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Summer Interdisciplinary Research Symposium

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Summer 7-29-2019

#### Propane Dehydrogenation Using Transition Metal Cluster Catalysts

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#### **Recommended** Citation

Bradash, Sarah E.; Hoisington, Andrew D.; Zygmunt, Stan A.; and Fadel, Louis, "Propane Dehydrogenation Using Transition Metal Cluster Catalysts" (2019). *Summer Interdisciplinary Research Symposium*. 58. https://scholar.valpo.edu/sires/58

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# Propane Dehydrogenation Using Transition Metal Cluster Catalysts Sarah Bradash, Andrew Hoisington\*, Louis Fadel\*, Stan Zygmunt Valparaiso University Department of Physics and Astronomy, \*Ivy Tech Community College

#### Abstract

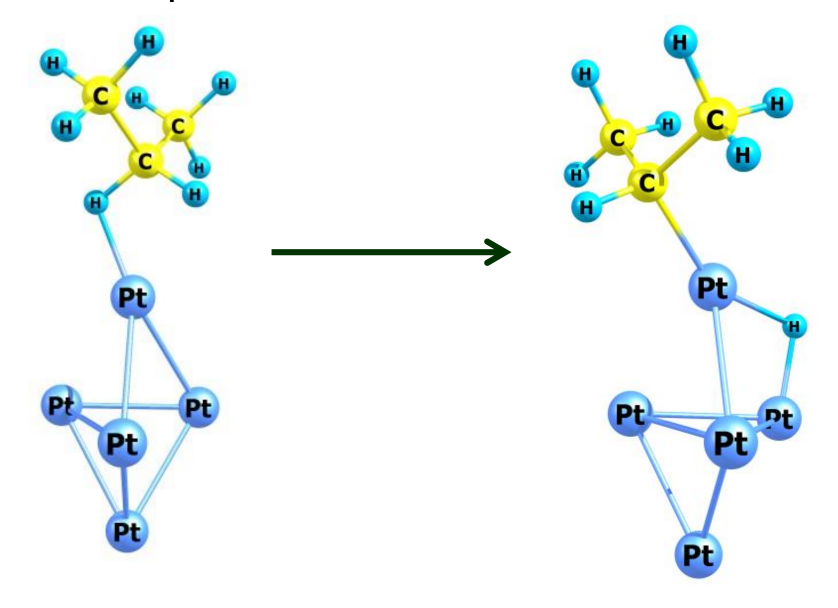
Our research seeks to determine the propane dehydrogenation (PDH) reaction pathways using various transition-metal cluster catalysts. We are studying the first step of the reaction, in which a C-H bond is broken. This has been previously shown to be the ratelimiting step of the PDH reaction. We are calculating the PDH activation energy (E<sub>a</sub>) using the Vienna Ab-Initio Simulation Package (VASP) in conjunction with the nudged elastic band algorithm. Thus far, we have studied Pt, Ta, and Ni clusters ranging in size from 2-10 atoms. Our goal is to better understand the dependence of  $E_a$  on metal type and cluster size.

#### **Computational Procedure**

- Use Vienna Ab-Initio Simulation Package (VASP) which implements density functional theory (DFT) to optimize the reactant and product structures
- Use the nudged elastic band (NEB) algorithm to calculate reaction pathways for the first step of PDH reaction: C-H bond cleavage
- Calculate E<sub>a</sub> for each TM by subtracting the initial energy from the transition state energy

