

Hydrogen from Water in a Novel Recombinant Oxygen-Tolerant Cyanobacterial System

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Overview

Timeline

- Project start: 5-01-05
- Project end: N/A
- Percent complete: 10%

Budget

- Total project funding
 - \$720K (Contractor share)
- Funding for FY06
 - \$0 for the Venter Institute
 - \$26K for NREL
- Funding for FY07
 - \$175K for the Venter Institute
 - \$100K for NREL

Barriers

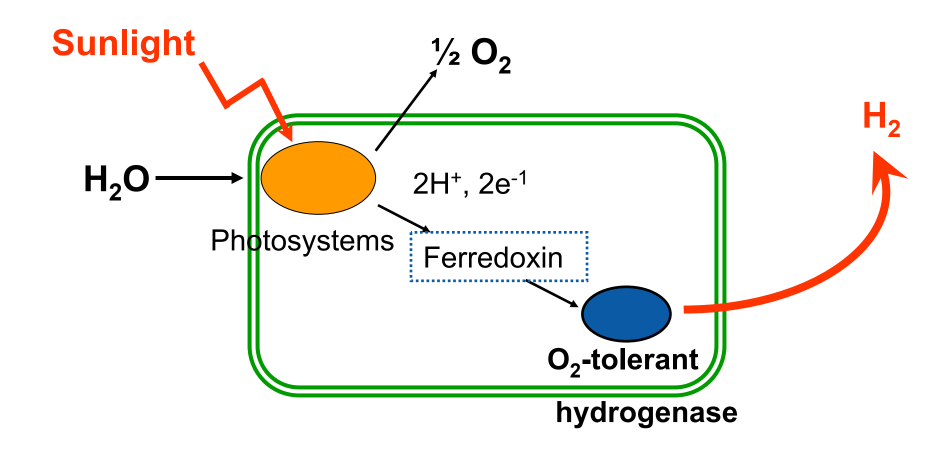
- Barriers addressed
 - Barrier Z, continuity of
 H₂ photo-production

Partner

National Renewable Energy Laboratory

Objective

Develop an O₂-tolerant cyanobacterial system for continuous light-driven H₂ production from water



The overall goal is to produce a cyanobacterial recombinant to produce H₂ continuously

Approaches to address hydrogenase O₂-sensitivity barrier

- Problem: Cyanobacteria have the ability to split water photolytically into O₂ and H₂, but their hydrogenases are highly O₂-sensitive. In contrast, certain bacteria have O₂-tolerant H₂-evolving hydrogenases, but they can not use water as the electron donor.
- Approach: transfer O₂-tolerant hydrogenases into cyanobacteria
 - Identifying novel O₂-tolerant hydrogenases from VI's sampling in international waters and transferring them into cyanobacteria
 - 2. Transferring known O₂-tolerant hydrogenases into cyanobacteria

Technical Approach 1 (Venter Institute)

 Identification of novel O₂-tolerant hydrogenases from environmental microbes in international water

(This task was not funded by DOE during 2006, and the following progress was made with internal funding from the Venter Institute.)



Sorcerer II Expedition: an Global Ocean Sampling Project accomplished at the Venter Institute

The Sargasso Sea Project: A Pilot Study for VI's Sorcerer II Expedition

- Generated 1.045 billion base pairs of non-redundant sequences
- Found 1800 genomic species, including 148 new bacterial species
- Identified 1.2 million new genes, including 782 new rhodopsin-like photoreceptors

The Sorcerer II Global Ocean Sampling Expdition

- Generated 7.7 million sequencing reads equivalent to 6.3 billion bps of non-redundant sequences
- Identified 6.12 million proteins (covering nearly all known prokaryotic protein families) that nearly double the number of current proteins
- Revealed 3995 GOS-only clusters that have no homology to known protein families
- The rate of new protein families discovered is linear with the addition of new sequences, implying we are still far from discovering all the protein families in nature.

Environmental microbes have significant potential in carrying new genes or pathways that may improve efficiency of renewable energy production.

