

Xiaoming Liu, Thomas Fitzgibbons, Paramita Ray, Enshi Xu, Gregory Larsen, Angela D. Lueking (PI), John V. Badding (co-PI), Vincent H. Crespi (co-Pl) *Pennsylvania State University*

Transformation of Hydrogenated Bucky-Balls at High pressure

- Abstract

Thermodynamics dictate that pre-formed C-H structures will rearrange with increased pressure, yet the final carbonhydrogen interactions may depend on the mechanism of hydrogen introduction.

•MD with ReaxFF used to model $C_{60}H_{36}$ and $C_{60}H_{18}$ •Investigate interplay between inter-ball polymerization into sp³ geometries and on-ball sp³ sites of H attachment •Compress multiple crystal structures up to 30 GPa •Develop a theoretical model for H release during hydrocarbon compression

•Test theory and models by comparison to experiment

Simulations imply that to automatically trap hydrogen gas during compression of $C_{60}H_{36}$ will require extreme conditions.





… and also hydrogen release

4 - Conclusions and On-Going Work

Molecular dynamics simulations suggest rapid compression will polymerize $C_{60}H_{36}$, while slow compression will not. Experimentally, FTIR of $C_{60}H_{14}$ that has been compressed up to 52 GPa confirms static high pressure does not modify the C-H stretching region. Raman spectroscopy studies are ongoing to analyze changes in the lattice modes and C-C bonding, and possible evolution of H_2 is being confirmed. Future work includes compression under non-hydrostatic conditions and compression of solid carbon materials in the presence of H_2 .

Exploration of Novel Carbon-Hydrogen Interactions

2 - Diamond Anvil Cell

Mao-Bell type piston-cylinder cells are used. Samples are loaded in a 250-300 µm hole in the metal gaskets and are compressed between the diamonds. High pressure gas can be loaded along with the sample as a pressure transmitting medium or reagent. Diamonds provide unique combination of optical clarity and high strength allowing for *in situ* Raman at GPa pressures

An FT-IR experiment at static high pressure indicates no change in the C-H stretching region up to 52 Gpa, other than the expected red shift with pressure. A candidate H_2 stretch (at ~4250 cm⁻¹, right) is due to either H_2 evolution and subsequent physisorption or etalonging effects. Subsequent Raman experiments (with 458, 488, and/or 633 nm excitation) will differentiate.

In-Situ Micro Raman Detection of Reversible Basal Plane Hydrogenation in Pt-doped Activated Carbon

- Abstract

Development of *in situ* spectroscopic measurement techniques capable of combined high-pressure and variable temperature measurements has allowed us to explore carbon-hydrogen interactions that are unresolved and debated in the literature. In particular we focused on the hydrogen spillover mechanism via Platinum catalyst.

•In situ Raman and DFT to investigate local interaction between atomic H and the graphite basal plane adjacent to a Pt dissociation catalyst •Demonstrate spectroscopic evidence for reversible hydrogenation of the carbon basal plane via the spillover mechanism

These results clarify C-H binding, mechanism for reversibility, and role of carbon structure and doping in mobility and reversibility.

3 - Calculated H Binding Energies and Modes

First principle calculations show the 1160/1180 peak is consistent with C-H wagging, and when H is replace by Deuterium, the observed peak shift is as expected from theory. The exact frequency and binding energy varies with absorption density, and that this mode has a significant Raman intensity.

Ten more configurations calculated but not shown here, among which this configuration has the lowest binding energy. Its Raman intensity ratio to that of the G peak is 0.086.

This work gives spectroscopic evidence of hydrogen spillover onto an activated carbon. Simulations and theoretical calculations are used to confirm the origin of the spectroscopically observed C-H wag mode and develop thermodynamic arguments for adsorption by spillover and subsequent stability and desorption mechanisms.

BES01

