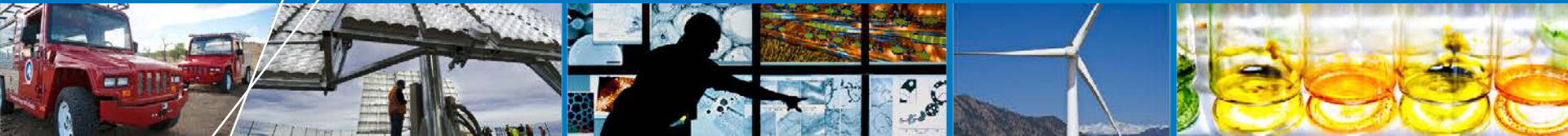


# Fermentation and Electrohydrogenic Approaches to Hydrogen Production



**2013 Annual Merit Review and Peer Evaluation  
Meeting; May 16, 2013**

**Pin-Ching Maness (PI; Presenter); National Renewable Energy  
Laboratory**

**Bruce Logan (Presenter); Penn State University**

**Project ID #: PD038**

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

# Overview



## Timeline

- Project start date: FY05
- Project not funded in FY06
- Project end date: 10/2013\*
- Percent complete: N/A

## Budget

- Total project funding: \$2,720K (includes \$387K subcontract)
- Funding received in FY12: \$350K
- Planned funding for FY13: \$410K

## Barriers

Barriers addressed

- H<sub>2</sub> molar yield (AX)
- Feedstock cost (AY)
- System engineering (AZ)

## Partners

- Dr. Bruce Logan  
Pennsylvania State University
- Drs. David Levin and Richard Sparling, University of Manitoba, Canada

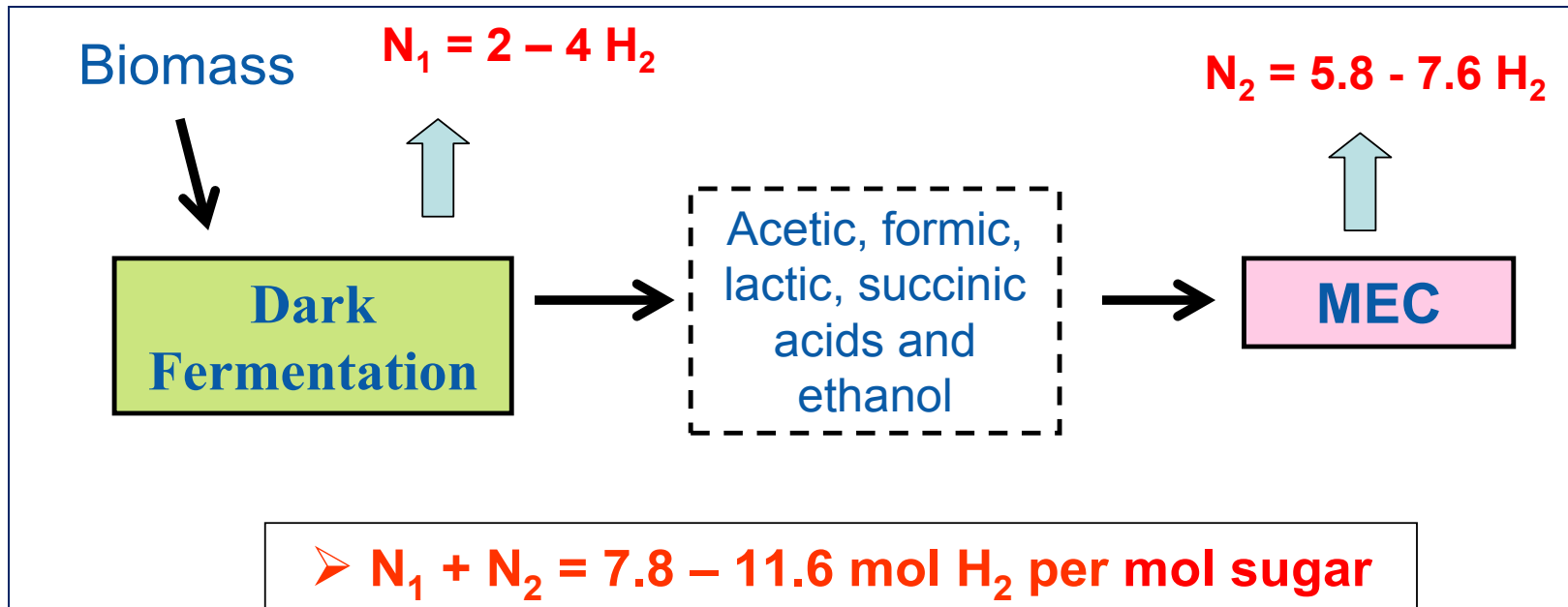
\*Project continuation and direction determined annually by DOE

# Objectives/Relevance

## Project Overview



**Objective:** Develop direct fermentation technologies to convert renewable lignocellulosic biomass resources to  $H_2$ .



Integrating fermentation with MEC dramatically improves combined  $H_2$  molar yield while reducing fermentation waste.

- MEC: microbial electrolysis cell;  $N_1$  and  $N_2$ :  $H_2$  molar yield (mol  $H_2$ /mol hexose).

# Relevance

**Relevance:** The project addresses directly two DOE barriers (feedstock cost and H<sub>2</sub> molar yield) aimed at improving the techno-economic feasibility of H<sub>2</sub> production via biomass fermentation.

- **Task 1. Bioreactor Performance:** Use cellulose (in lieu of sugars) and optimize parameters in sequencing fed-batch bioreactor to lower feedstock cost (Barrier AY).
- **Task 2. Metabolic Engineering:** Develop genetic tools to block competing pathways aimed at improving H<sub>2</sub> molar yield (Barrier AX).
- **Task 3. Electrochemically Assisted Microbial Fermentation:** Integrate microbial electrolysis cell (MEC) reactor with fermentation to improve H<sub>2</sub> molar yield (Barrier AX).

