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# Electrochemical Reduction of CO2 using Cu-Pd clusters on Graphene

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# ELECTROCHEMICAL REDUCTION OF CO, USING **Cu-Pd CLUSTERS ON GRAPHENE**

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## Motivation

Over the past decade, the amount of  $CO_2$  in the atmosphere has increased drastically, creating adverse climate effects. We must ensure that the level of  $CO_2$  in the atmosphere doesn't increase

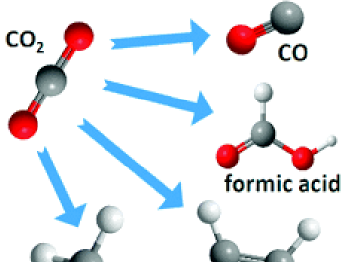




## **Computational Method**

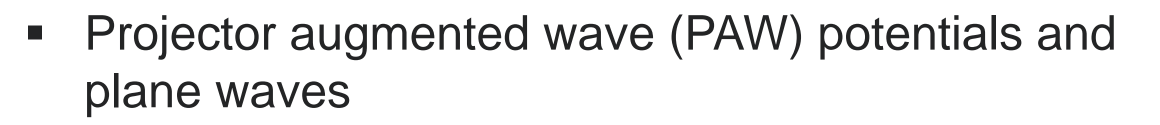
Five Cu-Pd alloy clusters were screened for the electrochemical CO<sub>2</sub> reduction based on density functional theory (DFT)

- Vienna Ab-initio Simulation Package (VASP)
- PBE exchange—correlation functional
- van der Waals (vdW) interaction (D2 method)

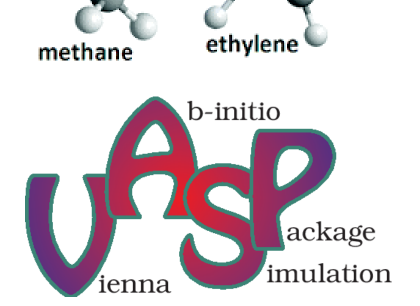




One way of preventing this increase, is to convert CO<sub>2</sub> into useful hydrocarbons such as methane. Current catalysts for this conversion are not efficient, and as such our group is screening for potential new catalysts for this conversion.

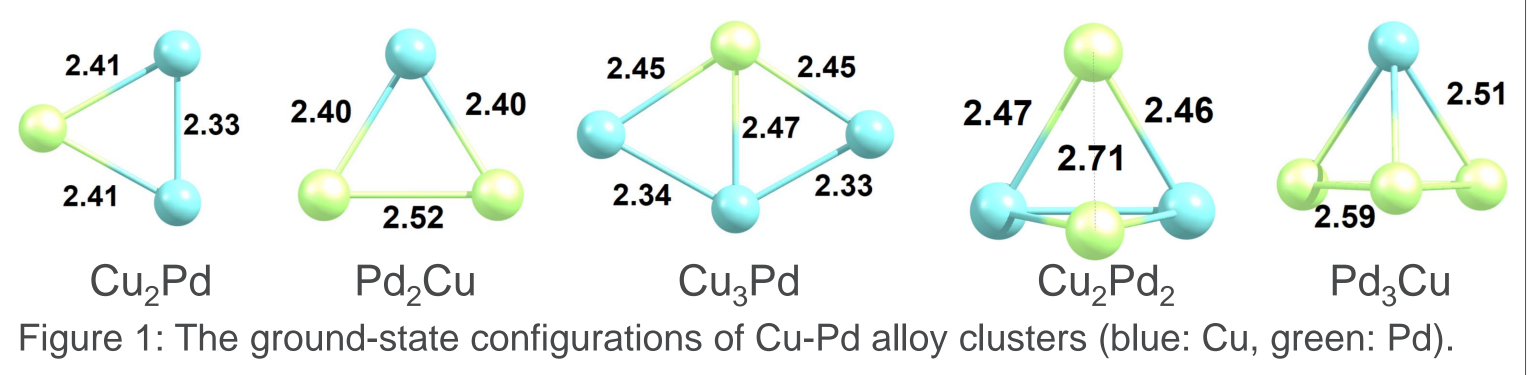


Free energies were calculated based on the computational hydrogen electrode model



# **Optimizing Clusters**

First, the cluster geometry in gas phase was optimized to find the stable state.



The clusters were then placed on a single vacancy graphene support. Different binding sites and orientations were tested.

