HyMARC: Hydrogen Storage Materials Advanced Research Consortium

2016 DOE Hydrogen Annual Merit Review June 8, 2016



<u>B. Wood</u>, T.W. Heo, J. Lee, S. Bonev, M. Morales, T. Baumann

Lawrence Livermore National Laboratory



LLNL-PRES-XXXXXX

This work was performed under the auspices of the U.S. Department of Energy by Lawrence Livermore National Laboratory under contract DE-AC52-07NA27344. Lawrence Livermore National Security, LLC



Timeline	Barriers addressed	
Project start date: 9/17/2015 Phase I end date: 9/30/2018	 Lack of understanding of hydrogen physisorption and chemisorption (Barrier O) System weight and volume (Barrier A) Charge/discharge rate (Barrier E) 	
Budget	Team	
FY15 DOE Funding: \$250K FY16 Planned DOE Funding: \$735K Total Funds Received: \$985K	Funded Partners: Sandia National Laboratories (lead) Lawrence Berkeley National Laboratory	



Relevance & objective

HyMARC will provide **community tools** and **foundational understanding** of phenomena governing thermodynamics and kinetics to enable development of solid-phase hydrogen storage materials

Theory, simulation, & data



- Quantum Monte Carlo for sorbents
- DFT/ab initio molecular dynamics for bulk/surface/interface chemistry
- Classical MD & kinetic Monte Carlo for non-equilibrium transport
- Phase-field modeling for solid-state phase transformation kinetics
- Computational spectroscopy
- Community software & databases

Controlled synthesis



- Metal organic frameworks (MOFs) Hierarchical integrated bulk and
- nanoscale metal hydrides
- High-pressure synthesis
- Functionalized carbon encapsulants and porous nanoconfining media
- Sorbent suite for model testing and validation

In situ characterization



- Ambient-pressure XPS for gassurface interactions
- Soft X-ray absorption and emission spectroscopy
- Electron microscopy and X-ray spectromicroscopy
- Low-energy ion scattering for surface hydrogen detection

Approach: HyMARC tasks target thermodynamics and kinetics





Approach: Validated multiscale modeling

Modeling approach focuses on integrating techniques at multiple scales to tackle complexities of **"real" (beyond-ideal) materials**



LLNL contributions to HyMARC

Multiscale modeling Porous carbon synthesis **Ted Baumann** DFT & ab initio molecular dynamics: **Brandon Wood** Phase-field mesoscale kinetic modeling: **Tae Wook Heo** Pat Campell **Kinetic Monte Carlo: Stanimir Bonev Quantum Monte Carlo: Miguel Morales** X-ray absorption/emission Postdocs: ShinYoung Kang, Keith Ray, **Roman Nazarov, Patrick Shea** Jon Lee 200



Photon Energy/eV

Progress towards milestones with key LLNL FY16 activities

Q1: (S) Synthesize library of bulk-phase model storage systems for T1-T5

Task 1: Synthesize and characterize high-surface area carbon frameworks with pore sizes < 2 nm (100%)

Q2: (S) Demonstrate size control method for one prototype complex hydride nanostructure *Task 1: Perform sensitivity studies to determine design rules for nanoconfined metal hydrides (50%)*

Q3: (C) Demonstrate in-situ soft X-ray AP-XPS, XAS, XES tools, with sample heating

Tasks 3 & 5: Performed XAS spectroscopy of hydrides and dopants at LBL/ALS, SLAC, and Canadian Light Source (75%)

Q4: (C+T) Identify hydride mobile species and diffusion pathways

Task 2: Perform ab initio molecular dynamics of surface and bulk model hydrides (80%) Task 2: Gather defect formation energies and migration barriers for model materials (70%) Task 2: Construct initial KMC framework for interface/amorphous transport (100%) Task 3: Compute H² dissociation energetics on model surfaces (75%) Task 3: Construct initial microkinetic model for surface reactions (100%) Task 4: Construct initial phase-field modeling code for solid-solid interface kinetics (100%)

Q4: (S+C) Synthesize library of nanoparticles: 1 – 5 nm, 5 – 10 nm, > 10 nm for one prototype hydride *Tasks 1-5: Select model materials for facilitating experiment-theory feedback within each Task (100%)*



