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Form Approved
OMB No. 0704-0188

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|--|------------------------------------|--|--|---|--|
| 1. REPORT DATE (DD-MM-YYYY) 25 May 2016 | | 2. REPORT TYPE Briefing Charts | | 3. DATES COVERED (From - To) 22 April 2016 - 25 May 2016 | |
| 4. TITLE AND SUBTITLE Theoretical Studies of nanocluster formation | | | | 5a. CONTRACT NUMBER | |
| | | | | 5b. GRANT NUMBER | |
| | | | | 5c. PROGRAM ELEMENT NUMBER | |
| 6. AUTHOR(S) Robert J. Buszek, Jerry A. Boatz | | | | 5d. PROJECT NUMBER | |
| | | | | 5e. TASK NUMBER | |
| | | | | 5f. WORK UNIT NUMBER Q188 | |
| 7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES) Air Force Research Laboratory (AFMC) AFRL/RQRP 10 E. Saturn Blvd. Edwards AFB, CA 93524-7680 | | | | 8. PERFORMING ORGANIZATION REPORT NO. | |
| 9. SPONSORING / MONITORING AGENCY NAME(S) AND ADDRESS(ES) Air Force Research Laboratory (AFMC) AFRL/RQR 5 Pollux Drive Edwards AFB, CA 93524-7048 | | | | 10. SPONSOR/MONITOR'S ACRONYM(S) | |
| | | | | 11. SPONSOR/MONITOR'S REPORT NUMBER(S) AFRL-RQ-ED-VG-2016-097 | |
| 12. DISTRIBUTION / AVAILABILITY STATEMENT Approved for Public Release; Distribution Unlimited. The U.S. Government is joint author of the work and has the right to use, modify, reproduce, release, perform, display, or disclose the work. | | | | | |
| 13. SUPPLEMENTARY NOTES For presentation at AFOSR Molecular Dynamics and Theoretical Chemistry Program Review; Arlington, VA (25 May 2016) PA Case Number: #16215; Clearance Date: 5/5/2016 | | | | | |
| 14. ABSTRACT Viewgraph/Briefing Charts | | | | | |
| 15. SUBJECT TERMS N/A | | | | | |
| 16. SECURITY CLASSIFICATION OF: | | | 17. LIMITATION OF ABSTRACT SAR | 18. NUMBER OF PAGES 17 | 19a. NAME OF RESPONSIBLE PERSON J. Boatz |
| a. REPORT Unclassified | b. ABSTRACT Unclassified | c. THIS PAGE Unclassified | | | 19b. TELEPHONE NO (include area code) N/A |

Standard Form
298 (Rev. 8-98)
Prescribed by ANSI
Std. Z39.18



Theoretical studies of nanocluster formation

26 May 2016

Jerry Boatz

Principal Research Chemist

Aerospace Systems Directorate, RQRP

Air Force Research Laboratory

This briefing contains information up to:

Distribution A: Approved for public release; Distribution Unlimited. PA Clearance No: 16215



Outline



1. Introduction

- background, technical approach

2. Core-shell nanoclusters (Mg/Cu, Si/Al, etc.)

- energetic additives for propellants, explosives
- gas generators
- biocidal defeat agents

3. Summary and conclusions

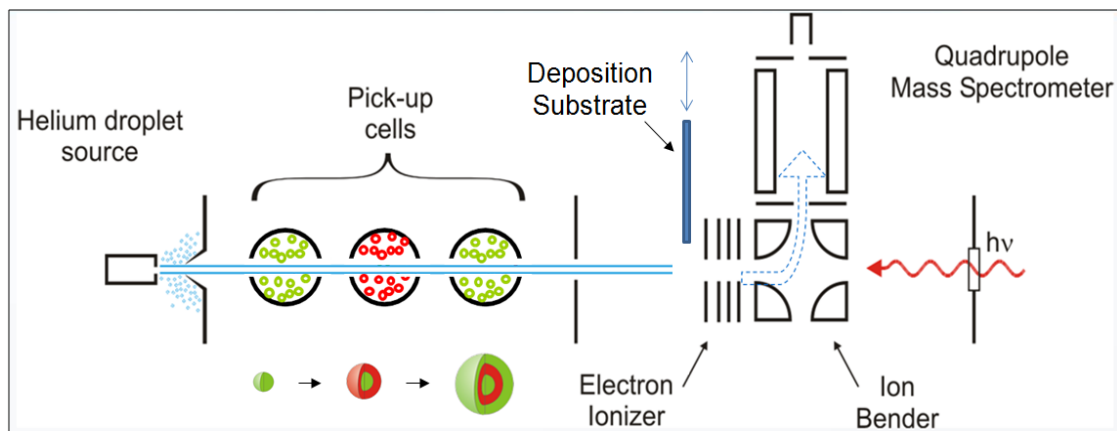


Core-shell nanocluster synthesis



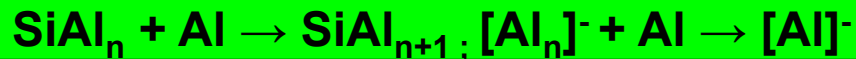
Core-shell nanoclusters such as SiAl_n , Ni_nAl_m , $\text{Al}_n(\text{CuO})_m$, etc. may be useful ingredients in propellants and explosives

- higher energy densities than organics (~ 3x RDX)
- some are resistant to surface oxidation (i.e., “magic clusters”)



Helium droplet experiments at AFRL/RW

Can core-shell nanoclusters be formed under cryogenic conditions (i.e., in helium droplet experiments) via stepwise condensation; i.e., what are the energy barriers (if any) to stepwise addition of atomic Al?

