2009 DOE Hydrogen Program

Montana Palladium Research Initiative:

Use of Biological Materials and Biologically Inspired Materials for H₂ Catalysis

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DOE Project ID#: PDP 19 Peters



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Overview

Timeline

- Start Aug. 2006
- End Dec. 2009

Budget

- Total project funding \$1,303,041
 - DOE \$1,031,433

Barriers addressed

- Stability/Durability
- Oxygen Sensitivity
- Electron Donors
- Coupling

Partners

Montana State University



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Approaches

Couple Different Catalyst Systems for Light Driven Hydrogen Generation

Biological catalysts (Hydrogenases)

Nanoparticle biomimetic catalysts



Objectives Addressed

- 1. Immobilize hydrogenase in gels
- 2. Determine basis of hydrogenase stability
- 3. Improve conductivity, mass transfer, and hydrogen production in gels
- 4. Biomimetic hydrogen production catalyst synthesis
- 5. Photocatalyst synthesis
- 6. Coupling catalysts to electrode surfaces
- 7. H₂ production device fabrication

Approach:

Biological and Biomimetic Catalysts for H₂ production





Hydrogenase Enzymes (protein architecture protecting Metal sulfide active site)

Protein encapsulated nano-catalyst

Coupled Reactions to Generate Hydrogen



GOAL: use **biological catalysts** and develop **biomimetic catalysts** with a variety of sacrificial electron donors or electrochemical source of e^- to produce H_2

Issues and Barriers: Catalyst Stability

- Durability shelf life
- Reusability
- Product based Inhibition
- Oxygen tolerance / resistance
- Susceptibility to proteolytic inactivation
- Optimization electron transfer, pH, ionic strength, mediators

Hydrogenases: Highly evolved finely tuned catalysts for *hydrogen oxidation and proton reduction (hydrogen production)*

C. pasteurianum



 $H_2 \cong 2H^+ + 2 e^-$

"H Cluster"

Desulfovibrio gigas



Cellular location

Membrane Associated Soluble Periplasmic Cytoplasmic



NiFe Cluster

Microorganisms:

hydrogen, acetategrown, methanogenic, green, purple, cyanobacteria; algae; fungus.

Stable NiFe hydrogenase from purple sulfur bacteria forms supermolecular structures





Electron microphotograph of hydrogenase complexes from *T. roseopersicina* negatively stained with 2% uranyl acetate

| Properties | Thiocapsa roseopersicina | |
|---------------------|-----------------------------|--|
| Large subunit | 64kDa | |
| Small subunit | 34kDa | |
| Temperature optimum | 80°C | |
| Stability to Oxygen | stable | |

Cryo reconstruction of hydrogenase from *T. roseopersicina* at ~33 Å.



Structural studies indicate a role for C-termini in the stability and super molecular complex formation of hydrogenase

T. r. AALEQAKRPSVIWLSFQECTGCTESLTRSH LTAKKRPSVVYLHNAECTGCSESVLRTV D.r. T.r. APTLEDLILDVISLDYHHTLQAAAGDAAEH D.r. DPYVDELILDVISMDYHETLMAGAGHAVEE T.r. AREQAMAANPGEYLVIVDGSIPGPDSNPGY D.r. ALHEAI --- KGDFVCVIEGGIPMGDGGY-W 100 110 120 T. r. STVAGHSNYAMLMETVENAAAVIAVGTCAT D.r. GKVGGRNMYDICAEVAPKAKAVIAIGTCAT 130 140 150 T. r. FGGLPGAN PNPTGAMSVMDLVKDKPV--IN D.r. YGGVQAAKPNPTGTVGVNEALGKLGVKAIN 160 170 180 T.T. VSGCPPIPMVITGVIAHYLTFGRLPELDAY D.r. IAGCPPNPMNFVGTVVHLLTKG-MPELDKQ 190 210 200 T.r. NRPMAFFGQSIHDRCYRRPFYDKGLFAKTF D.r. GRPVMFFGETVHDNCPRLKHFEAGEFATSF 220 230 240 T. r. DDEGARLGWCLYELGCKGPTTYNACATMRW D.r. GSPEAKKGYCLYELGCKGPDTYNNCPKQLF 260 270 T. r. NDGTSWPVEAGHPCLGCSEPRFWDA-GGFY D.r. NO-VNWPVOAGHPCIACSEPNFWDLYSPFY VSVPTSASGVNVLAAGAAGAIVGGA T.r. NT D.r. SA T.r. **OTKTAVAHROPVTVEELEAKI** D.r.

