011 Milestones

- Confirm thermodynamic viability of overall SA cycle with Aspen Plus<sup>®</sup> modeling <u>100% complete</u>
- Electrolytic H<sub>2</sub> production step: <0.8V, >50mA/cm<sup>2</sup> <u>100% complete</u>
- All-liquid O<sub>2</sub> production step validation <u>100% complete</u>
- Update solar concentrating system to match chemistry- <u>100% complete</u>

### 2012 Milestones

- Reduce energy consumption of electrolytic H<sub>2</sub> production step 20% by decreasing voltage <u>80% complete</u>
- Demonstrate the molten salt is liquid and will flow (low viscosity) so it is easily pumped-<u>100% complete</u>
- Demonstrate the NH<sub>3</sub> can be separated from the SO<sub>3</sub> by thermal decomposition thus avoiding potentially uneconomic gas separation processes - <u>85% complete</u>
- Develop a fully functioning and converging Aspen Plus modeling of the SA cycle <u>80% complete</u>
- Update solar concentrating system to match chemistry <u>50% complete</u>

#### 2012 Go/No-Go Decision Points

- Demonstrate the viability of the overall SA cycle, with no show stoppers
  - Electrolytic cell with current efficiency >90%, voltage 0.5-0.8V, current density 50-500mA/cm<sup>2</sup>, 500 + hour test
  - Thermodynamic and chemical plant analyses show the cycle can be closed and conversion yields for the O<sub>2</sub> production step are 90% + for all high temperature reactions



# **Electrolysis Progress**

- New catalysts and electrode materials have been tested over a range of current densities and temperatures (80 – 130°C). Promising materials include spinels M<sub>x</sub>N<sub>3-x</sub>O<sub>4</sub> (where M,N=Fe/Ni/Co), Pt/Co and vanadium oxides. To date, Pt/Co has shown the best catalytic performance.
- Quantitative hydrogen production with efficient sulfite oxidation has been verified with divided cells.
- Economic modeling has shown that the minimum annualized cost is at current densities <100 mA/cm<sup>2</sup>.



## **Electrolysis Progress**

Pressure reactor capable of 150 psi is being used to test electrolytic cells at 130°C.





Operation of the cell at 50 mA/cm<sup>2</sup> allows operation up to 90% conversion of sulfite with only a 60 mV penalty.

# **Oxygen Generation Half-Cycle**

- The all-fluid high-temperature sub-cycle using potassium sulfate-potassium pyrosulfate has been demonstrated by measuring the viscosity of the salts, which is low enough to be pumped
- Thermochemical reactor and Residual Gas Analysis (RGA) equipment is set up at UCSD and data have been collected and analyzed to show ammonia & sulfur trioxide gases can be evolved separately



1 to 5 g samples

# RGA Analysis of SaltsTech. Progressfor Oxygen Generation Half Cycle

 $(NH_4)_2SO_4 + K_2SO_4 + 4K_2S_2O_7 + Na_2SO_4 + 4Na_2S_2O_7$  (2 grams)



## **Molten Salts can be Easily Pumped**

 $K_2SO_4 + 4 K_2S_2O_7 + Na_2SO_4 + 4 Na_2S_2O_7$ 

K<sub>2</sub>SO<sub>4</sub> + 9 K<sub>2</sub>S<sub>2</sub>O<sub>7</sub> + Na<sub>2</sub>SO<sub>4</sub> + 9 Na<sub>2</sub>S<sub>2</sub>O<sub>7</sub>



- Viscosity of K<sub>2</sub>SO<sub>4</sub> + 4 K<sub>2</sub>S<sub>2</sub>O<sub>7</sub> + Na<sub>2</sub>SO<sub>4</sub> + 4 Na<sub>2</sub>S<sub>2</sub>O<sub>7</sub> ranged from 0.53 2.2 cP from 419 - 507°C
- Viscosity of K<sub>2</sub>SO<sub>4</sub> + 9 K<sub>2</sub>S<sub>2</sub>O<sub>7</sub> + Na<sub>2</sub>SO<sub>4</sub> + 9 Na<sub>2</sub>S<sub>2</sub>O<sub>7</sub> ranged from 0.29 2.3 cP from 393 510°C
- We measured melting points, densities, and viscosities

# Aspen Plus<sup>®</sup> Tech. Progress Process Model Schematic



	H2O	H2	O2	SULFITE	SULFATE	NH3-H2O	NH3H2O-2	K2SO7	K2SO4	SO3	SO2-O2
Г (°С)	25	100	155	155	100	400	65	400	835	835	1000
<sup>o</sup> (bar)	1	9	9	9	9	9	9	9	9	9	9
M (kg/hr)	1.04E+05	1.06E+04	1.01E+05	1.75E+06	1.74E+06	1.14E+06	1.14E+06	9.38E+07	9.32E+07	6.19E+05	6.19E+05
Phase	liquid	vapor	vapor	liquid	liquid	vapor	liquid	liquid	liquid	vapor	vapor