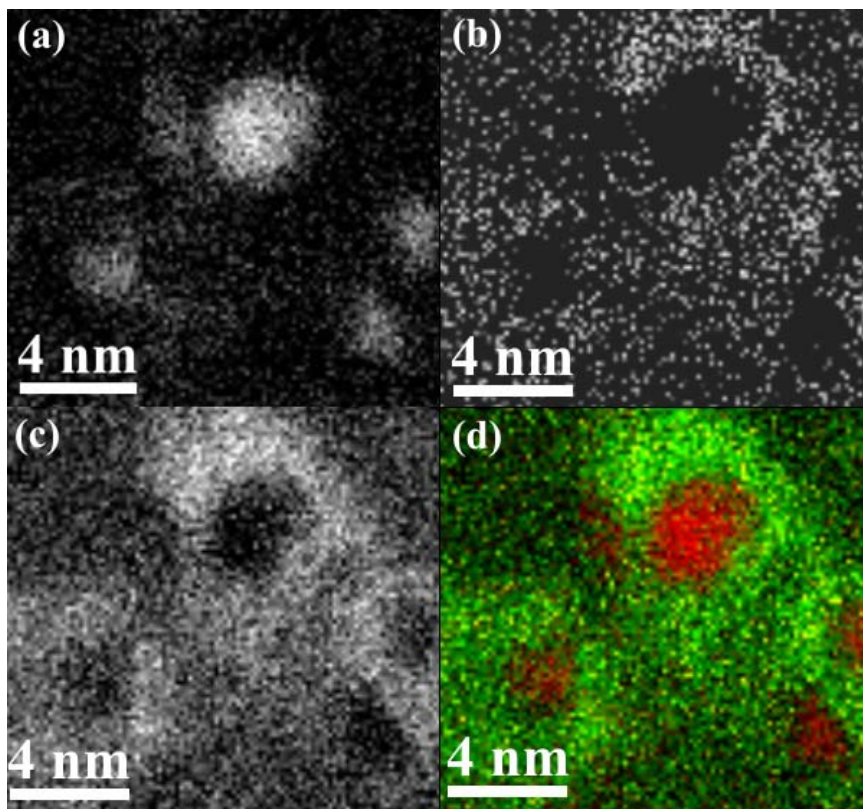




Cu_xMg_y core-shell nanocluster inversion



In helium droplet experiments, Mg atoms were captured in first pickup cell, followed by capture of Cu atoms to form Cu_xMg_y core-shell nanoclusters. However, scanning transmission electron microscopy (STEM) measures show cluster inversion occurred to produce Mg_yCu_x (!)