## Technical Targets

Characteristics	Units	2011 Status	2015 Target	2020 Target
Feedstock cost <sup>a</sup>	Cents/lb sugar	13.5	10	8
Yield of H <sub>2</sub> production from glucose	Mol H <sub>2</sub> /mol glucose	<b>3.2<sup>b</sup></b>	4	6
MEC production rate	L-H <sub>2</sub> /L-reactor-day	-	1	4

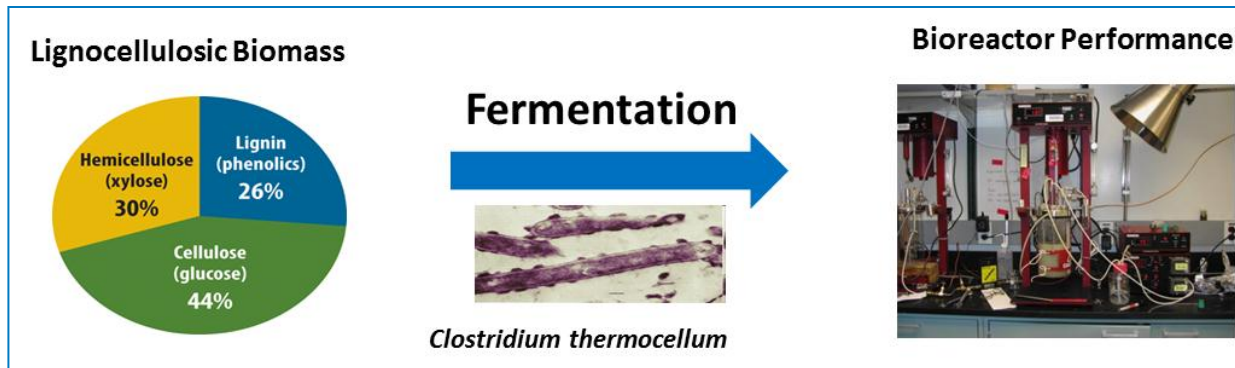
a. Status and target of the DOE Bioenergy Technology Office (formerly Office of the Biomass Program).

b. Low carbon substrate loading (1 g/L) led to high H<sub>2</sub> molar yield.

# Approach/Milestone

## Task 1: Bioreactor Performance

- Approach:** Optimize bioreactor in sequencing fed-batch mode by testing parameters such as cellulose loadings, hydraulic retention time, and liquid volume replacement and frequency using the cellulose-degrading bacterium *Clostridium thermocellum*.



	Milestone	Completion Date	Status
3.2.1-1	Design and build a feed tank capable of delivering a constant concentration of cellulose over the course of fermentation under variable volume conditions. A successful design will be defined by a standard deviation of less than 10%, evaluated by dry weight of cellulose exiting the vessel.	1/13	Completed
3.2.1-2	In order to demonstrate feasibility of H <sub>2</sub> production from biomass, improve H <sub>2</sub> molar yield by 20% compared to the current baseline (1.1 mol H <sub>2</sub> /mol hexose at 5 g/L/day cellulose feeding) by decreasing hydraulic retention time from 48 hr to 12-24 hr with more frequent liquid replacement using a constant cellulose loading of 5 g/L/day ( <i>CPS Agreement Milestone</i> )	5/13	On track

# Task 1 – Technical Accomplishments

## Scale Up “Automated” Sequencing Fed-Batch Bioreactor

- Automation features: Fermentation module executes the Feed, React, Settle, and Decant stages in a 5-liter bioreactor (2-liter working volume).
- New feed-tank design (impeller, high RPM) affords better stirring and reliable cellulose delivery.



Feed, React, Settle, Decant



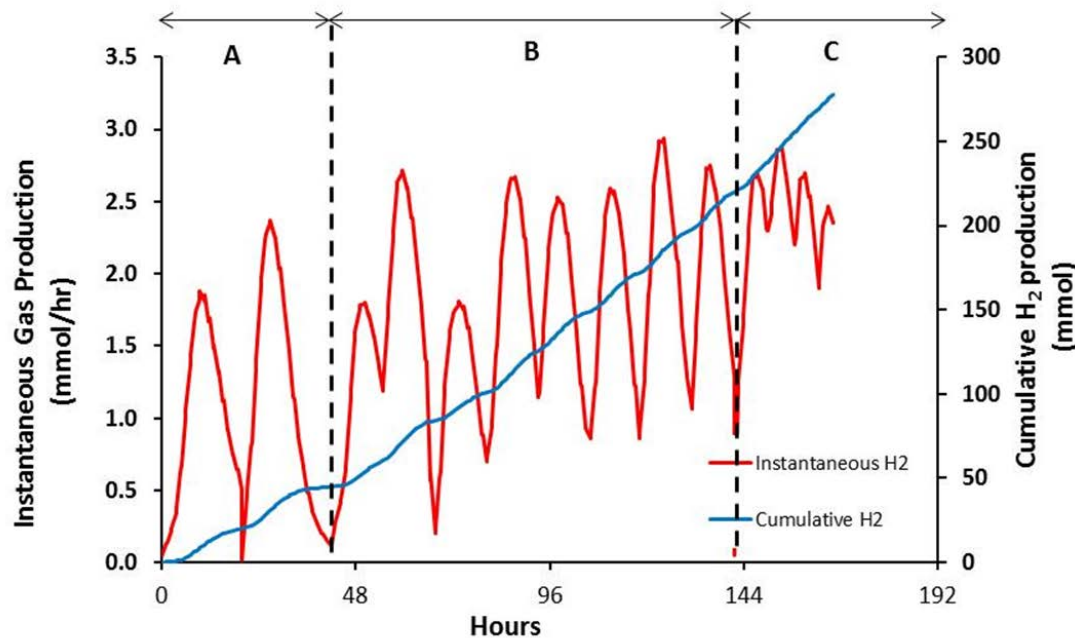
Cellulose settling retained acclimated microbes

***C. thermocellum* are attached to cellulose, hence allowing the bulk of the growth medium to be replenished while retaining the fully acclimated microbes.**

# Task 1 – Technical Accomplishments

## Increased Rate of H<sub>2</sub> Production by Two-fold

- Cellulose loading at 2.5 g/L/day with varying hydraulic retention time (HRT) and amount and frequency of liquid medium replacement.
- HRT: the length of time to replace the working volume (2 L) in a bioreactor.



	HRT (h)	Liquid Displacement (%)	Interval (h)	Rate of H <sub>2</sub> Production (ml/L/d)
A	48	50%	24	308 (1X)
B	24	50%	12	508 (1.7X)
C	24	25%	6	637 (2.1X)

- New feed-tank design delivered consistent amount of cellulose, resulting in <10% variations in H<sub>2</sub> production in each feeding cycle (completed Milestone 3.2.1-1).
- Increased rate of H<sub>2</sub> production by >2 fold with shorter HRT and more frequent liquid medium replacement.**

# Task 1 – Technical Accomplishments

## Increased Hydrogen Molar Yield by 53%

- Cellulose loading at 5 g/L/day with varying HRT and amount and frequency of liquid medium replacement.

HRT (h)	Liquid Displacement (%)	Interval (h)	H <sub>2</sub> Molar Yield (mol H <sub>2</sub> /mol hexose)	Rate of H <sub>2</sub> Production (ml/L/d)
48	75%	36	0.79 (1X)	764.6 (1x)
48	25%	12	1.09 (1.38X)	1057 (1.38X)
12	50%	6	<b>1.21 (1.53X)</b>	<b>1179 (1.54X)</b>

- Increased both H<sub>2</sub> molar yield (by 53%) and the rate of H<sub>2</sub> production (by 54%; by **3.8-fold** if compared to rate in the previous slide).
- At fixed HRT, more frequent liquid replacement (interval) is key to improvement.
- HRT of 12-24 h may be more optimal since shorter HRT increases water usage.