Searched 1.2 million peptides in the Sargasso Sea Dataset for putative novel NiFe-hydrogenases

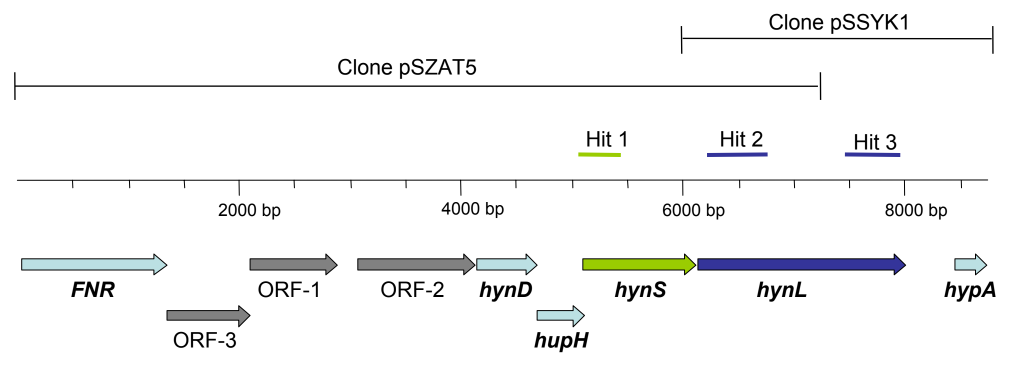
- 20 peptide hits representing the sequences from 10 NiFe-hydrogenases were identified by using 14 Hidden Markov Models (HMMs) of hydrogenases that we constructed.
- Three of 10 NiFe-hydrogenases are known hydrogenases, which are

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\begin{cases} \textit{Shewanella oneidensis} \text{ quinone-reactive } H_2 \text{ase (8 hits)} \\ \textit{Citrobacter freundii } H_2 \text{ase-1 (2 hits)} \\ \textit{E. coli/S. enterica } H_2 \text{ase-3 (1 hit)} \end{cases}
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• The remaining 7 H_2 ases appear to be novel, with homology to

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E. coli/S. enterica H<sub>2</sub>ase-2 (3 hits)
E. coli/S. enterica H<sub>2</sub>ase-3 (1 hit)
E. coli/S. enterica H<sub>2</sub>ase-4 (1 hit)
Gloeothece sp. uptake H<sub>2</sub>ase (1 hit)
Thiocapsa roseopersicina O<sub>2</sub>-stable H<sub>2</sub>ase (3 hits)
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Cloned genes of a putative novel hydrogenase from the Sargasso Sea with strong homology to a *Thiocapsa* O₂-tolerant hydrogenase



hynL: Large subunit of novel hydrogenase

hynS: Small subunit of novel hydrogenase

hynD: NiFe-hydrogenase maturation factor

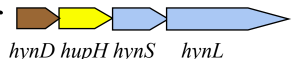
hupH and hypA: NiFe-hydrogenase expression protein

Systems Used to Heterologously Express the Novel Hydrogenase

Host organism: Thiocapsa roseopersicina

- Well defined wild-type strain contains 3 different hydrogenases.
- Hydrogenase biosynthesis is understood in detail.
- A suitable mutant ($\triangle hynSL$, $\triangle hupSL$, $\triangle hoxH$) is constructed.
- Genetic techniques have been developed for this organism.

Expression vector pAmDHSL, for expression of



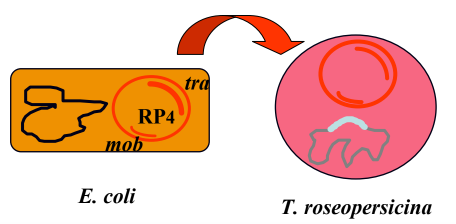
- pAmDHSL can self-replicate in *T. roseopersicina*
- It is transferred from *E. coli* into *T. roseopersicina* through conjugation.

Hydrogenase activity of *T. roseopersicina*

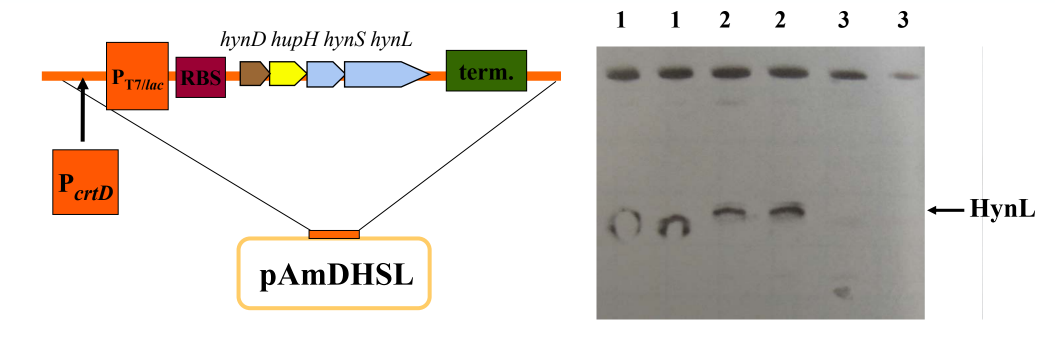
WT strain vs. the mutant $(\Delta hynSL\Delta hupSL\Delta hoxH)$