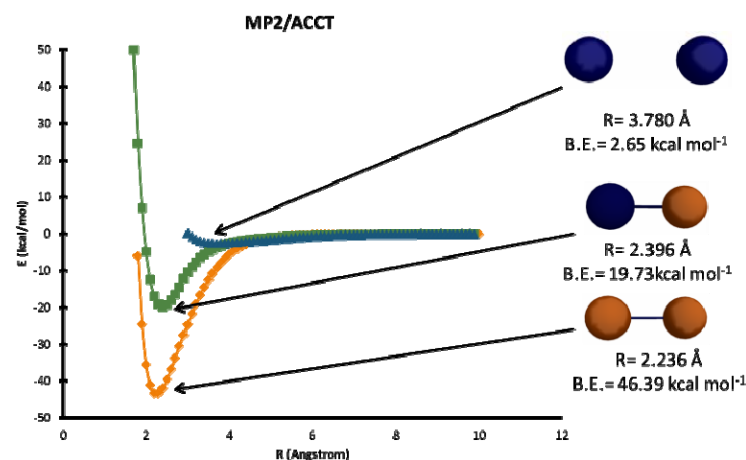


a) copper atoms

b) magnesium atoms

c) oxygen atoms

d) composite image

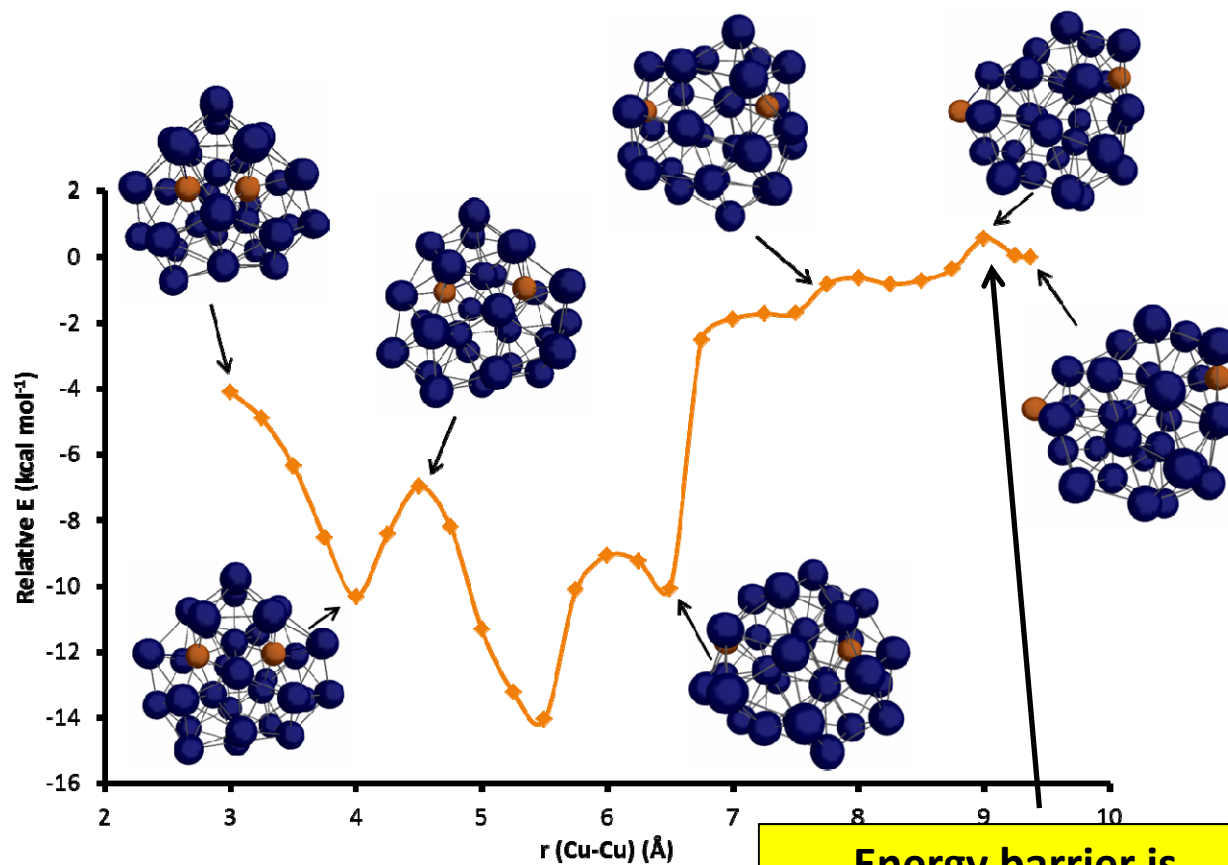




Cu_2Mg_{30}



1. Structure of Mg_{30} cluster was fully optimized.
2. Two Cu atoms were placed on opposite sides of Mg_{30} and structure reoptimized.