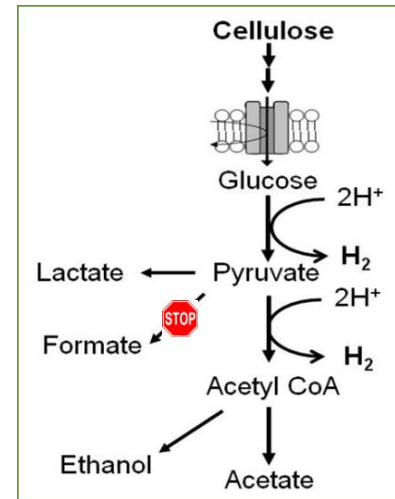
	Milestone	Completion Date	Status
<b>3.2.1-2</b>	Improve H <sub>2</sub> molar yield by 20% compared to the current baseline (1.1 mol H <sub>2</sub> /mol hexose at 5 g/L/day cellulose feeding) by decreasing HRT from 48 hr to 12-24 hr with more frequent liquid replacement ( <i>CPS Agreement Milestone</i> )	5/13	On Track



# Approach/Milestone

## Task 2 – Develop Genetic Methods for Metabolic Engineering

- **Approach:** Redirect metabolic pathways to improve H<sub>2</sub> molar yield via the development of genetic methods. The immediate goal is to delete the pyruvate-to-formate step.



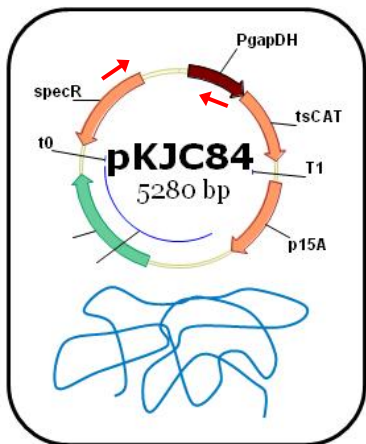
Katherine Chou

	Milestones	Completion Date	Status
<b>3.2.2 (FY12)</b>	Produce at least one plasmid for future use in knock-out strain production and other genetic manipulations, and demonstrate successful transformation into <i>C. thermocellum</i> via electroporation, with plasmid transfer confirmed by antibiotic selection and PCR (CPS Agreement Milestone 51458)	9/12	Completed
<b>3.2.2-1 (FY13)</b>	Produce a <i>C. thermocellum</i> mutant lacking its <i>hpt</i> gene to serve as the host for metabolic pathway mutant	4/13	Completed
<b>3.2.2-2 (FY13)</b>	Use $\Delta hpt$ mutant as the platform strain to further knockout the <b>pyruvate-to-formate</b> competing pathways. This approach aims at increasing H <sub>2</sub> molar yield by redirecting resources such as carbon and electrons towards H <sub>2</sub> production by reducing side-products production.	9/13	Completed

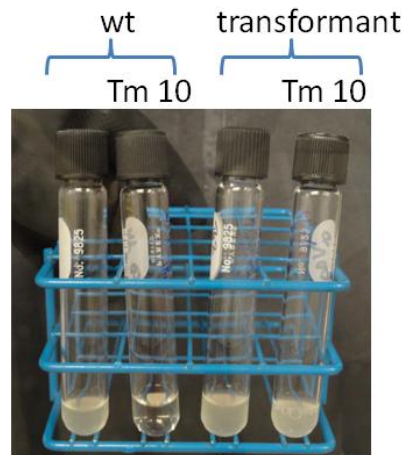
# Task 2 – Technical Accomplishments

## Generated a Stable Plasmid for Genetic Transformation

- NREL developed proprietary plasmid and stably transformed *C. thermocellum*.
  - We are one of the only two labs we know of that can transform *C. thermocellum*.
- PCR analysis verified presence of the plasmid (red arrow) which conferred growth in the antibiotic thiamphenicol (Tm, 10 µg/ml) (green arrow) whereas wild type (WT) cannot.

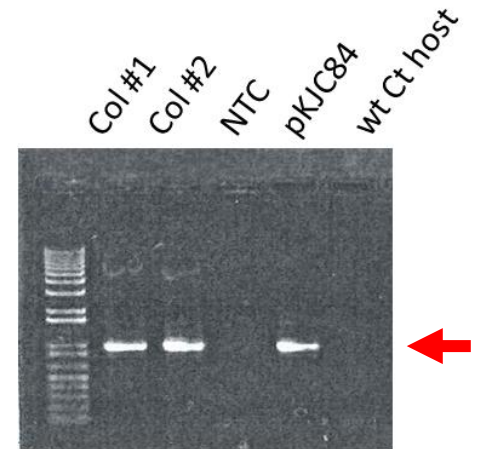


### Antibiotic Selection



10 subcultures

### PCR Analysis

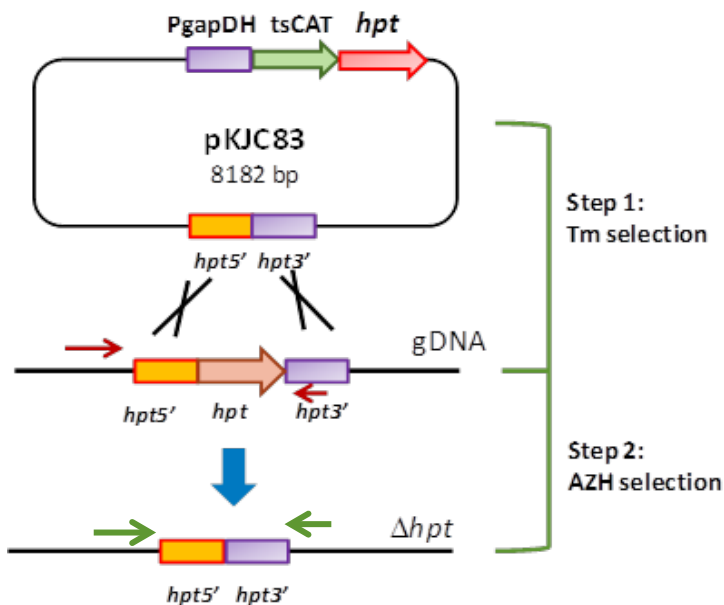


	Milestones	Completion Date	Status
3.2.2 (FY12)	Produce at least one plasmid for future use in knock-out strain production and other genetic manipulations, and demonstrate successful transformation into <i>C. thermocellum</i> via electroporation, with plasmid transfer confirmed by <b>antibiotic selection</b> and <b>PCR analysis</b> (CPS Agreement Milestone)	9/12	Completed