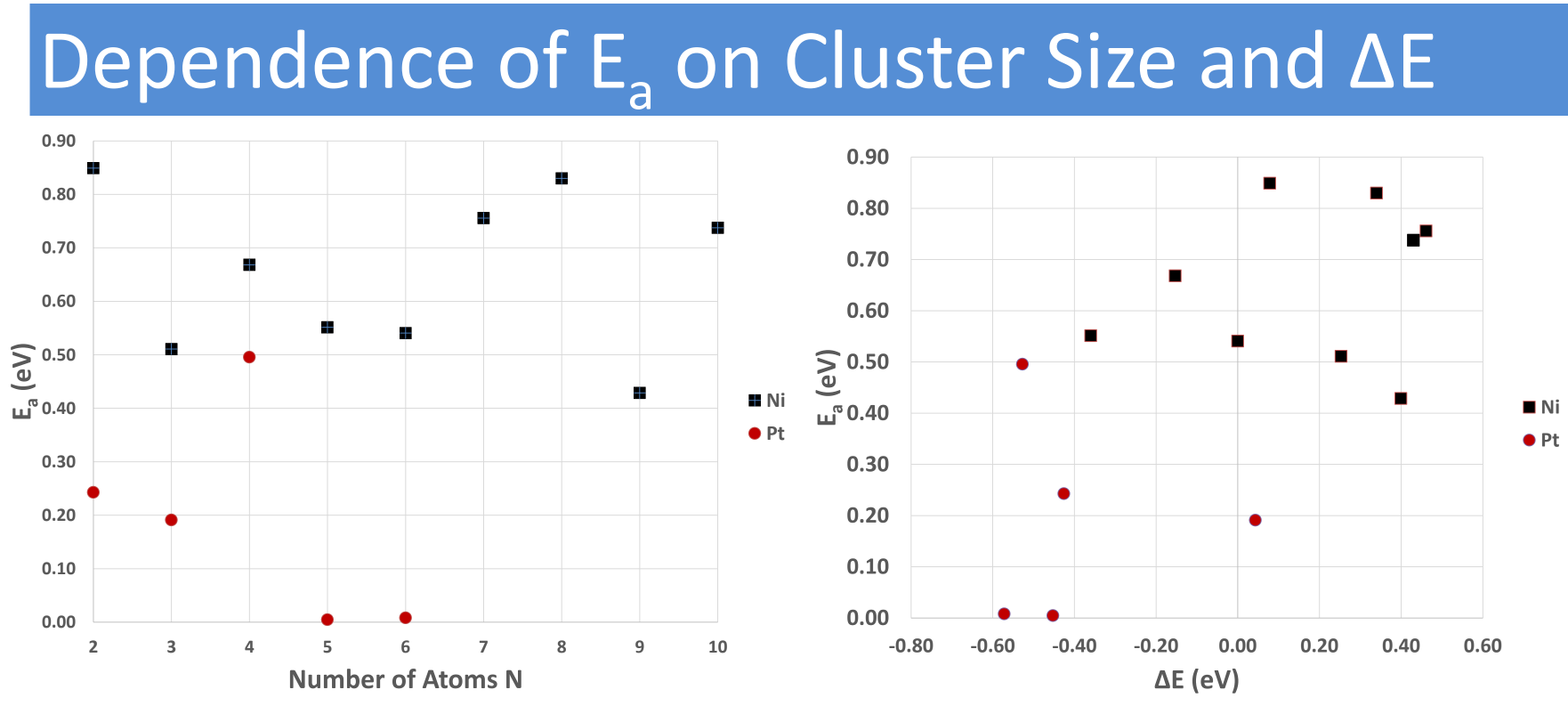
#### NEB Algorithm

- VASP implements DFT and the NEB algorithm to calculate minimum energy path between known reactants and products
- The NEB algorithm functions by adding spring forces along the band between images and by projecting out the component of the force due to the potential perpendicular to the band
- We use a total of 7 images (5 intermediate structures) to model the reaction path