3. Distance between Cu atoms was decreased in steps of 0.25 \AA , held fixed, and remaining DOF reoptimized.
4. Total energy plotted as function of fixed Cu-Cu distance.



Energy barrier is
~0.5 kcal/mol

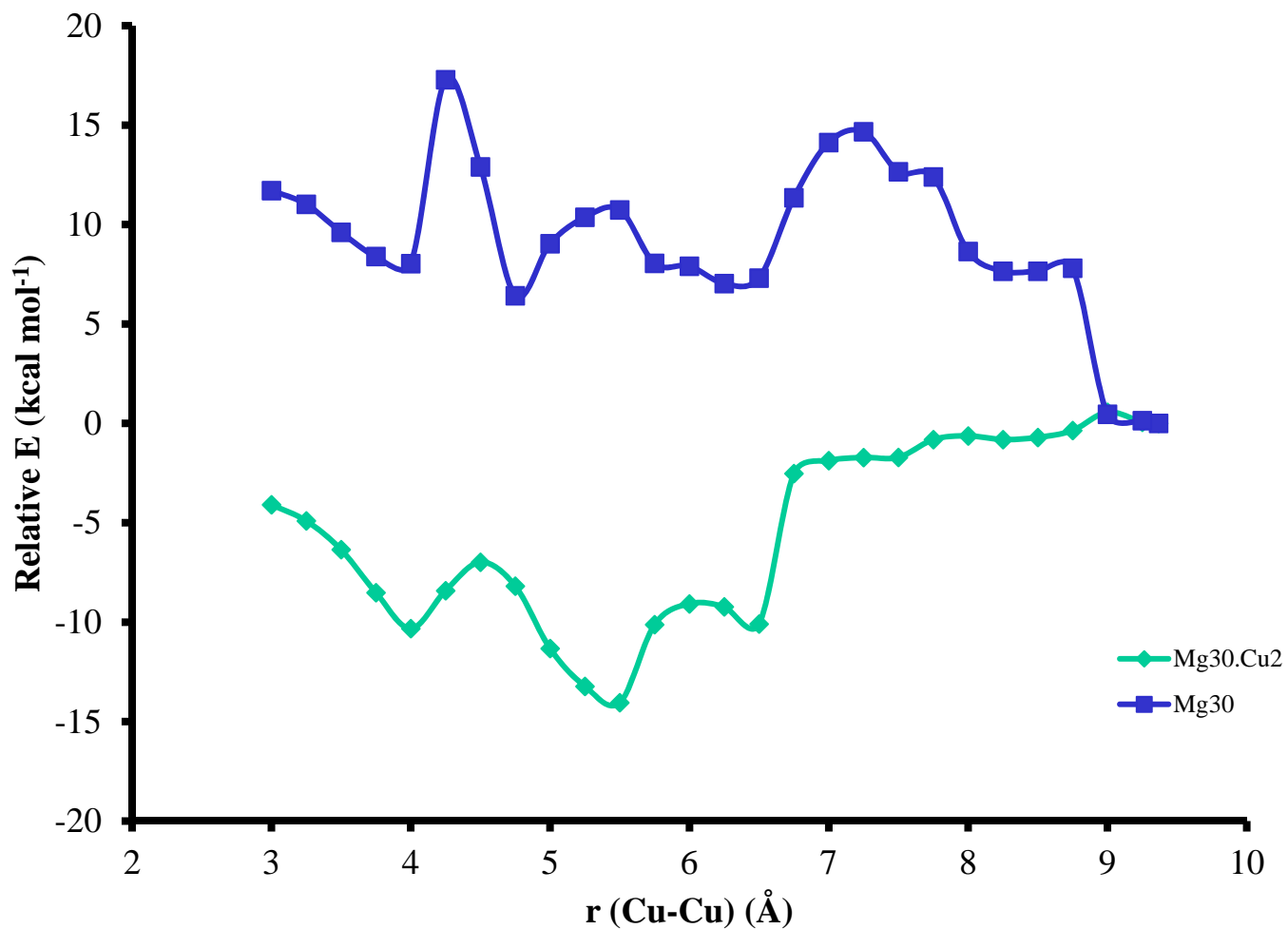
DFT calculations: B3PW91/aug-cc-pwCVTZ(-PP) level