# Screening

Free energies of the first hydrogenation steps of competing reactions was compared.

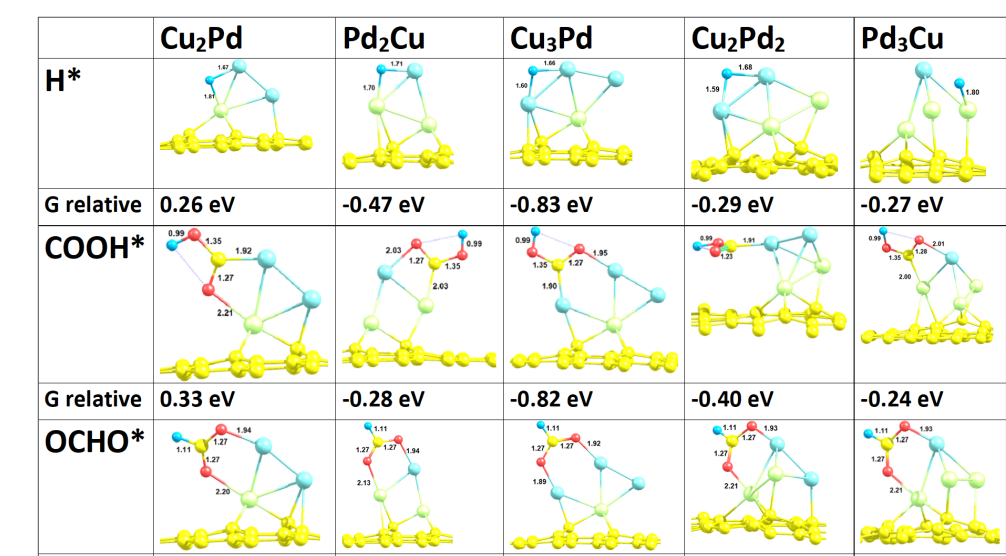
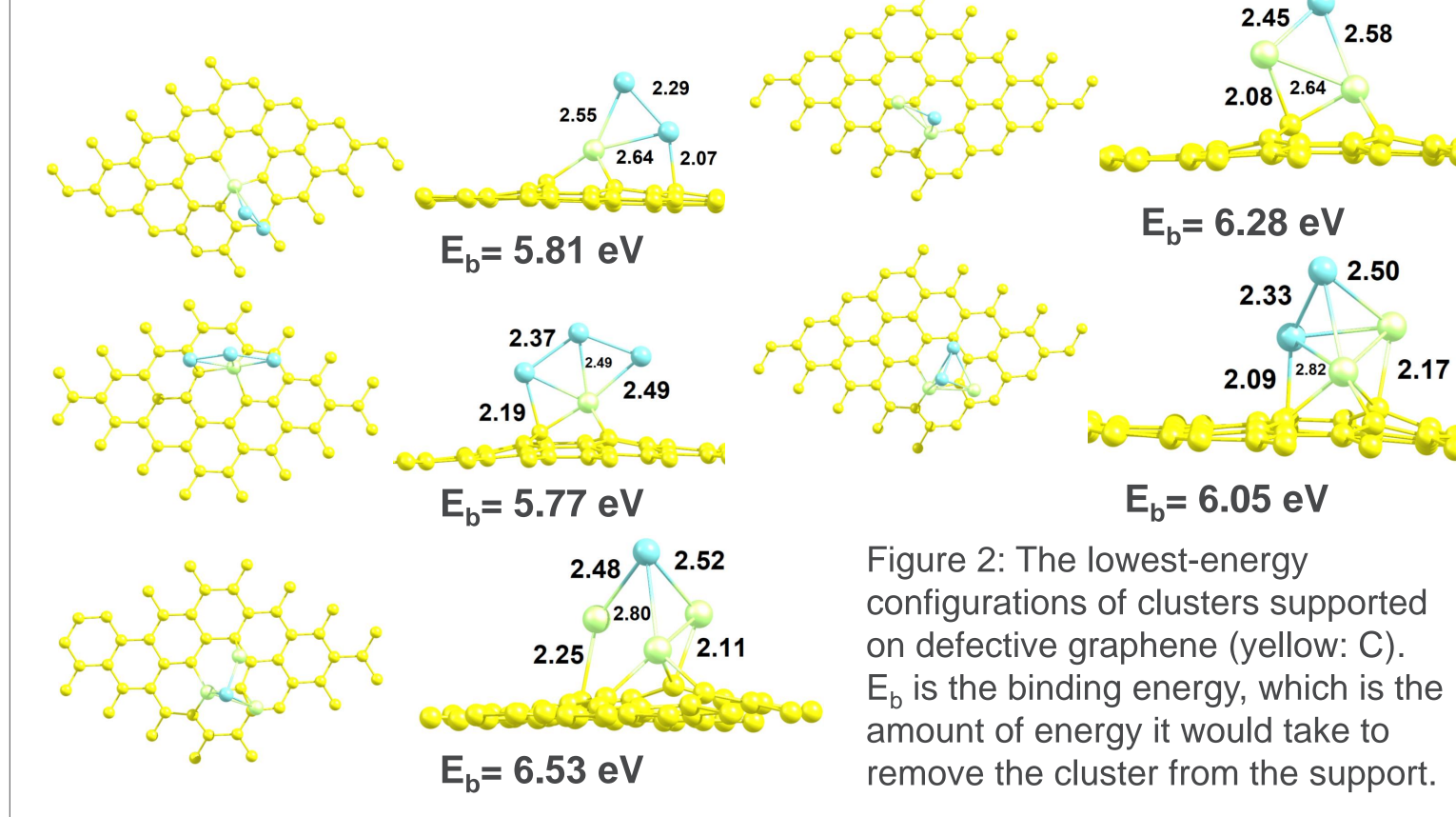