# Aspen Plus®Tech. ProgressProcess Model Improvements

- Low temperature reactor:
  - A design specification was implemented to regulate flow of salt stream to keep reactor at specified temperature (e.g., 400°C).
- Mid temperature reactor:
  - A design specification was implemented to input molten salt thermodynamics from Lindberg et al<sup>1</sup> paper.
  - This allows for more realistic values of K<sub>2</sub>S<sub>2</sub>O<sub>7</sub> decomposition at plant temperature (835°C) and pressure (9 bar).
- Electrolyzer:
  - A calculator block was placed to compute the power input needed at 0.8 V and an output concentration of a 2.5 M solution; values supplied by Electrosynthesis Company, Inc.
- Efficiency calculator:
  - A calculator block was placed in the flowsheet to compute the efficiency, as defined by DOE.
- Sensitivity analyses:
  - With the thermodynamics and physical parameter values, sensitivity analyses were conducted to obtain realistic values for plant operating conditions, i.e. pressures, temperatures, compositions.

<sup>&</sup>lt;sup>1</sup> Lindberg, D., R. Backman, and P. Chartrand. "Thermodynamic Evaluation and Optimization of the (Na2SO4+K2SO4+Na2S2O7+K2S2O7) System." *The Journal of Chemical Thermodynamics* 38.12 (2006): 1568-583. Web.

# **Alternate Flowsheet**



- Complete process at 9 bar
- Recovers heat of solution of SO<sub>2</sub> and NH<sub>3</sub>
- Hydrogen compressed to 300 psig
- Rankine power recovery system
- 24/7 production of hydrogen

- Excess electricity produced, or optionally partially used to Joule Boost SO<sub>3</sub> decomposition, simplifying solar system
- Overall efficiency ~40% (JB ~36%)
- Hydrogen only efficiency ~20% (JB ~22%)

# Solar Receiver Development

- SAIC cost-share supported two student interns in 2011
- Developed conceptual design of a receiver suitable for the SAIC 115 sq.m. dish concentrator system.
  - Approximately 100kW peak thermal input
  - Sufficient for production of ~0.8 kg/hr of hydrogen
- Design effort focused on the high-temperature SO<sub>3</sub> decomposition reactor
- Design based on Sandia bayonet reactor
- Reactor consists of two concentric Vycor<sup>®</sup> tubes with catalyst between them
  - Reactant flows up the annulus and products flow out the center tube
  - Heat transfer between inlet and outlet flows provides heat recuperation
  - Seals are at reduced temperature
  - Back reaction is minimized as products are cooled away from contact with catalyst

## **Conceptual High-Temperature Receiver**



Sandia Bayonet Reactor originally designed for H<sub>2</sub>SO<sub>4</sub> decomposition Proposed configuration for dish concentrator SO<sub>3</sub> decomposition

## Collaborations

# **Project Partners & Key Personnel**

#### Science Applications International Corp. (SAIC)- Prime (Industry)

- Key Personnel: Rob Taylor, Roger Davenport
- Project management & system integration
- Solar concentrator/receiver evaluation & design
- Thermal reactor optimization & design
- Economic evaluation & hydrogen production cost estimates

#### University of California, San Diego (UCSD) – Sub (Academia)

- Key Personnel: Jan Talbot, Richard Herz
- Laboratory testing & thermochemical cycle development
- Chemical plant analysis

#### Thermochemical Engineering Solutions (TChemE) - Sub (Industry)

- Key Personnel: Lloyd Brown
- Thermochemical cycle evaluation & analysis

#### Electrosynthesis Company, Inc. – Subcontractor (Industry)

- Key Personnel: David Genders, Peter Symons
- Electrolytic cell development & optimization

## Proposed Future Work

# FY12/FY13 Activities

- Complete Optimization of Electrolytic Oxidation Process
  - Continue development of better catalysts that lower the overpotential at the anode & allow operation at high current densities
  - Maintain pH at anode and cathode to avoid formation of reduced sulfur species
  - Perform 500 hour durability test & cell design improvement
  - Electrochemical cell design optimization and scale-up
- Complete Evaluation of High-Temperature K<sub>2</sub>SO<sub>4</sub> Sub-Cycle
  - Complete decomposition and gas separation studies
  - Develop bench-scale, pressurized molten salt flow system
- Economic and Solar System Analysis
  - Continued Refinement of the Solar Field, Aspen Plus<sup>i</sup>, and H2A Analyses to Incorporate Chemical Process Modifications
  - Evaluation of receiver materials, storage and control strategies

# Summary

#### • Electrolytic SA Cycle Step

- Improvements to electrocatalysts and high temperature operation have achieved cell voltages as low as 0.64 V at 50 mA/cm<sup>2</sup> and 0.85 V at 300 mA/cm<sup>2</sup>.
- Long term stability of these materials still needs to be demonstrated.
- Economic modeling has shown that the minimum annualized cost is at current densities <100 mA/cm<sup>2</sup>.