Cu_2Mg_{30} vs. Mg_{30}



Drop in energy in Cu_2Mg_{30} is not due to Mg_{30} rearranging to more stable structure.

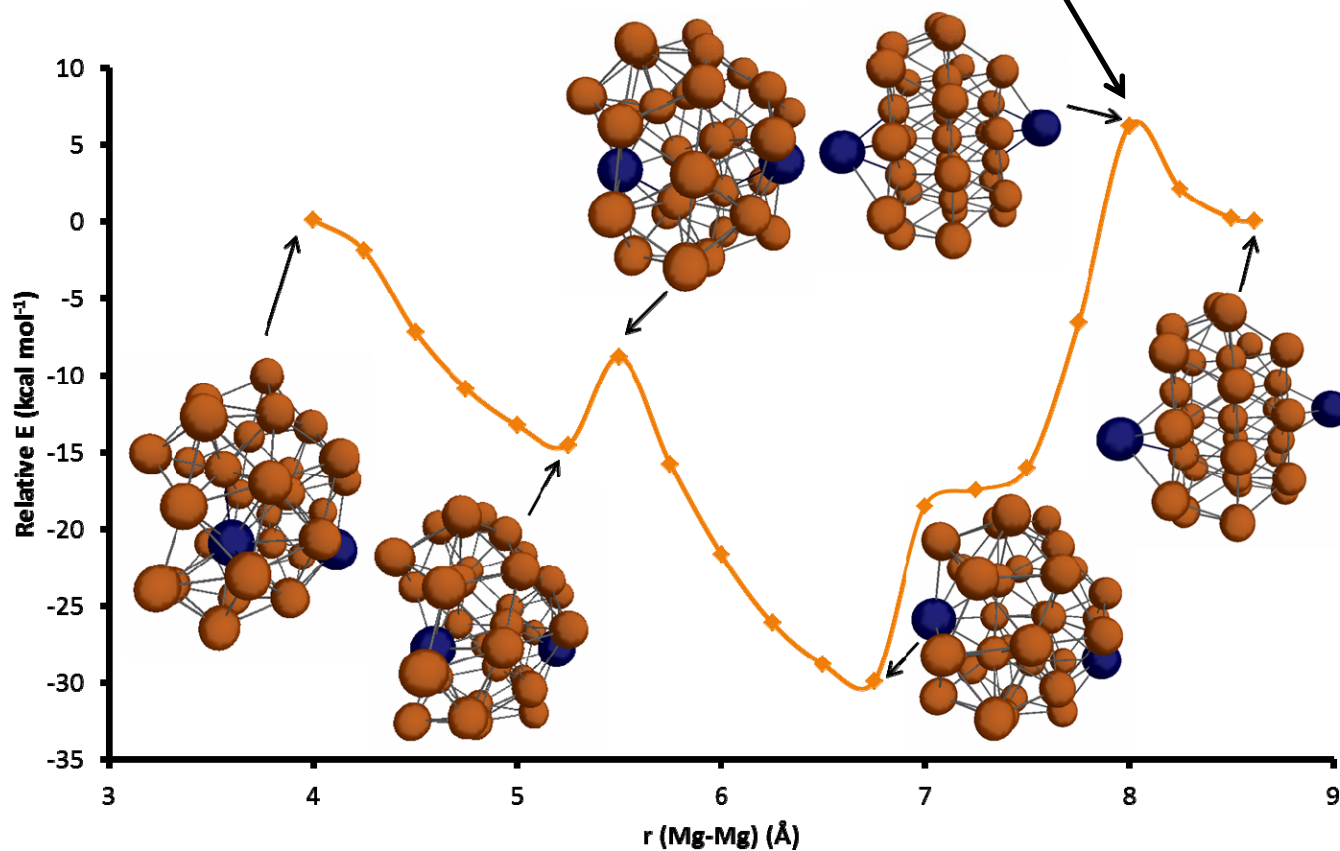




$Cu_{30}Mg_2$



Energy barrier is
~6 kcal/mol

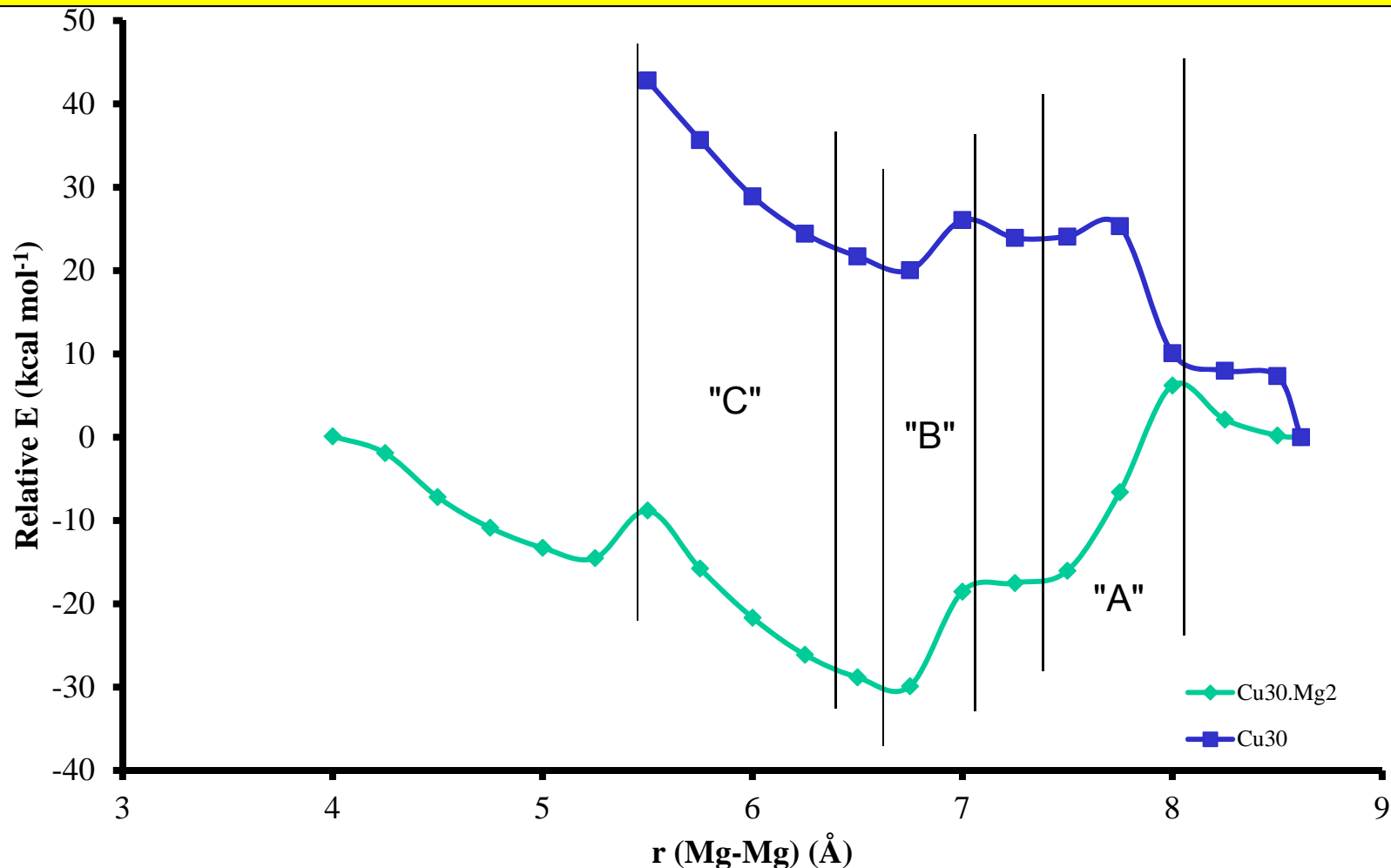




Mg_2Cu_{30} vs. Cu_{30}



Drop in energy in Mg_2Cu_{30} is not due to Cu_{30} rearranging to more stable structure.