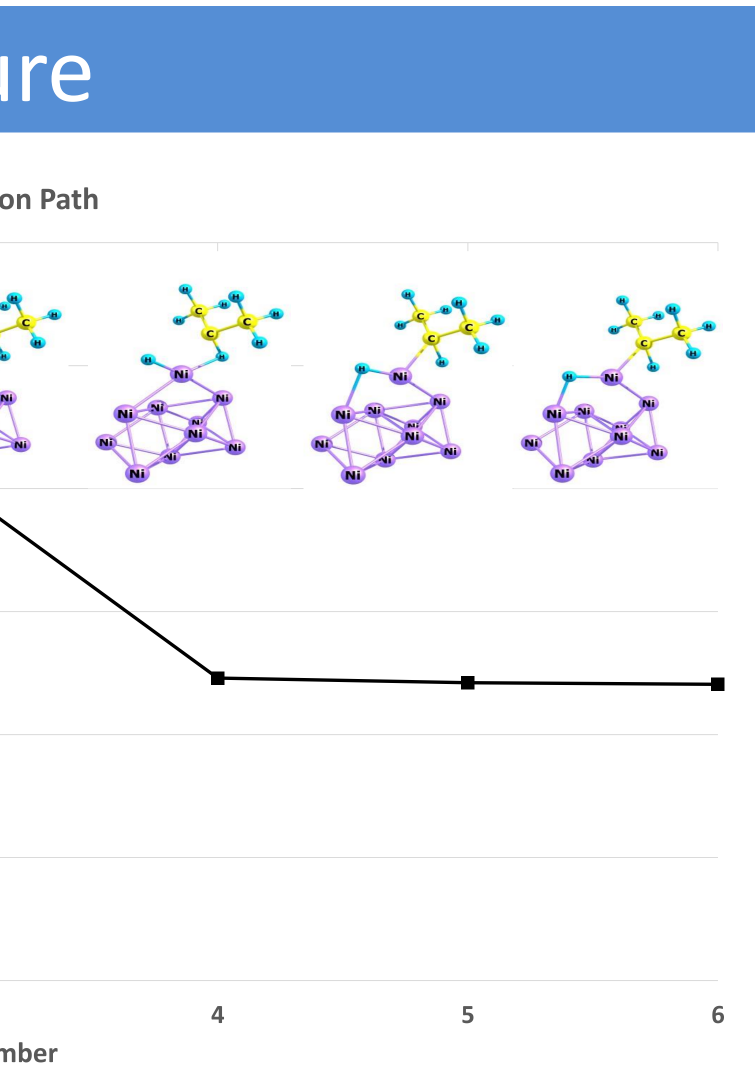


### **NEB Calculation Procedure**

- Ni<sub>10</sub> + C<sub>3</sub>H<sub>8</sub> NEB Reaction Path -93.10 -93.70 -93.90 -94.10 Image Number
- Each structure was initially optimized as an isolated TM cluster, then a propane molecule was added to begin NEB calculation
- Each NEB calculation maps the energetic pathway of the first step of the PDH reaction (C-H bond cleavage)
- Our goal: calculate E<sub>a</sub> and investigate correlations with reaction energies and TM characteristics



E<sub>a</sub> varies according to cluster size, but not in a linear fashion No linear relation between  $E_{a}$  and  $\Delta E$ ; Bell-Evans-Polanyi (BEP) principle does not hold for these results

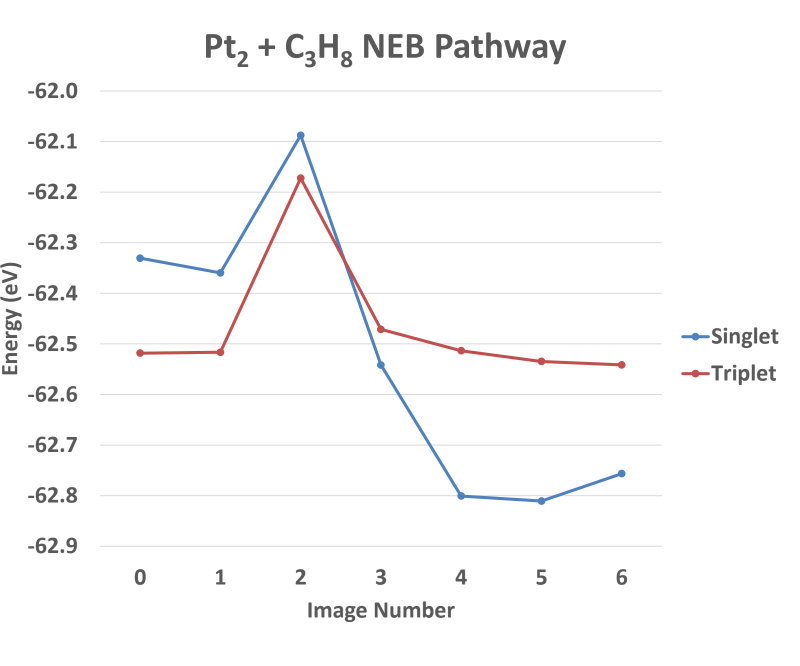


### Graphene-Supported Pt Clusters

Cluster	∆E (eV)	E <sub>a</sub> (eV)	Pt-Pt (Å)
Pt	0.15	0.75	
Pt <sub>2</sub>	-0.06	0.51	2.362
Pt <sub>3</sub>	-0.95	0.46	2.503 <mark>,</mark> 2.587
Pt <sub>4</sub>	-0.16	0.08	2.580, 2.642, 2.773
Pt <sub>6</sub>	-0.34	0.45	2.491, 2.606, 2.660

- No clear trend between  $E_a$  and  $\Delta E$
- E<sub>a</sub> depends on cluster size and is a minimum for Pt<sub>4</sub>
- Local geometry of interacting Pt

# Spin State Crossing



### Conclusions

- contrary to BEP principle

## Acknowledgements

- Indiana Space Grant Consortium
- Prof. Zygmunt



atom may influence catalytic activity

- Spin state crossing occurs in some structures
- Position at which the crossing occurs is currently unknown

E<sub>2</sub> is dependent on spin state, metal type, and size of cluster • Spin state crossing occurs for some clusters • No clear dependence of  $E_a$  on  $\Delta E$  exists for both Ni and Pt,

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