Conjugational system



The novel hydrogenase from environmental samples in the Sargasso Sea is heterologously expressed in *Thiocapsa roseopersicina*



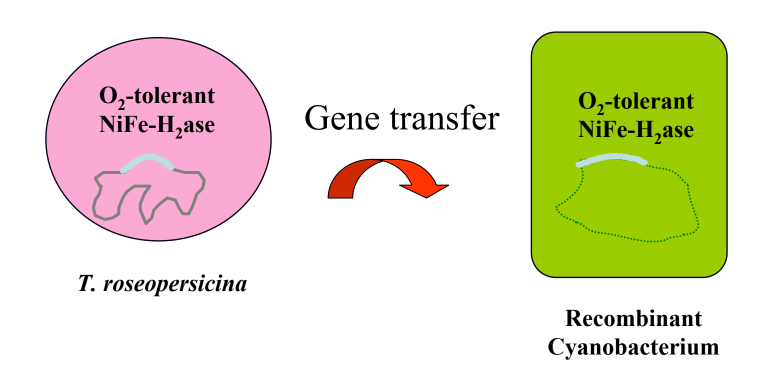
- 1. *T. roseopersicina* wild-type strain (BBS), positive control
- 2. T. roseopersicina Δ Hyn, Δ Hup, Δ Hox strain containing pAmDHSL (hynDhupHhynSL)
- 3. T. roseopersicina Δ Hyn, Δ Hup, Δ Hox strain (GB112131), negative control

Rabbit polyclonal antibody specific for *T. roseopersicina* HynL was used for Western blotting

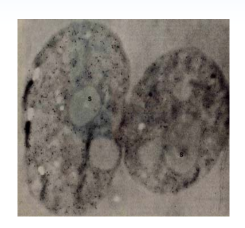
Technical Approach 2.1 (Venter Institute)

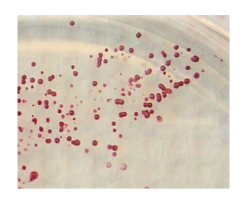
Transferring a known O₂-tolerant NiFe-hydrogenase from bacterium *Thiocapsa roseopersicina* into cyanobacterium *Synechococcus* PCC7942

(This task was not funded by DOE during 2006, and the following progress was made with internal funding from the Venter Institute.)

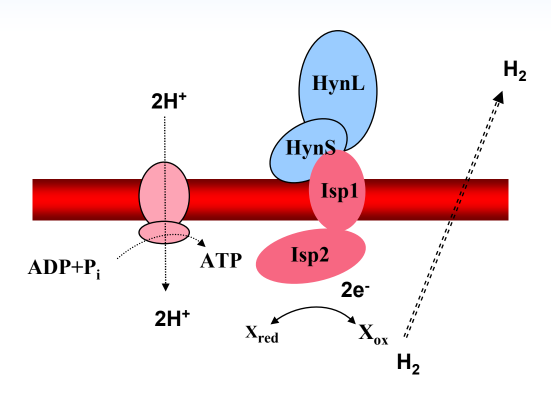


Phototrophic purple sulfur bacteria *Thiocapsa roseopersicina* carries an O₂-tolerant hydrogenase (Hyn)





