

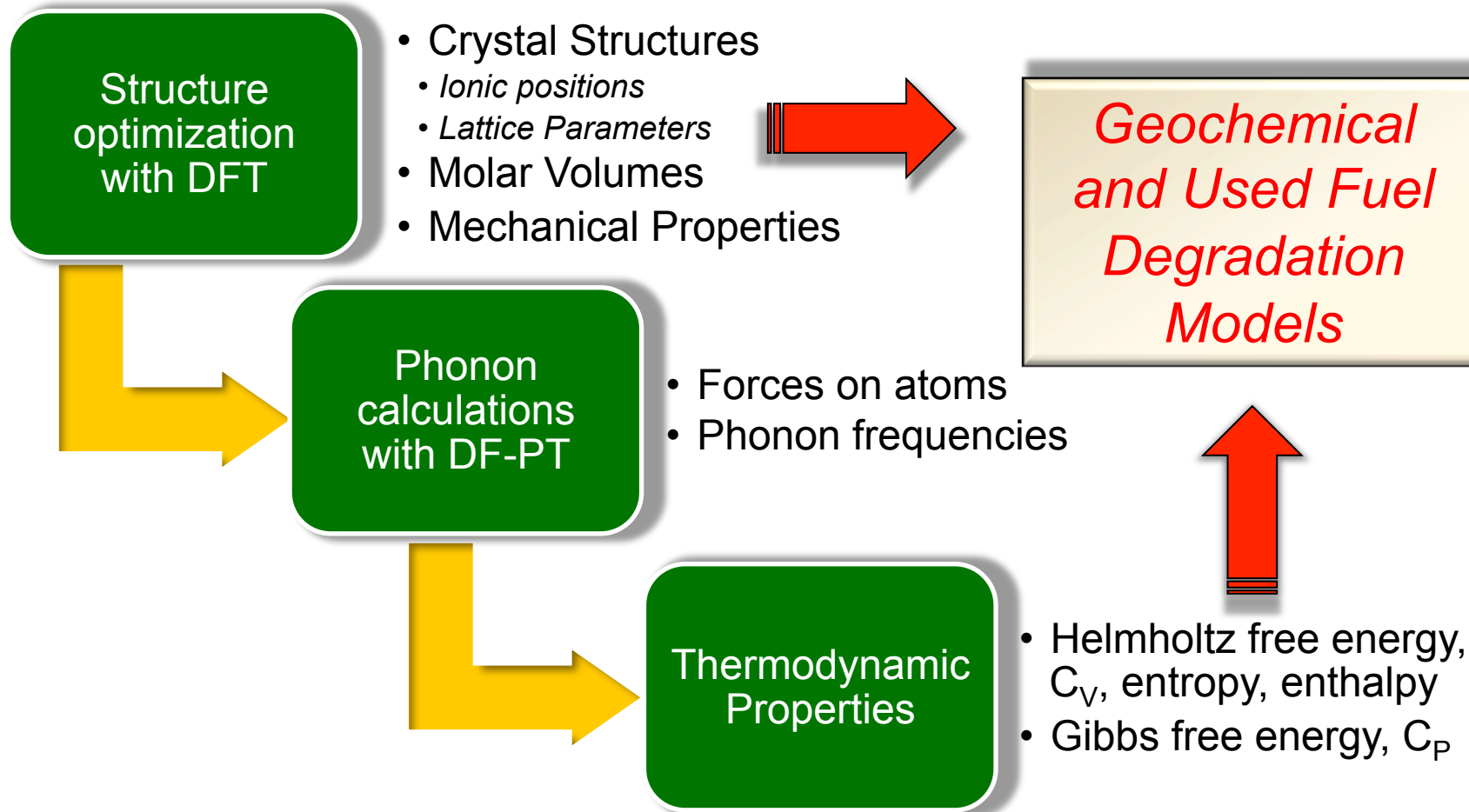
# Used Fuel Disposition Campaign

## Thermal-Mechanical Data of Argillite and Crystalline Disposal Systems: A First-Principles Approach

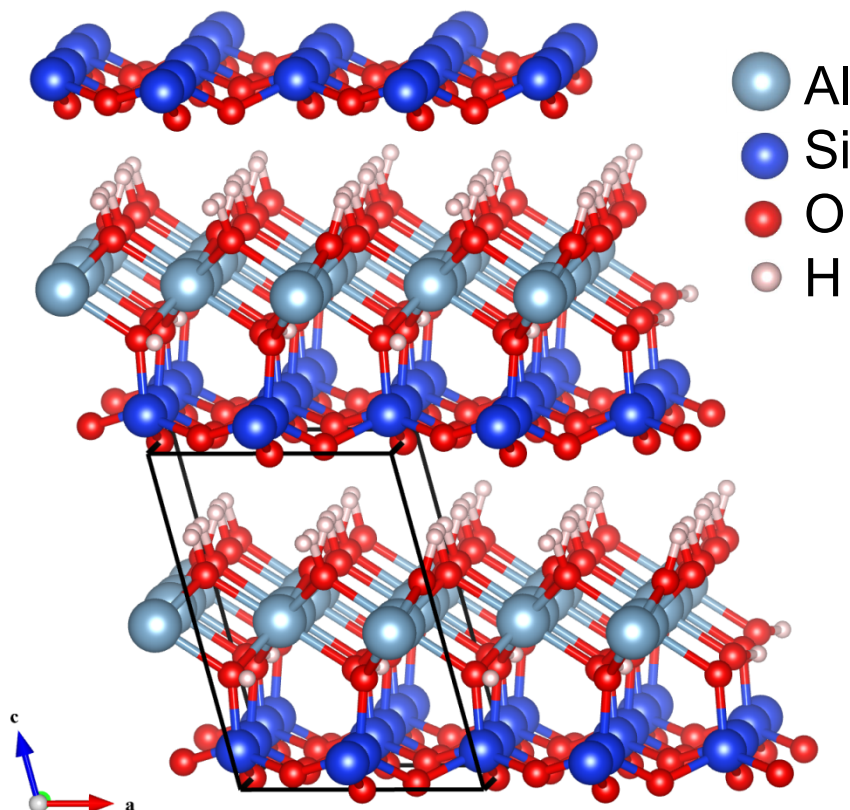
**Philippe F. Weck, Carlos F. Jove-Colon**  
**Sandia National Laboratories**

Used Fuel Disposition Working Group Meeting  
Las Vegas, June 4-6, 2014

- **Thermal-mechanical data for natural system (NS) minerals and engineered barrier systems (EBS) materials** are critical to assess their stability and behavior in geologic disposal environments for safety assessments.
- **Thermodynamic Data Gaps and Research Needs:** NS minerals surrounding the waste package (e.g. clays, complex salts, granite...).
- **Objectives:** Using parameter-free first-principles methods to:
  - Calculate missing thermodynamic data needed for geochemical & SNF degradation models, as a **fast and systematic way to predict materials properties and to complement experiments**.
  - Provide an **independent assessment** of existing experimental thermodynamic data and resolve contradictions in existing calorimetric data.
  - Validate our computational approach using high-quality calorimetric data.
- **Approach**
  - Structural optimization using density functional theory (DFT) [VASP code].
  - Use density functional perturbation theory (DF-PT) to calculate the phonon properties of materials relaxed with DFT and derive their thermal properties.



# Clays: Structure & Thermal-Mechanical Properties of Kaolinite ( $\text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4$ )



■ Expt.:  $V = 328.708 \text{ \AA}^3$

$a = 5.153 \text{ \AA}$ ,  $b = 8.942 \text{ \AA}$ ,  $c = 7.391 \text{ \AA}$   
 $\alpha = 91.93^\circ$ ,  $\beta = 105.05^\circ$ ,  $\gamma = 89.80^\circ$ .

■ Standard DFT:  $V = 340.11 \text{ \AA}^3$

$a = 5.21 \text{ \AA}$ ,  $b = 9.05 \text{ \AA}$ ,  $c = 7.48 \text{ \AA}$   
 $\alpha = 91.8^\circ$ ,  $\beta = 105.1^\circ$ ,  $\gamma = 89.7^\circ$ .