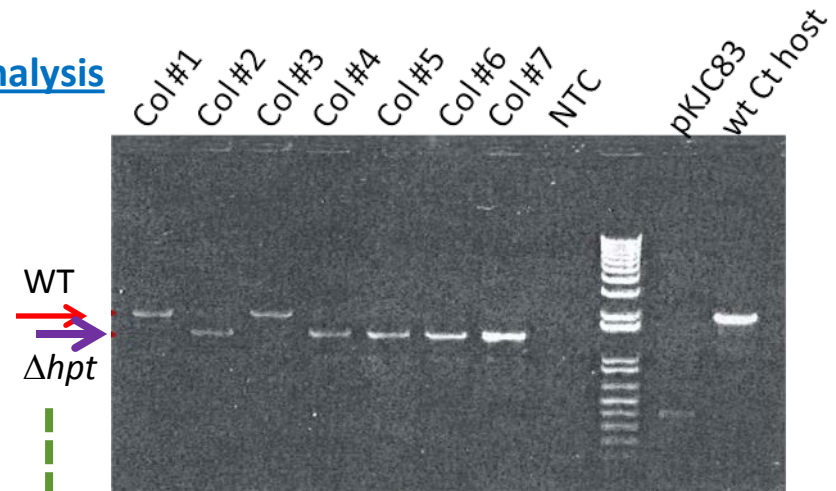
# Task 2 – Technical Accomplishments

## Generated Five *hpt* Knockout Mutants using Proprietary pKJC83 Plasmid

- *hpt* encodes hypoxanthine phosphoribosyl transferase, which converts AZH (8-azahypoxanthine) to a toxic product that kills the wild-type cells.
- $\Delta hpt$  mutants survive in AZH, which forms the basis for a  $\Delta hpt$  counter-selection strategy.
- **Five  $\Delta hpt$  mutants were verified by PCR (purple arrow, right figure) and DNA sequencing.**
- $\Delta hpt$  mutant will serve as the host to generate metabolic pathway mutants.



### PCR Analysis



Only  $\Delta hpt$  mutants will yield this PCR product (lower band; lanes 2, 4-7).

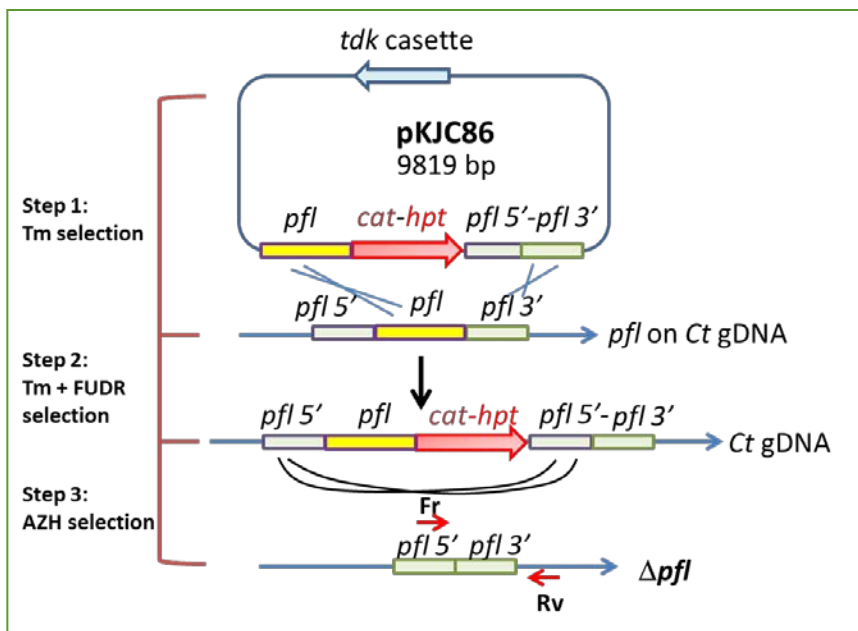
	Milestone	Completion Date	Status
<b>3.2.2-1 (FY13)</b>	Produce a <i>C. thermocellum</i> mutant lacking its <i>hpt</i> gene to serve as the host for metabolic pathway mutant	4/13	Completed

# Task 2 – Technical Accomplishments

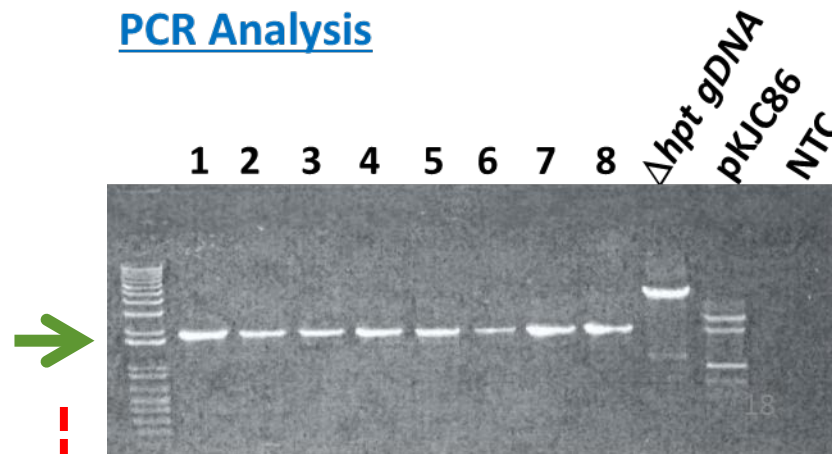
## Generated Eight Pyruvate-to-formate (PFL) Pathway Mutants

Generated **eight**  $\Delta pfl$  mutants, verified by PCR using  $\Delta hpt$  mutant as the host (green arrow in the right figure).

- pKJC86 plasmid design included both *hpt* and *tdk* (thymidine kinase) genes. *tdk* leads to toxicity in fluoro-deoxyuracil (FUDR) – a double selection strategy (combining with AZH).