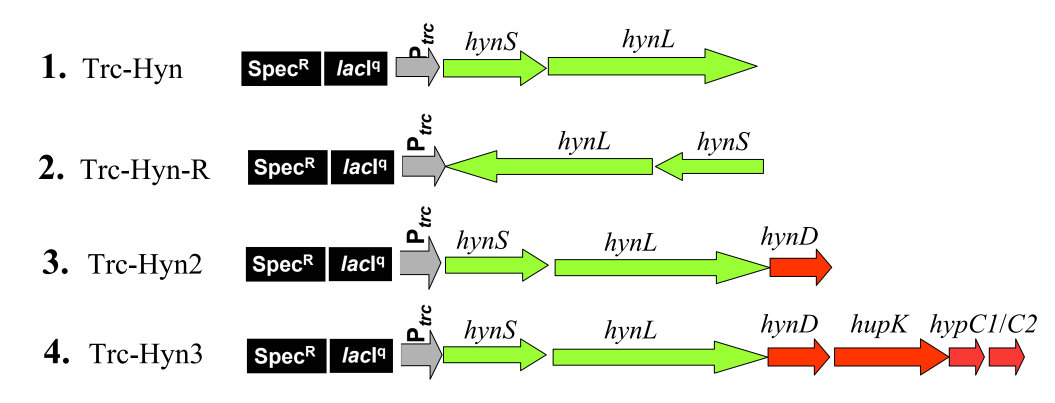
T. roseopersicina



High O₂ / thermal stability, and resistance to proteolysis

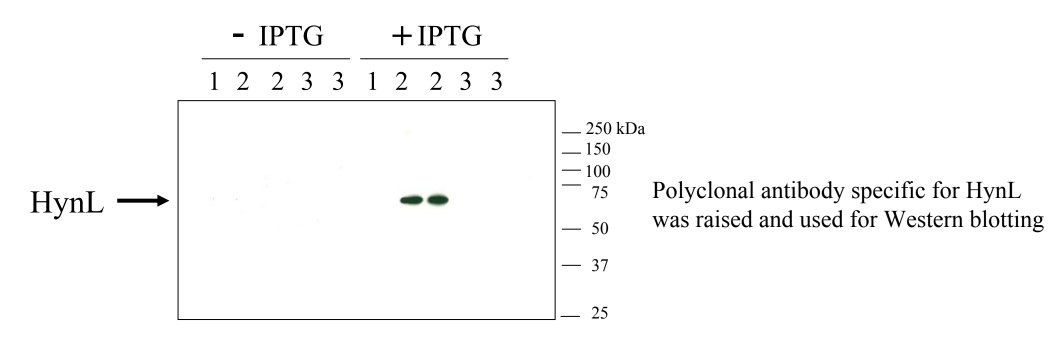
- ➤ Structural subunit: HynS and HynL
- ➤ Electron transfer subunit: Isp1 and Isp2

Transferred Hyn's structural and accessory genes into the genome of cyanobacterium *Synechococcus sp* PCC7942 strain



P_{Trc:} an IPTG-inducible promoter; *hynS* and *hynL: Hyn's* structural genes (green); *hynD*, *hypC1*, *hypC2*, and *hupK:* accessory genes essential for Hyn's assembly (red)

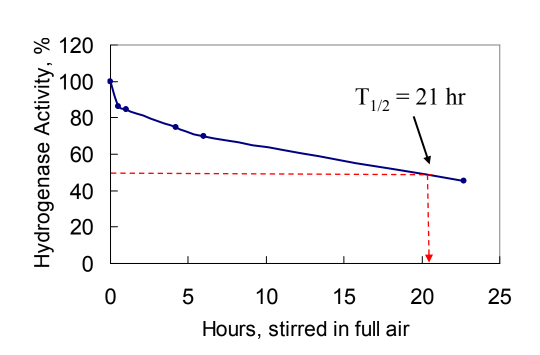
Confirmed IPTG-inducible expression of O₂-tolerant hydrogenase Hyn in constructed recombinant *Synechococcus sp* PCC7942 strains

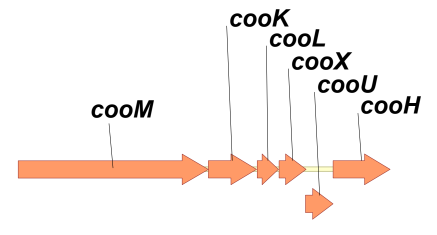


- 1. Synechococcus sp PCC7942 wild type strain
- 2. Synechococcus sp PCC7942 Trc-Hyn
- 3. Synechococcus sp PCC7942 Trc-R-Hyn

Technical Approach 2.2 (NREL)

- Goal: Transfer the O₂-tolerant hydrogenase from Rubrivivax gelatinosus CBS into the cyanobacterial hosts Synechocystis sp. PCC6803
- > Purple non-sulfur photosynthetic bacterium Rx. gelatinosus CBS was isolated from soils in the metro Denver area
- > Rubrivivax evolving hydrogenase displays a half-life of 21 hours in air
- > Structural and accessory genes encoding the Rubrivivax hydrogenase have been cloned