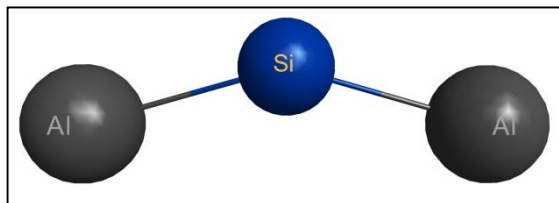


Formation of SiAl_2

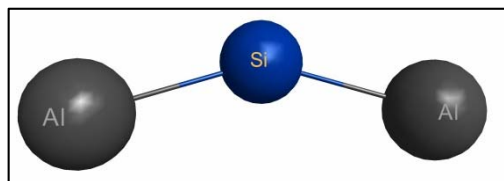


MRMP2(10e,12o)/aug-cc-pvtz level of theory

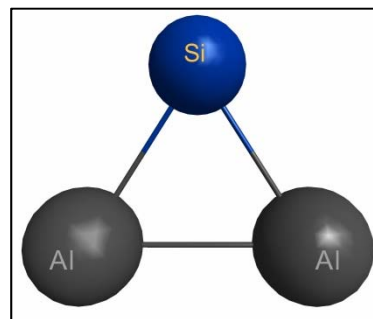
Cyclic isomers



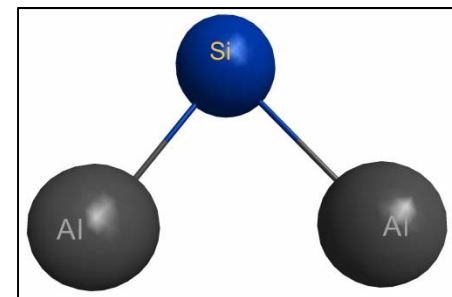
$^5\text{A}_1$, -0.63 eV



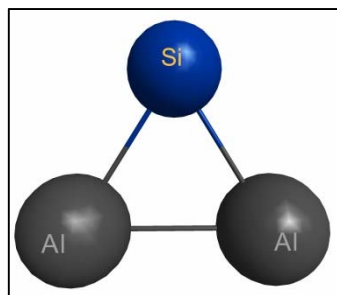
$^3\text{B}_1$, -1.59 eV



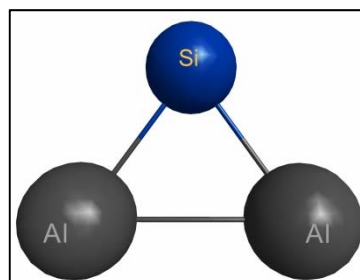
$^5\text{A}_2$, -1.55 eV



$^3\text{A}''$, -1.80 eV



$^1\text{A}_1$, -2.00 eV



$^3\text{B}_2$, -2.04 eV

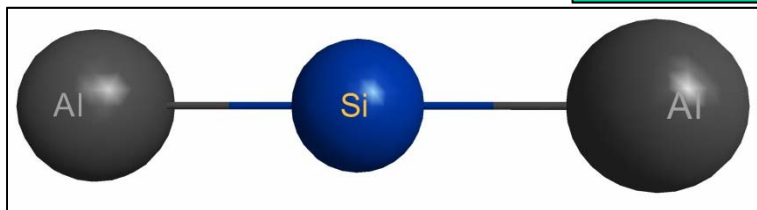


Formation of SiAl_2

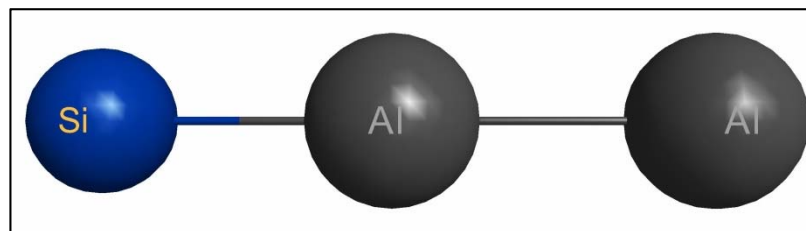


MRMP2(10e,12o)/aug-cc-pvtz level of theory

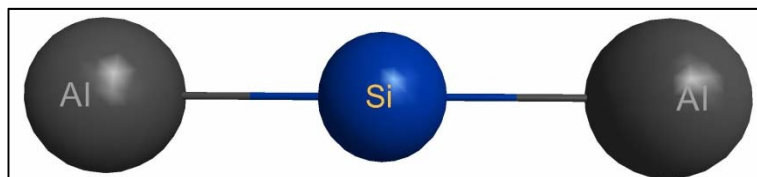
Linear isomers



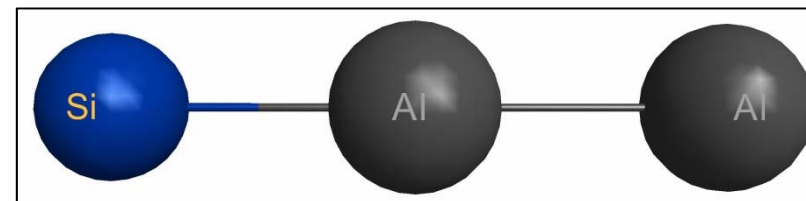
$^5\Pi_u$, -0.58 eV



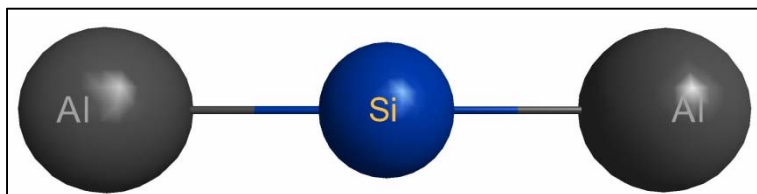
$^1\Pi$, -0.64 eV



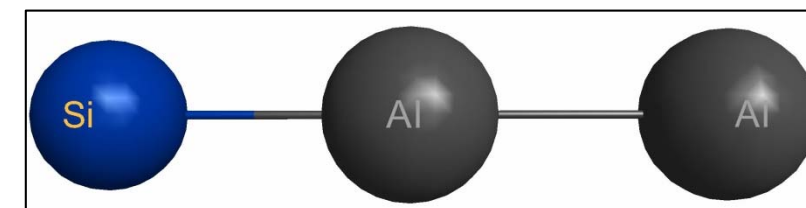
$^1\Sigma_g^+$, -0.89 eV



$^5\Pi$, -0.70 eV



$^1\Pi_u$, -1.28 eV



$^3\Pi$, -0.94 eV



Summary and Conclusions



- **Mg/Cu core-shell nanoclusters**
 - Helium droplet experiments show inversion of Cu_xMg_y clusters to Mg_yCu_x .
 - Cu atoms diffusing into Mg_{30} , and vice-versa, have been modeled using DFT.
 - Estimated barrier for Cu atoms to migrate into Mg_n is < 1 kcal/mol.
 - Estimated barrier for Mg atoms to migrate into Cu_n is 6 kcal/mol.
 - Calculations are consistent with observed Cu/Mg inversion.
- **SiAl_n clusters**
 - SiAl_2 has multiple local minima which are more stable than ground state SiAl ($^4\Sigma^-$) + Al (2P)
 - Cyclic (C_{2v} and C_s) and linear ($D_{\infty h}$ and $C_{\infty v}$)
 - Singlet, triplet, quintet states
 - At long SiAl -----Al separations, preferred approach is linear
 - Al-Si-Al: $^5\Pi_u$ can form without a barrier
 - Si-Al-Al: $^5\Pi$ can form without a barrier