Task 1 Accomplishment: Accurate sorbent energetics

Quantum Monte Carlo (QMC) has been demonstrated as a viable approach for improving and bechmarking the accuracy of predicted H_2 sorption on model MOFs

DFT functionals give very disparate energies, but geometries are reliable and can be used as QMC inputs



Will also be used to parameterize classical potentials for future materials screening (Task 6) and predict isotherms that can compare against Sandia measurements **PMARC**

Task 1 Accomplishment: Porous carbon synthesis

Progress in reproducible synthesis pathways for carbons with tailored porosity & chemistry



Heat-treated and compressed samples delivered to Sandia for high-pressure gas sorption measurements for benchmarking isotherm predictions

Task 1 Accomplishment: Modeling sorbent charge/field effects

Developed first-principles modeling framework for simulating charge and field effects on H₂ physisorption on conductive sorbents for guiding synthesis

Uses Effective Screening Medium Method* to compute physisorption as function of Fermi level



Can be applied to functionalized materials and any 2D conductive sorbents, and will be used to assess polarization effects in HyMARC experiments



*Otani & Sugino, Phys. Rev. B 73, 115407 (2006)

Task 1 Accomplishment: More accurate hydride thermodynamics

Established protocols for more accurate DFT computations of hydride thermodynamics that can improve predictive power beyond established methods by accounting for:

- Dynamical (anharmonic) contributions to free energy via explicit finite-T simulations
- Microstructure, including phase coexistence and interface/surface contributions
- Phase morphology & nucleation: dispersed molecular defects vs. amorphous vs. crystalline



Task 1&4 Accomplishment: Hydride phase fraction predictions

Demonstrated thermodynamic phase fraction prediction as a function of pressure, temperature, and size, accounting for **microstructure**, **phase coexistence**, and **nucleation**



Predicted size-dependent phase fractions for hydrogenation of $Li_3N/[LiNH_2+2LiH]$ system in core-shell microstructure



Successfully predicted Sandia-measured reaction pathway in nanoconfined Li-N-H and confirmed relevance of interfaces in determining phase stability

Task 1,4&5 Accomplishment: Phase nucleation in hydrides

Computed critical nucleus sizes for model hydrides to guide nanoscale synthesis as a means of kinetically suppressing undesirable intermediate phases

Interface energy approximation: $\gamma_{ij} = p_{ij} \cdot (\sigma_i + \sigma_j)$



Computed critical nucleus size for Li-N-H and Mg-B-H systems, including intermediates



Task 1,4&5 Accomplishment: Statistical approach to interfaces

Demonstrated non-equilibrium statistical approach to quantify effects of non-ideal interfaces and additives on reaction pathways

Probability P of Li₂NH phase suppression in nanoscale [LiNH₂+2LiH]/Li₃N sampled over interfacial energies



Sensitivity analysis can steer towards most promising engineering avenues

Task 2 Accomplishment: Ab initio molecular dynamics (AIMD)

AIMD is being used to elucidate transport/reaction pathways and enable finitetemperature calculations of thermodynamic and spectroscopic quantities



Task 2&3 Accomplishment: Multiscale surface hydrogen kinetics

Demonstrated multiscale modeling of surface dissociation and diffusion of hydrogen by synthesizing DFT and continuum approaches

$$\frac{\partial c}{\partial t} = \vec{\nabla} \cdot D(r) \vec{\nabla} c + \psi(\vec{r}) j_0$$



Combining with low-energy ion scattering (Sandia) to analyze surface diffusion



Task 4 Accomplishment: Solid phase transformation kinetics

Demonstrated coupled atomistic and mesoscale framework combining thermodynamics, mechanical stress, and phase nucleation/growth to model solid phase transition kinetics



encapsulated hydrides show predictions capture size-dependent kinetic behavior

500

(slower)

300

Time steps

400

200

Ĭ00

Task 3&5 Accomplishment: XAS spectroscopy of hydrides

Demonstrated X-ray absorption spectroscopy (XAS) for probing catalyst chemistry and surface vs. bulk hydrogenation effects on catalysts within doped hydrides