### All-Liquid High-Temperature Oxygen Generation

- Lab results prove the subcycle feasibility; salts have low viscosity and can be pumped
- Thermochemical reactor and Residual Gas Analysis (RGA) equipment used to show ammonia and sulfur trioxide can be evolved separately
- Optimization of process and integration into viable solar concentrating system and receivers needs to be further evaluated
- The Aspen Plus<sup>®</sup> SA process modeling has been significantly improved and is now a robust fully functioning process tool.
- Aspen Plus<sup>®</sup> and H2A Continue to be Used to Optimize and Trade-off SA Cycle Configurations
- Solar Configuration Focus is Central Receiver System with Molten Salt Storage to Allow 24/7 Operation
- Next-Phase Program is to Perform the SA Cycle Bench-Scale Tests and Prepare for On-sun Tests

# **Technical Back-Up Slides**

# Effect of pH on Anode Overpotential/ Cell Voltage

#### 160 2M NH4HSO3, 1M (NH4)2SO4, pH=4.9 140 2M (NH4)2SO3, pH=8.9 120 2M (NH4)2SO3 with 1 M NaOH, pH=12.5 Current density mA/cm<sup>2</sup> 100 80 60 40 20 0 -0.2 0 0.2 0.4 0.6 0.8 1 1.2 1.4 1.6 Potential vs NHE (V)

Potential for Sulfite Oxidation on Graphite in Various Solutions.

- Oxidation is kinetically slow. Potential needed to drive current is much higher than thermodynamic potential (more than 1 V overpotential).
- Anodic oxidation of sulfite and the cathodic hydrogen evolution reaction are pH dependent.

E° = 0.000 V

E° = -0.828 V

#### In acidic media

- anode reaction:  $SO_2 + 2H_2O \rightarrow SO_4^{2-} + 4H^+ + 2e^- E^\circ = 0.138 V$
- cathode reaction:  $2H^+ + 2e^- \rightarrow H_2$
- overall:  $SO_2 + 2H_2O \rightarrow SO_4^{2-} + 2H^+ + H_2$  E°cell = -0.138 V

#### In basic media

- anode reaction:  $SO_3^{2-} + 2OH^- \rightarrow SO_4^{2-} + H_2O + 2e^- E^0 = -0.936 V$
- cathode reaction:  $2H_2O + 2e^- \rightarrow H_2 + 2OH^-$
- overall:  $SO_3^{2-} + H_2O \rightarrow SO_4^{2-} + H_2$  E°cell = 0.108 V

# **Improvements in Electrocatalysis**



- All data shown is at 300 mA/cm<sup>2</sup>.
- Improvements in operating cell voltage can be attained by improving the catalyst and reducing the membrane resistance.
- Improvements in cell design also eliminate some of the resistive losses in the cell.

# Efficiency

- First Law efficiency  $\eta = -(\Delta H^{o}_{f[H^{2}O]} + E)/Q$
- Second Law efficiency  $\eta = -(\Delta G^{o}_{f[H^{2}O]} + E)/Q$
- DOE working definition  $\eta = -\Delta H^{o}_{f[H^{2}O(g)]}/(Q + E/\eta_{e})$

where

- $\eta$  = Plant efficiency
- $\Delta H^{o}_{f[H^{2}O]}$  = enthalpy of formation of water in standard state = -hhv
- $\Delta H^{o}_{f[H^{2}O(g)]}$  = enthalpy of formation of water vapor in ideal gas state = -lhv
- $\Delta G^{o}_{f[H_{2}O]}$  = Gibbs energy of formation of water in standard state
- Q = Total heat input to cycle
- E = Total electrical input to process or if negative the electrical output of the process
- $\eta_e$  = Efficiency by which consumed electricity is produced

Note: The First Law efficiency is the efficiency calculated from a heat balance around the total plant. The Second Law efficiency is the efficiency that, for a reversible process would be given by Carnot's Equation. The DOE working definition was designed to estimate the relative ranking of a cycle before the complete plant design was complete.