 - Barriers for linear \leftrightarrow cyclic isomerizations TBD
 - In helium droplet environment, linear quintet states may be formed.



Acknowledgements



Core-shell nanoparticles

- Dr. Robert Buszek (ERC, Inc.)
- Dr. Sam Emery(NSWC-IH), Y. Xin (Florida State Univ.),
C.J. Ridge, B.K. Little, C.M. Lindsay (AFRL/RW)
J.M. Boyle (Dublin School, Dublin, New Hampshire)

AFOSR

Dr. Mike Berman

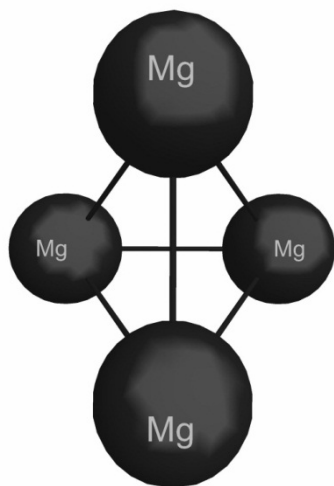
DoD HPCMP



Backup Slides



Mg_n benchmark calculations



Mg_n clusters

“Closed shell” atomic configuration $[(1s)^2(2s)^2(2p)^6(3s)^2]$ suggests that weak dispersion interactions will be important. Need to consider

- core-core and core-valence correlation
- correlation method (MP2, CC, DFT)
 - “active” electrons to be correlated in MP2, CC
 - suitable DFT functional for larger Mg_n clusters (up to $n \approx 100$)

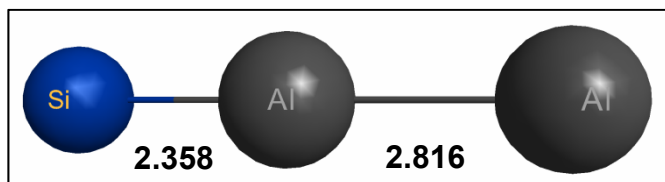
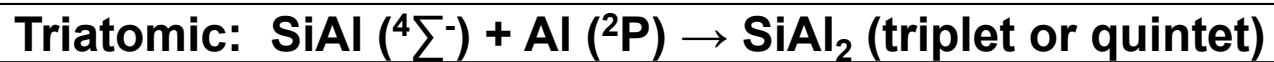
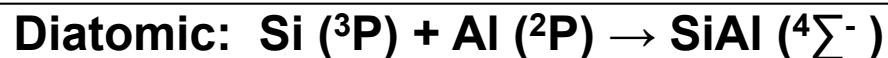
| Method | cc-pwCVDZ | cc-pwCVTZ | cc-pwCVQZ |
|------------|--------------|--------------|--------------|
| MP2 | 23.1 / 3.042 | 28.5 / 3.013 | 29.1 / 3.011 |
| CCSD(T) | 16.7 / 3.100 | tbd / 3.064 | tbd / 3.065 |
| DFT/B3PW91 | 26.5 / 3.092 | 26.5 / 3.091 | 26.4 / 3.092 |
| DFT/PBE | 34.6 / 3.070 | 34.5 / 3.070 | 34.3 / 3.070 |
| DFT/PBE0 | 31.5 / 3.078 | 31.5 / 3.078 | 31.4 / 3.078 |
| DFT/M06 | 30.8 / 3.028 | 30.4 / 3.025 | TBD / TBD |
| DFT/M11 | 19.2 / 3.134 | TBD / TBD | TBD / TBD |

Calculated binding energies used to determine size of helium droplet needed for evaporative cooling

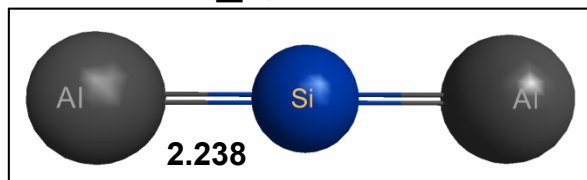


$SiAl_n$

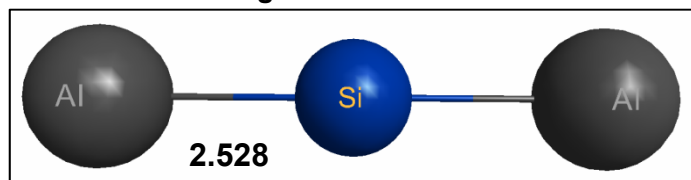
Calculate potential energy surfaces of stepwise atomic addition reactions;
e.g., $SiAl_n + Al \rightarrow SiAl_{n+1}$