XAS results on Na-Al-H and Mg-B-H systems are used to devise more realistic surface hydrogenation models, which can be directly validated by computing XAS (LBNL) **PMARC** Collaborations are crucial for realizing HyMARC goals

- NMR and borohydride chemistry: T. Autrey, M. Bowden, B. Ginovska-Pangovska (PNNL; H₂ Storage Characterization Group)
 - Several discussions, plus site visits in November 2015 and April 2016
 - HyMARC will focus on solid-state aspects and PNNL on borohydride chemistry
- Neutron diffraction/spectroscopy: T. Udovic (NIST)
- DFT computations of H₂ physisorption on MOFs: M. Head-Gordon (LBNL; H₂ Storage Characterization Group)
- Li-N-H system: N. Poonyayant, N. Angboonpong, and P. Pakawatpanurut (Mahidol University, Thailand)
- Kinetic Monte-Carlo for solid-state diffusion: H. Kreuzer (Dalhousie U.)

Also extensive collaborations within HyMARC



Remaining challenges/barriers & mitigation strategies

- Understanding internal microstructure evolution is challenging
 - Working with HyMARC team and collaborators to develop protocols for imaging low-Z materials with minimal beam damage
 - Focusing early efforts on materials (e.g., MgH_2) that can be more easily characterized
- Phase transformations of complex hydrides combine composition & phase changes, yet we are currently treating these separately
 - Separate treatment allows each to be independently validated, but integration of both will be explored in collaboration with the $Mg(BH_4)_2$ individual FOA project in FY17

• Need better validation of thermodynamics and transport for amorphous materials

- Working with HyMARC team to synthesize amorphous materials for testing our amorphous transport and thermodynamics theory framework
- Improved interface energy estimates are desirable
 - HyMARC data will be used to benchmark new approaches for dealing with chemical and elastic contributions to the interface free energy
 - Statistical approach allows us to quantify robustness of predictions with respect to interface energy assumptions
- QMC needs reliable input geometries for sorbent energetics
 - HyMARC will collaborate with M. Head-Gordon (LBNL, part of NREL-led consortium) to use high-throughput DFT with corrected functionals on clusters to obtain geometries



Proposed future work

	Milestone	Description & LLNL subtask	% complete
	Q3 FY16	Demonstrate in situ soft X-ray AP-XPS, XAS, XES tools with sample heating <i>Perform synchrotron XAS/XES of hydrogenation on catalyzed materials</i>	100
	Q4 FY16	Identify hydride mobile species and diffusion pathways Compute defect formation energies and ab initio dynamics of mobile species in model systems	25
	Q4 FY16 SMART	Synthesize library of nanoparticles: 1-5 nm, 5-10 nm, > 10 nm for prototype hydride Demonstrate porous carbon nanoconfining medium synthesis with controlled pore sizes	50
	Q1 FY17	Compute H ₂ binding curves by QMC	15
	Q2 FY17	Perform sensitivity analysis of local binding and second-sphere effects	0
~	Q2 FY17 G/NG	Rank improvement strategies: open metal sites, acid sites, polarization effects, phase change materials Perform analysis of potential of open metal sites (via QMC), acid sites/polarization (via DFT), and phase change materials (thermodynamic analysis)	25
t	MARC		

Summary

- Integrated **theory/synthesis/characterization** framework of HyMARC aims to provide foundational understanding and new tools for solid-state hydrogen storage
- FY16 LLNL modeling tasks focused on establishing framework for **multiscale integration** and approaches for **beyond-ideal materials modeling**
- FY16 LLNL synthesis tasks focused on establishing key **strategies for tailored porous carbon**
- FY16 LLNL characterization tasks focused on **spectroscopic changes upon hydrogenation** (complements LBNL activities)
- Early feedback between experiments and theory on the Pd-H and Mg-H systems demonstrates **feasibility of multiscale (de)hydrogenation kinetics simulation**
- Will be moving towards testing on complex hydrides and more complicated sorbents in FY17
- Flexible modeling, synthesis, and characterization protocols are being developed in preparation for application to upcoming FOA projects



Thank you for your attention!