Figure 3: The optimized configurations of the first hydrogenation for the reduction of  $CO_2$  and the production of  $H_2$  (red: O, small blue: H). The relative energy associated with each of the configuration is the free energy of the configuration with respect to the sum of energies of reactants.



All the clusters are found to bind strongly to the vacancy site and are very unlikely to move and aggregate.

#### **Results and Discussion**

Elementary limiting potentials and overpotentials (in V)

G relative -0.59 eV -1.12 eV -1.59 eV -0.99 eV -0.91 eV

- In all cases the formation of OCHO\* is preferred to the formation of H\*
- CO<sub>2</sub> reduction is more favorable than production of hydrogen gas

Most probable rate limiting steps to CH<sub>4</sub>  $OH^* + H^+ + e^- \rightarrow H_2O$ OH\* ----- $CO_2 + H^+ + e^- \rightarrow OCHO^*$ CHO<sup>\*</sup>  $OCHO^* + H^+ + e^- \rightarrow HCOOH^*$  $HCOOH^* \rightarrow CHO^* + H_2O$ 

The optimal binding configuration for all of the intermediates for possible rate

limiting steps were found. Cu<sub>2</sub>Pd Pd<sub>2</sub>Cu **Cu**<sub>3</sub>Pd  $Cu_2Pd_2$ Pd₃Cu HCOOH\* Figure 4: The optimized CHO\* configurations for the possible rate limiting steps of the reduction of CO<sub>2</sub> OH\*

- Carbon prefers to bond to Pd, and oxygen prefers to bond to Cu
- The bonding of C and O also depends on an atom's coordination number

|                        | Cu <sub>2</sub> Pd | Pd <sub>2</sub> Cu | Cu <sub>3</sub> Pd | Cu <sub>2</sub> Pd <sub>2</sub> | Pd <sub>3</sub> Cu |
|------------------------|--------------------|--------------------|--------------------|---------------------------------|--------------------|
| -U <sub>L</sub> [OH*]  | 0.30               | 0.75               | 1.53               | 0.95                            | 0.98               |
| -U <sub>L</sub> [CHO*] | 0.76               | 0.92               | 1.29               | 0.64                            | 0.58               |
| Overpotential          | 0.93               | 1.09               | 1.70               | 1.12                            | 1.15               |

- Rate limiting step: CHO\* formation for triatomic clusters OH\* desorption for other clusters
- The cluster that appears to make the best catalyst is Cu<sub>2</sub>Pd

## **Future Direction**

- The electronic structure of the Cu<sub>2</sub>Pd cluster on graphene will be analyzed to better understand the bonding that occurs during the reaction
- More clusters doped on graphene will be screened to determine if they could be good catalysts for the electrochemical reduction of CO<sub>2</sub>

#### References

Rapid growth in carbon dioxide emissions breaks in 2015." CSIRO. https://www.csiro.au/en/News/News-releases/2015/Rapid-growth-in-carbon-dioxide-emissions-breaks-in-2015.



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