В



Sequence alignment of small subunits of stable hydrogenase from *T. roseopersicina* (*T. r.*) and *D. gigas* (*D. g.*) (A) and cryoEM m (B). In the red box the C-termini residues, which could be involve in the cap formation are shown.

With Liang Tang – University of Kansas

Encapsulation of purified active hydrogenases in tetramethyl ortho silicate gels



- Nanoscopic encapsulation;
- Immobilization of unaltered enzyme
- "Heterogeneous material"



Screening electron microscopy photographs of sol-gel

Recovery of hydrogenase activity* encapsulated in Sol-Gel

| Hydrogenase | Solution | Gel | Solution/Gel (%) |
|-----------------------------|----------|------|------------------|
| C. pasterianum (extract) | 12550 | 7581 | 60.4±16 |
| L. modestogalophilus | 9150 | 6175 | 67.5±9 |
| T. roseopersicina | 12600 | 8834 | 70.1±3 |

•Activity measure at 25° C indicated in nmol/min/mg protein. Values represent average rate over a four-hour period.

Carbon nano tubes incorporated into Sol Gels





Scanning electron microscopy photographs of sol-gel in presence of carbon nanotubes

- → Enhance electron transfer
- → Facilitate electron transfer between immobilized mediators and hydrogenase
- → Facilitate electron transfer between electrodes and hydrogenase in devices

Formation of electro active matrix by encapsulation of the hydrogenase with multiwall carbon nanotubes

Hydrogen production by the silica gel matrix containing hydrogenase (H2ase), carbon nano tubes (NT) and methyl viologen (MV)



Hydrogenase activity highest in gels with hydrogenase and carbon nanotubes coencapsulated and activity is observed without addition of the redox mediator methyl viologen The addition of polyethylene glycol (PEG) to the sol gel enhances the hydrogen production by encapsulated hydrogenase



Poly anions and cations modulate the pore size in gels and mass transfer

Controlled synthesis of electroactive polymer gels – controlling protein adsorption, mass transfer, and conductivity



Biomimetic Catalysts - Synthesis of Pt⁰ Encapsulated Within a Protein Cage Architecture



Control of Pt cluster size (monitored by NCMS) correlation between activity and cluster size - defining the minimum catalytic cluster

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Observed # of Pt²

20

200

60



Relative abundance (%)

Moving beyond Pt... Pd Nanoparticles encapsulated with Ferritin as H₂ Catalysts





Pd particles show significantly lower activity than Pt

Schematic of Pd encapsulated within Ferritin

Stable to 80 °C Oxygen insensitive

Photocatalyst synthesis - Hematite (α -Fe₂O₃) in ferritin





Attachment of catalysts to electrode surfaces – Cyclic voltammetry to probe e⁻ transfer to catalysts



Design and fabrication of prototype devices



Testing - Measuring a Photocurrent with α -Fe₂O₃ Ferritin



Summary

- 1) Hydrogenase can be immobilized in gels and retain activity
- 2) C-termini contributes to stability of stable hydrogenases
- 3) Carbon nanotubes enhance performance of hydrogenase/catalyst in gels
- 4) Pore size in gels can be controlled effecting mass transfer and hydrogenase/catalyst activity
- 5) Pt and Pd hydrogen production catalysts can be synthesized using biological templates
- 6) Photocataysts can be synthesized using biological templates
- 7) Catalysts can be attached to conducting surfaces
- 8) Prototype device has been fabricated and initial testing is underway

Future Work

- 1) Device testing and optimization
- 2) Establish benchmarks for hydrogen production efficiency
- 3) Evaluate hydrogen production efficiency (electrochemical, photochemical, chemical reducing equivalents)
- 4) Evaluate device for durability and sustained H₂ production