### PCR Analysis



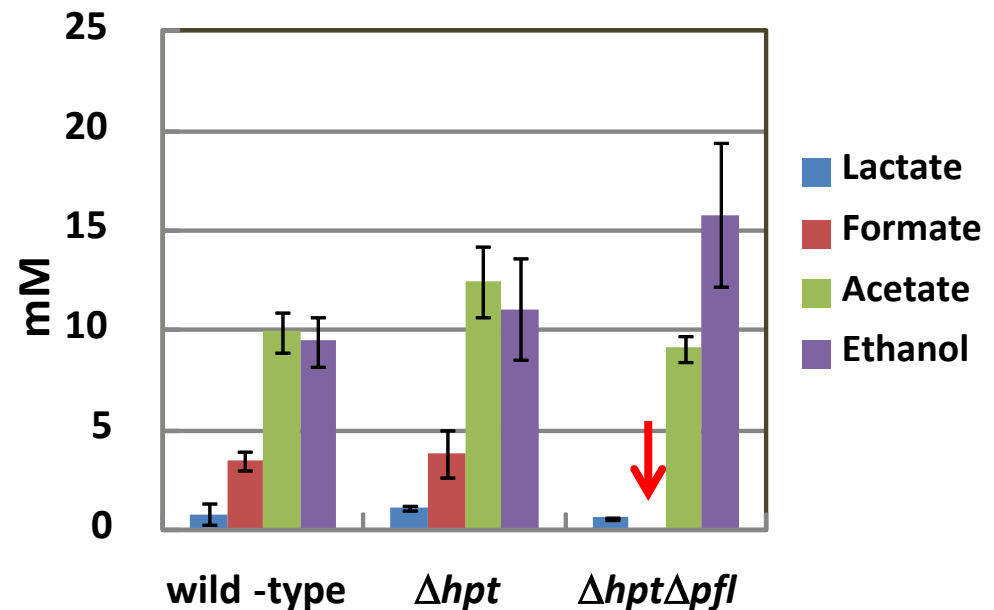
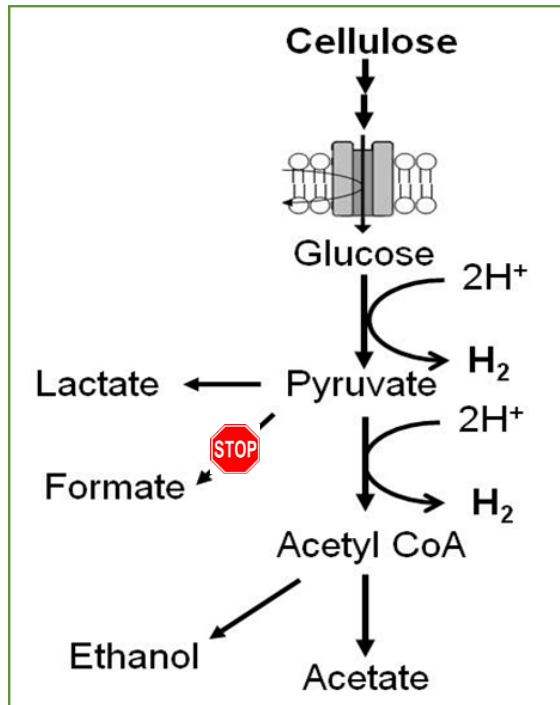
**Only  $\Delta pfl$  mutants will yield this PCR product (lanes 1-8).**

	Milestone	Completion Date	Status
<b>3.2.2-2 (FY13)</b>	Use $\Delta hpt$ mutant as the platform strain to further knockout the pyruvate-to-formate competing pathways aims at increasing H <sub>2</sub> molar yield by redirecting carbon and electrons toward H <sub>2</sub> production.	9/13	<b>Completed</b>

# Task 2 – Technical Accomplishments

## Confirmed $\Delta Pfl$ mutant by Metabolite Analysis

- $\Delta pfl$  mutant did not produce formate as expected based on deletion of *pfl* gene encoding the pyruvate-to-formate pathway.
- $\Delta pfl$  mutant produced ~ **60% more ethanol** than the parent strains.
- Knocking out the ethanol pathway may increase  $H_2$  molar yield in future experiments since both pathways compete for the reductant NAD(P)H.



- Work is underway to determine  $H_2$  production from various strains in bioreactors.