$^3\Sigma^-$, -1.28 eV

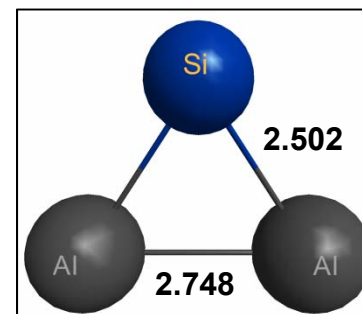


$^1\Sigma_g^+$, -1.70 eV

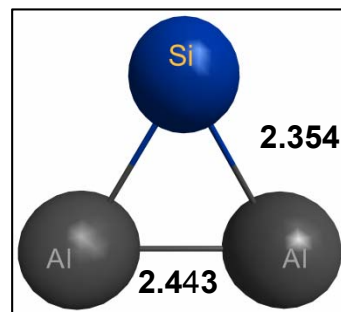


$^3\Sigma_g^-$, -1.93 eV

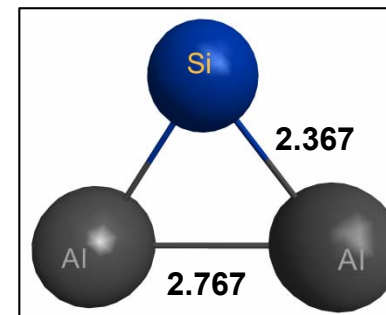
ZAPT(2)/aug-cc-pvtz level
of theory



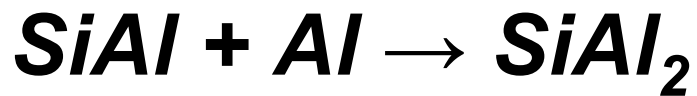
5A_2 , -1.77 eV



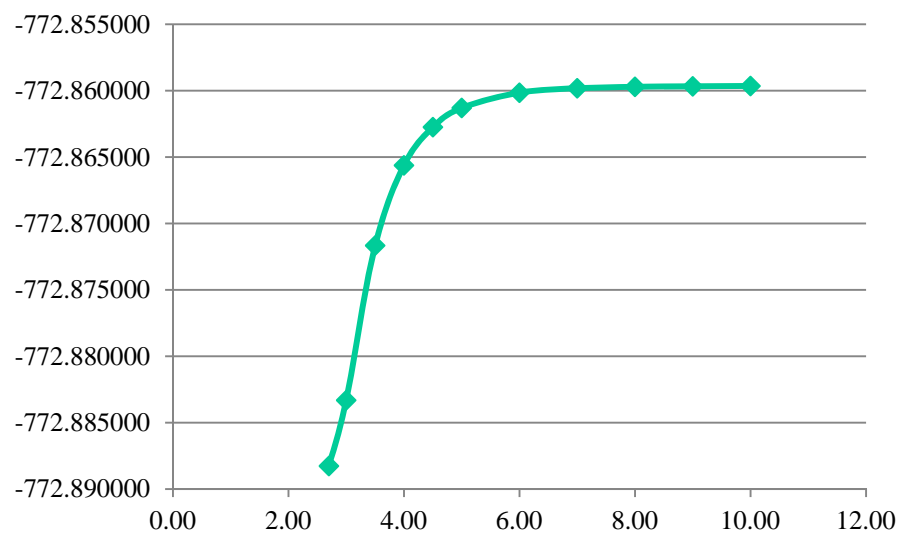
1A_1 , -2.65 eV



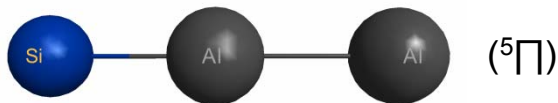
3B_2 , -2.67 eV



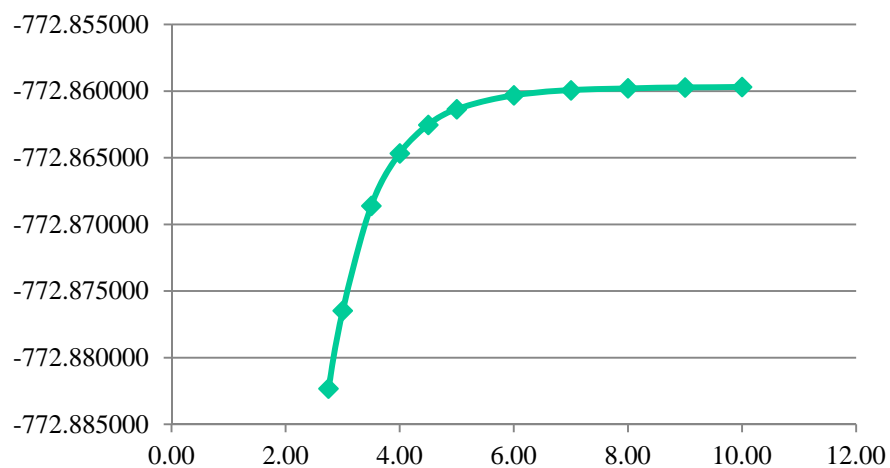
SiAl ($^4\Sigma$) + Al (2P)



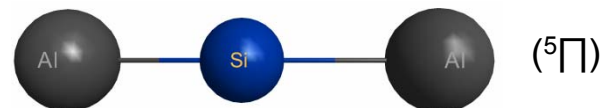
R(Al-Al)



AlSi ($^4\Sigma$) + Al (2P)



R(Si-Al)



MRMP2(4e,5o)/aug-cc-pVTZ