# **Computational Designing and Screening of Solid Materials for CO<sub>2</sub>** Capture

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**Research &** Innovation Center



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- > Due to the environmental issues that the world faces today, there are significant interests to develop materials capable to capture CO<sub>2</sub> with optimal performances.
- Solid materials are potential candidates for CO<sub>2</sub> sorbents. By combining the database mining with ab initio thermodynamic calculation, we implemented a novel theoretical methodology to screen solid sorbents from known material databank and to synthesize new materials with improved CO<sub>2</sub> capture capabilities for further experimental validations.

### **Theoretical Methods**

For the reaction of a solid to absorb CO<sub>2</sub>([...] are optional):

solid sorbent + CO<sub>2</sub>  $\pm$  [H<sub>2</sub>O]  $\rightarrow$  sorbent CO<sub>2</sub> + [solid]

The chemical potential ( $\Delta \mu$ ) of the reaction can be calculated as:

$$\Delta \mu(\mathbf{T},\mathbf{P}) = \Delta \mu^0(\mathbf{T}) - \mathbf{RT}(\ln \mathbf{P}_{\rm CO_2} \pm \ln \mathbf{P}_{\rm H,O})$$

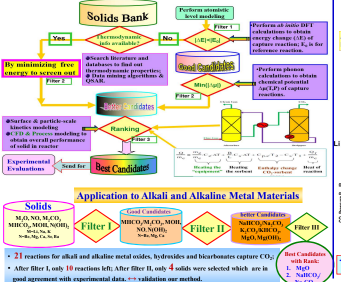
A. Intensive search for thermodynamic properties of solids from known databases and literatures. If all of them are known, obtain the better candidates by minimizing free energies based on operating conditions. B. If their thermodynamic properties are unknown, calculate by

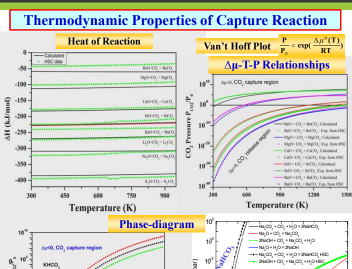
$$\begin{array}{l} \Delta \mu^{0}(T) \approx \Delta E^{DPT} + \Delta E_{ZPE} + \Delta F_{PH}(T) - (G_{CO_{1}}(T) \pm G_{H,O}(T)) \\ \Delta H(T,P) = \Delta \mu(T,P) + T(\Delta S_{PH}(T,P) - S_{CO_{1}}(T,P) \mp S_{H,O}(T,P)) \end{array}$$

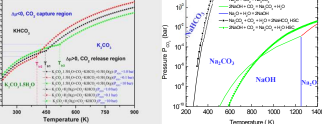
•  $G_{CO_2}$ ,  $S_{CO2}$ , and  $G_{H_{2O}}$  and  $S_{H_{2O}}$  evaluated from statistical mechanics;

- where  $\Delta \mathbf{E}^{DFT}$  are calculated by density functional theory;
  - $\Delta E_{ZPE}$ ,  $\Delta S_{PH}(T)$  and  $\Delta F_{PH}(T)$  evaluated through phonon dispersions

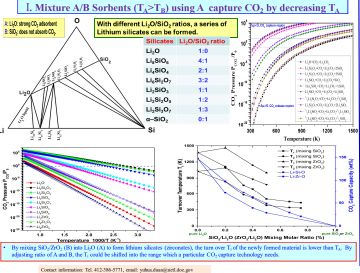
## **High-Throughput Screening Methodology**

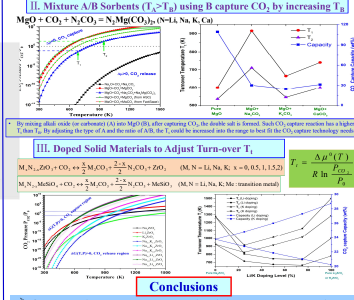






#### Synthesis New Sorbents to Fit Operation Needs





- Our theoretical approach can obtain similar results as experimental measurements and can be used to identify better candidates.
- The strength of our method is to screen complex sorbent materials for which their thermodynamic properties are not available.
- > Hundreds of solid materials have been investigated. Now, we are working on screening of multi-components, substituted, doped, and mixed materials to search for good CO<sub>2</sub> sorbents.
- By mixing/doping different solids, we can theoretically synthesize new materials which may fit the industrial operating conditions with optimal CO<sub>2</sub> capture performance.
- $\triangleright$ As a long time goal, we'll build a database of solid sorbents for CO<sub>2</sub> capture to satisfy industrial operational requirements.

#### Publications

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