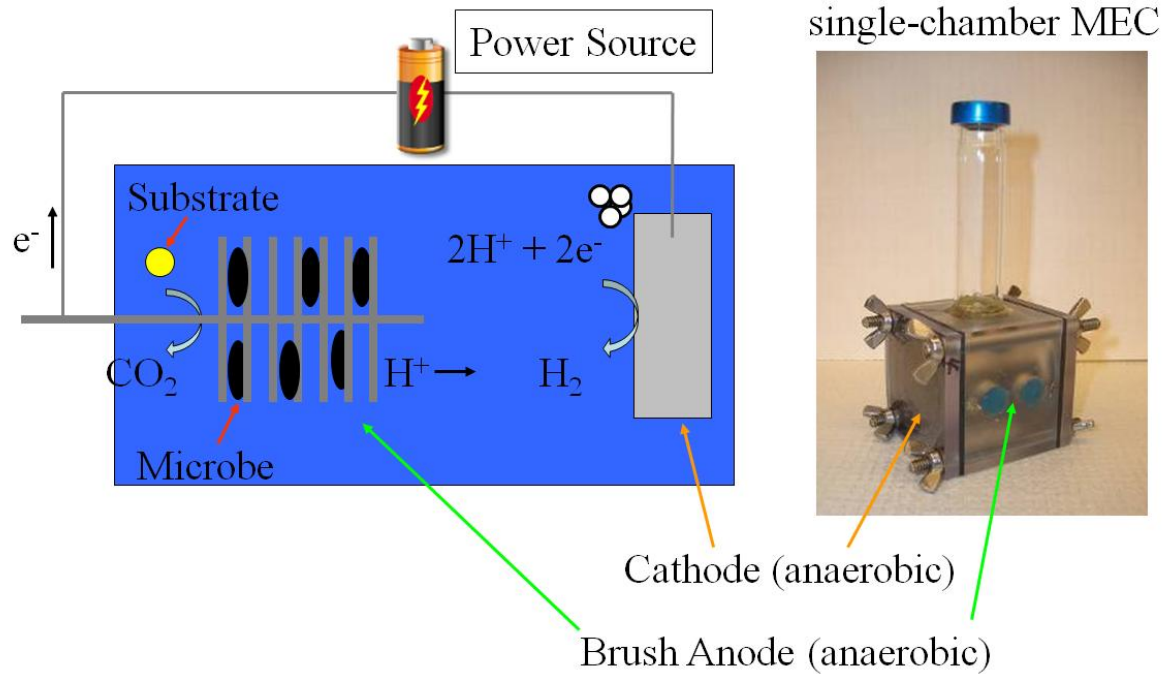
# Approach/Milestone



## Task 3 – Electrochemically Assisted Microbial Fermentation

-Produce H<sub>2</sub>

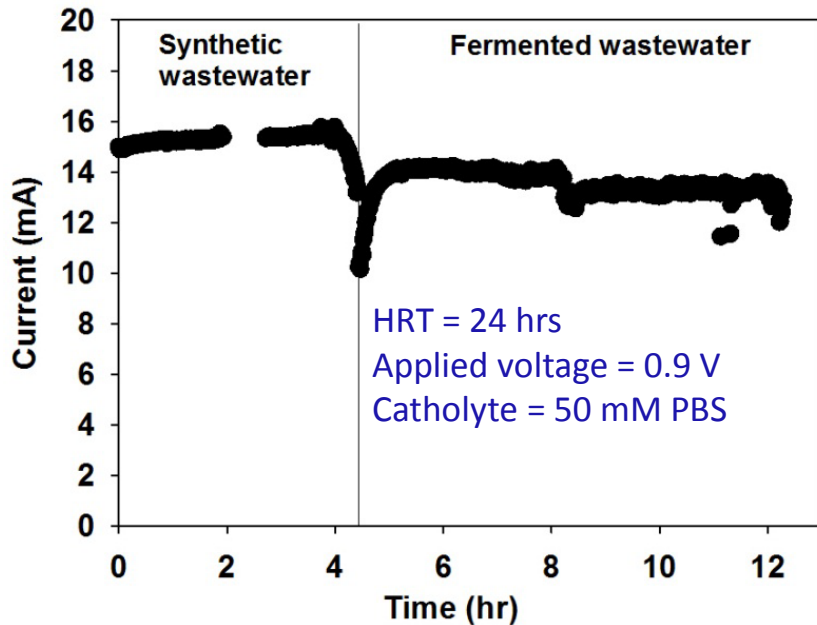
-Non-spontaneous (+ΔG) reaction



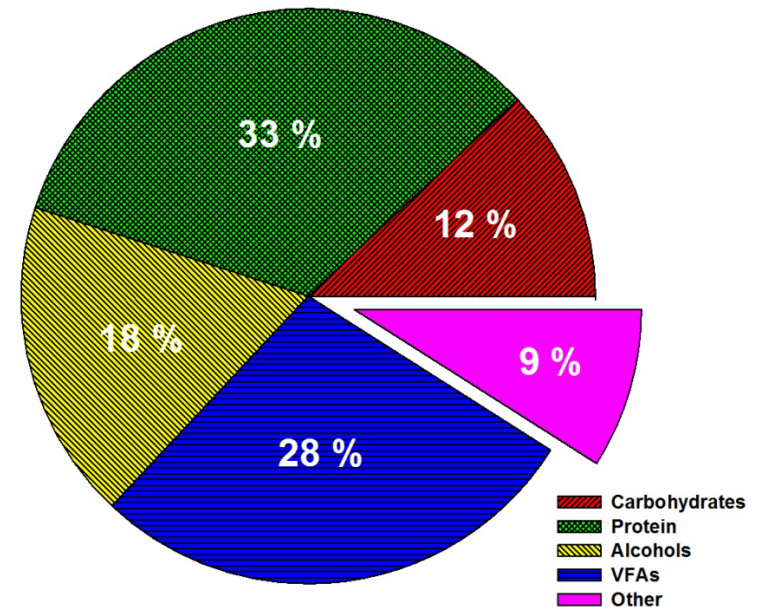
	Milestones	Completion Date	Status
<b>3.2.3 (FY12)</b>	Correlate removal of the subcomponents of the NREL fermentation effluent with current density and H <sub>2</sub> production	9/12	Completed
<b>3.2.3-1 (FY13)</b>	Build prototype MREC reactor and evaluate H <sub>2</sub> production using NREL fermentation effluent with zero electrical grid energy; demonstrate production rate of ≥0.5 L H <sub>2</sub> L <sup>-1</sup> reactor day <sup>-1</sup> over 3 hydraulic retention times with continuous flow	9/13	On Track

# Task 3.1 – Technical Accomplishments

## Hydrogen Generation from Fermentation Wastewater



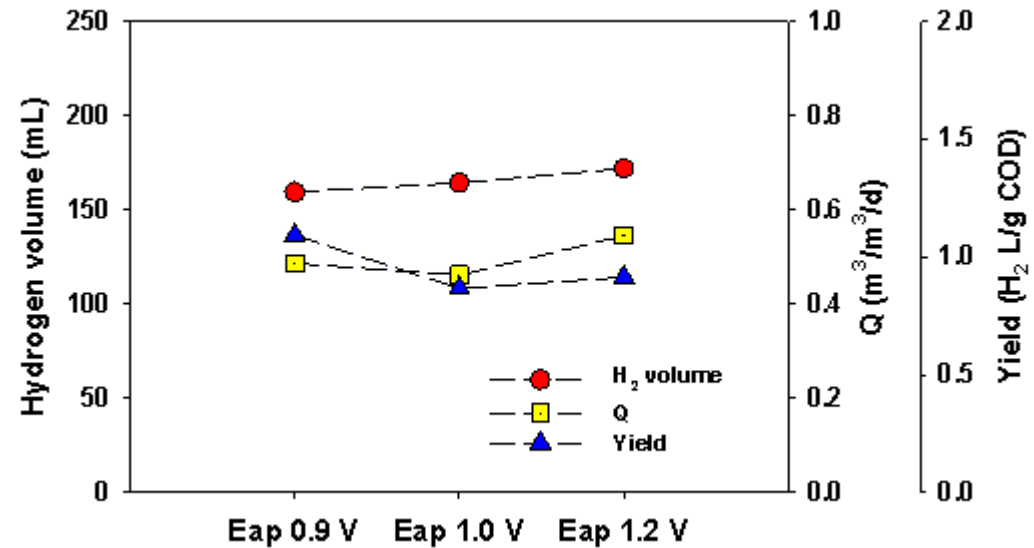
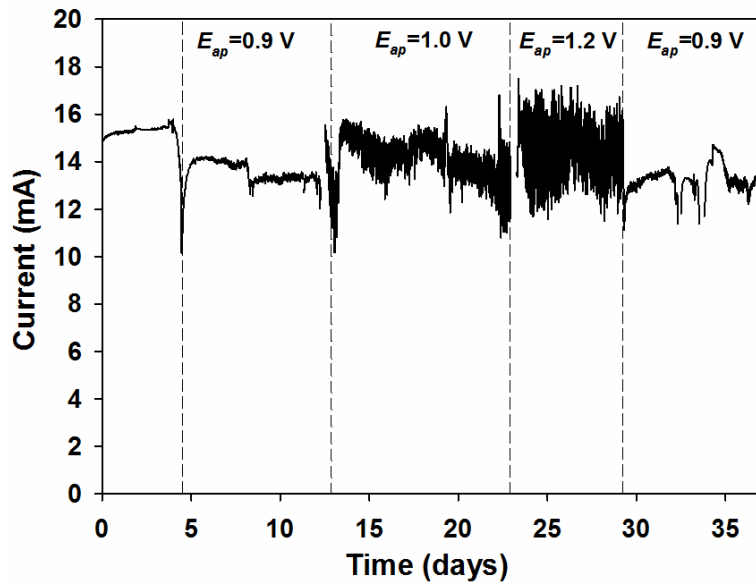
### Fermentation Effluent Composition



- Lower performance with fermentation wastewater (ww) due to substrate complexity
- **Current:** Synthetic ww = 51 A/m<sup>3</sup>; Fermented ww = 44 A/m<sup>3</sup> (no protein in synthetic ww).
- **COD (chemical oxygen demand) removal:** Synthetic ww = 87%; Fermented ww = 73%.
- **Gas volume:** Synthetic = 183 mL; Fermented = 159 mL
- Gas production rates: Synthetic, 0.6 L-H<sub>2</sub> L<sup>-1</sup> d<sup>-1</sup> ; Fermented, 0.5 L-H<sub>2</sub> L<sup>-1</sup> d<sup>-1</sup>.
- **Protein removal (48%) was lower than alcohols and VFAs (>90%) and carbohydrate (89 %).**