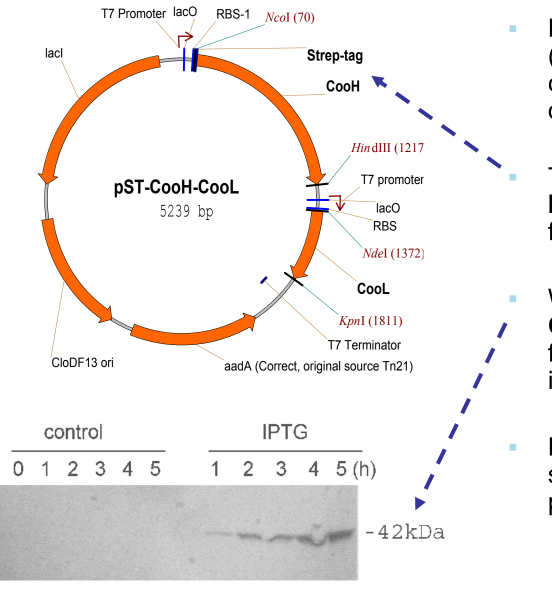




Structural subunits: CooL and CooH

Electron-transfer subunits: CooX and CooU

Expression of Rubrivivax Hydrogenase (CooLH) in E. coli



- Hydrogenase subunits: CooL (small subunit) and CooH (large catalytic subunit) are cloned into a duet vector pCDFDuet-1
- The CooH catalytic subunit is labeled with a **Strep II tag** to facilitate affinity purification
- When expressed in *E. coli*, tagged CooH (42kDa) is detected in cell-free extracts following IPTG induction
- Protease digestion/mass spectrometry confirmed the 42 kDa protein as CooH

Future Work (Venter Institute)

Remainder of FY2007:

- Purify hetero-expressed novel NiFe-hydrogenase (identified from the Sargasso Sea environmental samples) from *T.* roseopersicina
- Clone more accessory genes of Thiocapsa O₂-tolerant hydrogenase into cyanobacterial expression vector pTrc-NSI

• FY2008:

- Continue to transfer the accessory genes of *Thiocapsa* O₂-tolerant hydrogenase into cyanobacterium *Synechococcus* PCC7942
- Characterize purified novel hydrogenases that was identified from the Sargasso Sea environmental samples

Future Work (NREL)

Remainder of FY2007:

The Strep II-tagged as well as non-tagged Rubrivivax hydrogenase (CooLXUH) will be inserted into the pPETE_sigD plasmid (Dr. R. Burnap of Oklahoma State Univ.) for delivery into Synechocystis sp. PCC6803

FY2008:

 Systematically express Rubrivivax hydrogenase structural genes (cooMKLXUH) and accessory genes (hypABCDEF) for expressing foreign O₂-tolerant hydrogenases in Synechocystis sp. PCC6803

Summary

Approach 1 (Venter Institute)

- 1. 1.2 millions of peptide sequences in the Sargasso Sea dataset were searched for NiFe-hydrogenases by using 14 hydrogenases HMMs we constructed. 10 NiFe-hydrogenases were identified from this search (VI).
- 2. One novel NiFe-hydrogenases identified has strong homology (64% identity and 75% similarity) to *Thiocapsa* O₂-tolerant hydrogenase Hyn. Its structural and accessory genes have been cloned and sequenced. This novel hydrogenase was successfully hetero-expressed in *T. roseopersicina*, as demonstrated by Western blotting results. This study represents the first conversion of an environmental DNA into a hydrogenase (VI).

Approach 2 (Venter Institute and NREL)

- **1.** Both structural and essential accessory genes of *Thiocapsa* O₂-tolerant NiFe-hydrogenase, such as *hynS*, *hynL*, *hynD*, *hypC1*, *hypC2*, and *hupK*, have been transferred into the genome of cyanobacterium *Synechococcus sp* PCC7942 (VI).
- 2. Thiocapsa O₂-tolerant hydrogenase Hyn is heterologously expressed in Synechococcus sp PCC7942 upon IPTG induction, as demonstrated by Western blotting in which HynL-specific polyclonal antibodies were used (VI).
- 3. The structural genes encoding the *Rubrivivax* O₂-tolerant hydrogenase have been modified with a Strep II affinity tag to simplify purification. These genes were cloned in a pCDFDuet-1 vector and the CooH catalytic subunit was heterologously expressed in *E. coli* as the host initially, as demonstrated by Western blotting. Work is underway to express *Rubrivivax* hydrogenase genes in the cyanobacterium *Synechocystis sp.* PCC 6803 (NREL).