# Task 3.2 & 3.3 – Technical Accomplishments

## Different Applied Voltages, Gas Production Rates

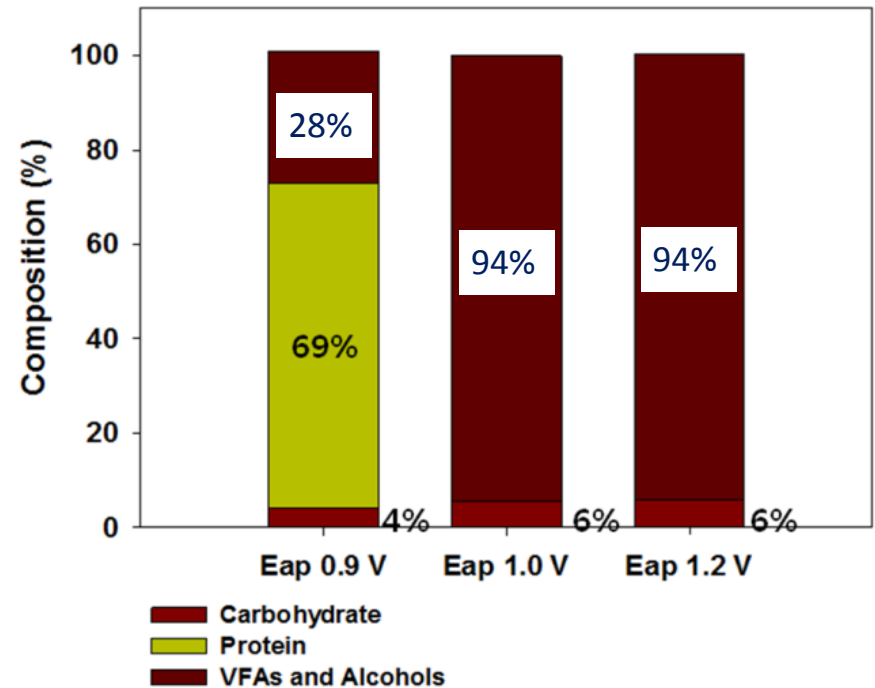
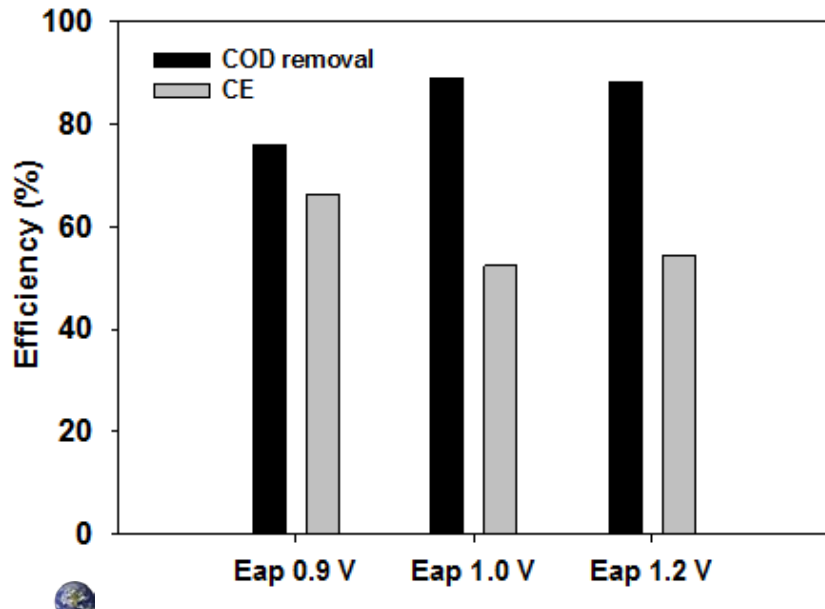


- Higher applied voltages resulted in more positive anode potentials which reduces performance
  - ( $P_{An} = -429 \pm 4$  mV at  $E_{ap} = 0.9$  V,  $P_{An} = -338 \pm 38$  mV at  $E_{ap} = 1.0$  V, and  $P_{An} = -113 \pm 38$  mV at  $E_{ap} = 1.2$  V).
  - Cathode potentials relatively constant (from  $-997$  to  $-1031$  mV).
- Current fluctuated at  $E_{ap} \geq 1.0$  V. Likely due to substrate depletion near the reactor outlet
- Energy recovery based on electrical energy added and hydrogen gas recovered ( $\eta_E = 223\%$ ) was highest at  $E_{ap} = 0.9$  V; overall energy recovery that includes the substrates ( $\eta_{E+S}$ ) was 64%.
- **More hydrogen gas was generated at increased applied voltages, but  $H_2$  yields were lower.** Highest  $H_2$  yield was 1.1  $H_2$  L/g COD at  $E_{ap} = 0.9$  V.



# Task 3 – Technical Accomplishments

## Removal of Subcomponents in Effluent at Different Applied Voltages



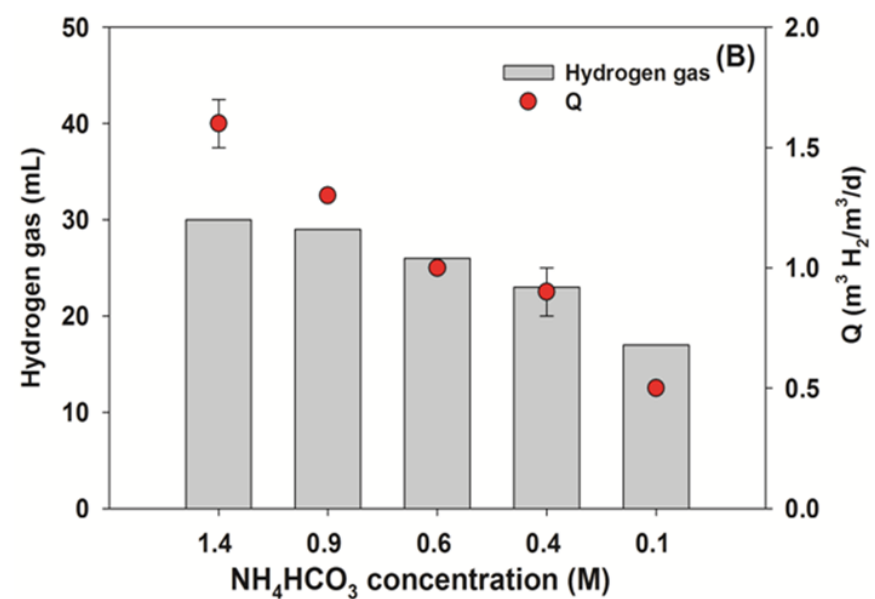
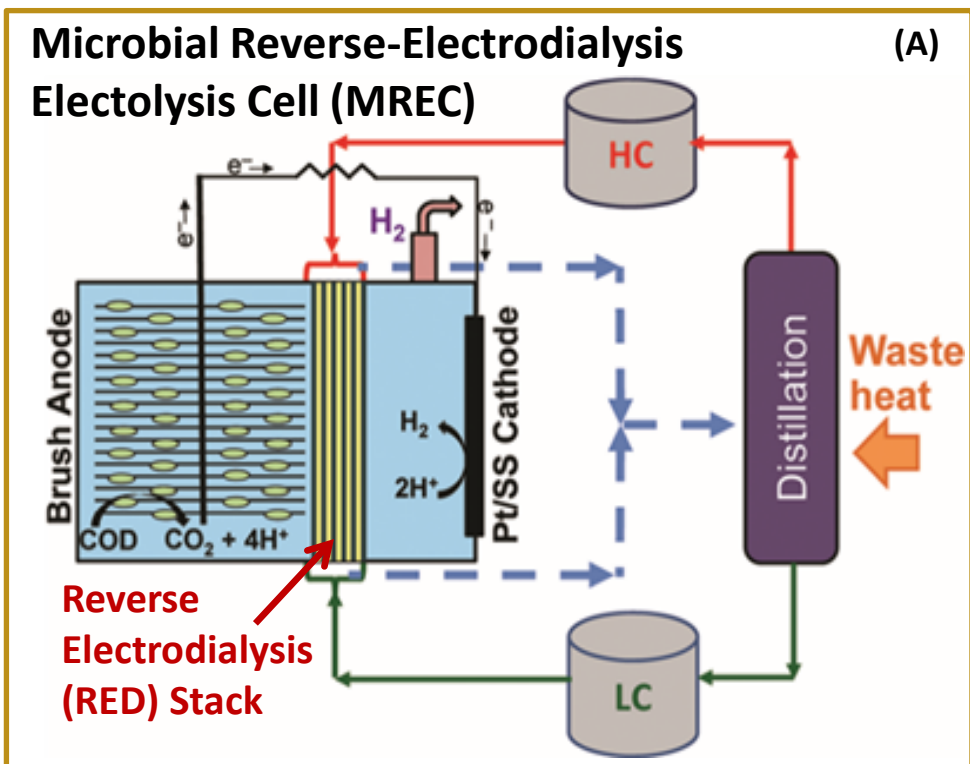
- Higher applied voltages increased COD removal, but lowered coulombic efficiencies (CE).
- Protein was almost completely removed at the higher applied voltages.
- **As CEs were lower at higher applied voltage, degraded protein seemed to be used more for cell growth than electricity generation.**

	Milestone	Completion Date	Status
3.2.3 (FY12)	Correlate removal of the subcomponents of the NREL fermentation effluent with current density and H <sub>2</sub> production	9/12	Completed

# Task 3 – Technical Accomplishments



## Hydrogen Generation in an MREC – No External Power Source



Decreasing HC concentration (1.4 M to 0.1 M) with fixed LC solution (distilled water)

- **Power source (A):** RED stack using high concentration (HC) and low concentrations (LC) of Ammonium bicarbonate,  $\text{NH}_4\text{HCO}_3$ , which can be regenerated. (No electrical grid energy used).
- **Performance (B):** Gas recovery ranged from 30 mL- $\text{H}_2$  to 17 mL- $\text{H}_2$  (93–94%  $\text{H}_2$ , 6–7%  $\text{CO}_2$ ) [bars].
- Gas production rate ranged from 1.6  $\text{m}^3 \text{H}_2/\text{m}^3\cdot\text{d}$  to 0.5  $\text{m}^3 \text{H}_2/\text{m}^3\cdot\text{d}$  [circles] [60 mL reactor, both chambers].
- Coulombic efficiency using acetate  $CE = 72\%$  to 50%. Maximum Yield = 3.4 mol  $\text{H}_2/\text{mol}$  acetate.
- **Current and  $\text{H}_2$  were successfully generated in the MREC using  $\text{NH}_4\text{HCO}_3$  solutions and no external power supply. Construction of a larger reactor (several hundred mL, with multiple anodes) for testing NREL fermentation effluent for  $\text{H}_2$  production is ongoing toward meeting Milestone 3.2.3-1 (9/13).**

# Collaborations



- **Task 1 (Bioreactor):**

Dr. Ali Mohagheghi, National Bioenergy Center at NREL (biomass pretreatment and characterization).

- **Task 2 (Genetic Methods):**

Drs. David Levin and Richard Sparling at the University of Manitoba, Canada. NREL is an international collaborator of the Genome Canada Grant award to co-develop genetic tools for pathway engineering in *C. thermocellum*.

- **Task 3 (MEC):**

Dr. Bruce Logan, Penn State University (microbial electrolysis cells to improve H<sub>2</sub> molar yield). Task 3 was cost shared by other projects of the investigator.

# Proposed Work



## Task 1 (NREL):

- Complete the H<sub>2</sub> production profiles from cellulose (rate, yield) using shorter HRT, varying frequency and volume of medium replacement in sequencing fed batch reactor (FY13, FY14)
- Guided by the optimal parameters derived from above, determine H<sub>2</sub> production rate and H<sub>2</sub> molar yield using lignocellulosic biomass (FY14)

## Task 2 (NREL):

- Test the pyruvate-to-formate pathway mutants for H<sub>2</sub> production rates, H<sub>2</sub> molar yield, and metabolites in bioreactors (FY13, FY14).
- Generate double knockout mutants by also deleting either lactic acid or the ethanol pathway (FY14).
- Using the optimized bioreactor parameters, test the above mutants for improved H<sub>2</sub> molar yield (FY14).

## Task 3 (Penn State):

- Design, build, and test larger prototype MREC reactor with acetate solution for scalability (FY13).
- Evaluate performance of new MREC with NREL fermentation effluent as to H<sub>2</sub> yields, H<sub>2</sub> production rates, and gas production (FY13).
- Acclimate MECs separately to protein (bovine serum albumin) and acetate in batch-fed reactors, followed by continuous mode, and then with NREL fermentation effluent in terms of H<sub>2</sub> yields and gas production rates (FY14).

# Summary



## Task 1:

- Designed and built a feed tank that delivered cellulose reliably to the bioreactors, with less than 10% standard deviation of H<sub>2</sub> output in the bioreactor.
- Increased rate of H<sub>2</sub> production by 3.8-fold and H<sub>2</sub> molar yield by 53% via manipulating HRT, frequency, and liquid volume replacement, at either 2.5 g/L or 5 g/L cellulose loading.

## Task 2:

- Generated proprietary plasmids and optimized protocols for reliable transformation in *C. thermocellum*.
- Produced five *C. thermocellum* mutants lacking the *hpt* gene, which served as the base strain for targeted pathway mutagenesis without leaving an antibiotic marker.
- Produced eight pyruvate-to-formate pathway mutants with ~60% increase in ethanol production.

## Task 3:

- Evaluated performance of MEC using fermentation effluent in terms of H<sub>2</sub> yields, H<sub>2</sub> production rates, and gas composition.
- Verified boosted voltages can increase protein utilization in the fermentation effluent.
- Successfully produced H<sub>2</sub> gas without an external energy input in an MREC using salinity gradient energy.