■ DFT + van der Waals correction  
(DFT-D2):  $V = 329.03 \text{ \AA}^3$

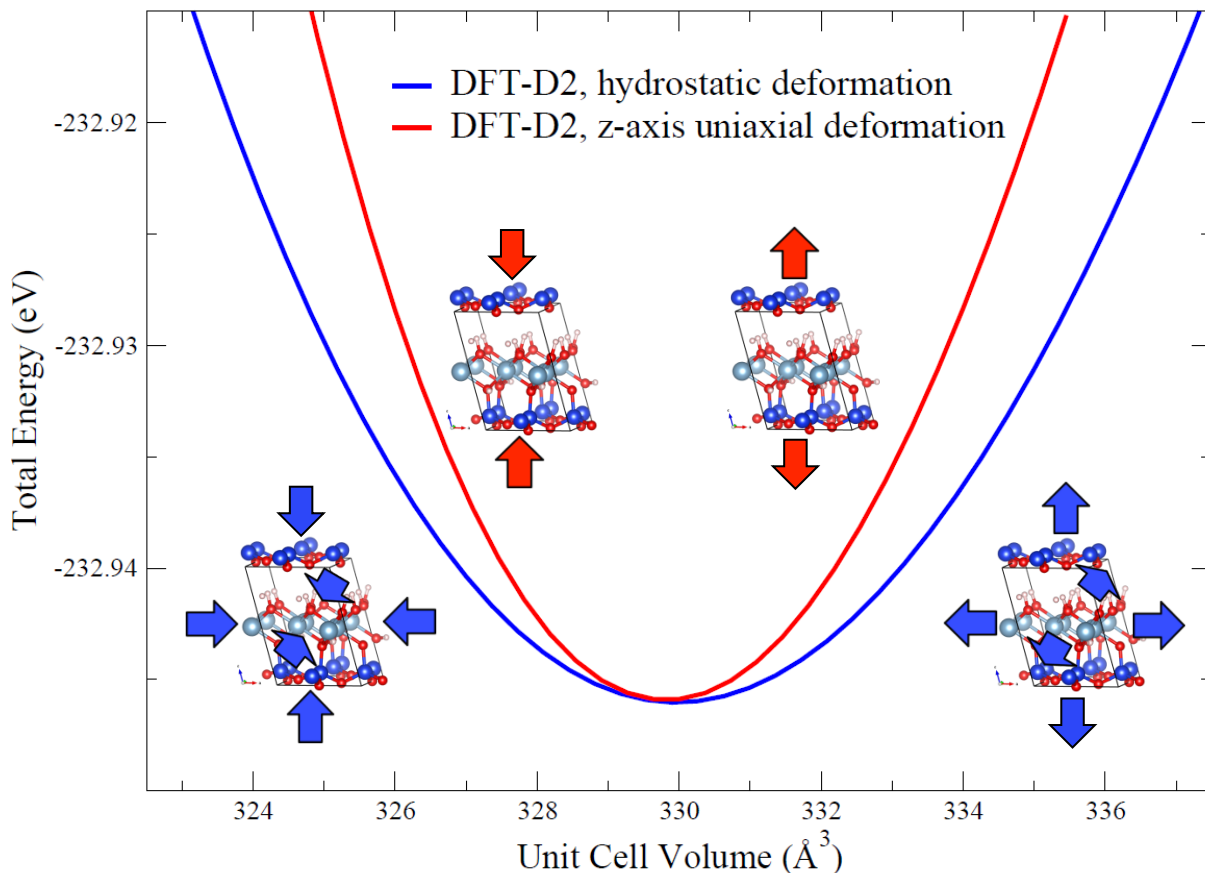
$a = 5.18 \text{ \AA}$ ,  $b = 8.99 \text{ \AA}$ ,  $c = 7.33 \text{ \AA}$   
 $\alpha = 91.6^\circ$ ,  $\beta = 105.1^\circ$ ,  $\gamma = 89.8^\circ$ .

Volume calculated with DFT-D2 overestimates experiment by less than 1%, while standard DFT overestimates expt. by ca. 3.5 %.

→ All the following calculations were carried out at the DFT-D2 level.

→ Important to use DFT corrected for dispersion interaction for layered systems such as clays.

# Clays: Structure & Thermal-Mechanical Properties of Kaolinite ( $\text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4$ )



Two types of cell deformation:

- **Hydrostatic deformation**
- **Uniaxial deformation** (along the z-axis normal of the layers)

**Bulk modulus** (Birch-Murnaghan 3rd-order equation of state):

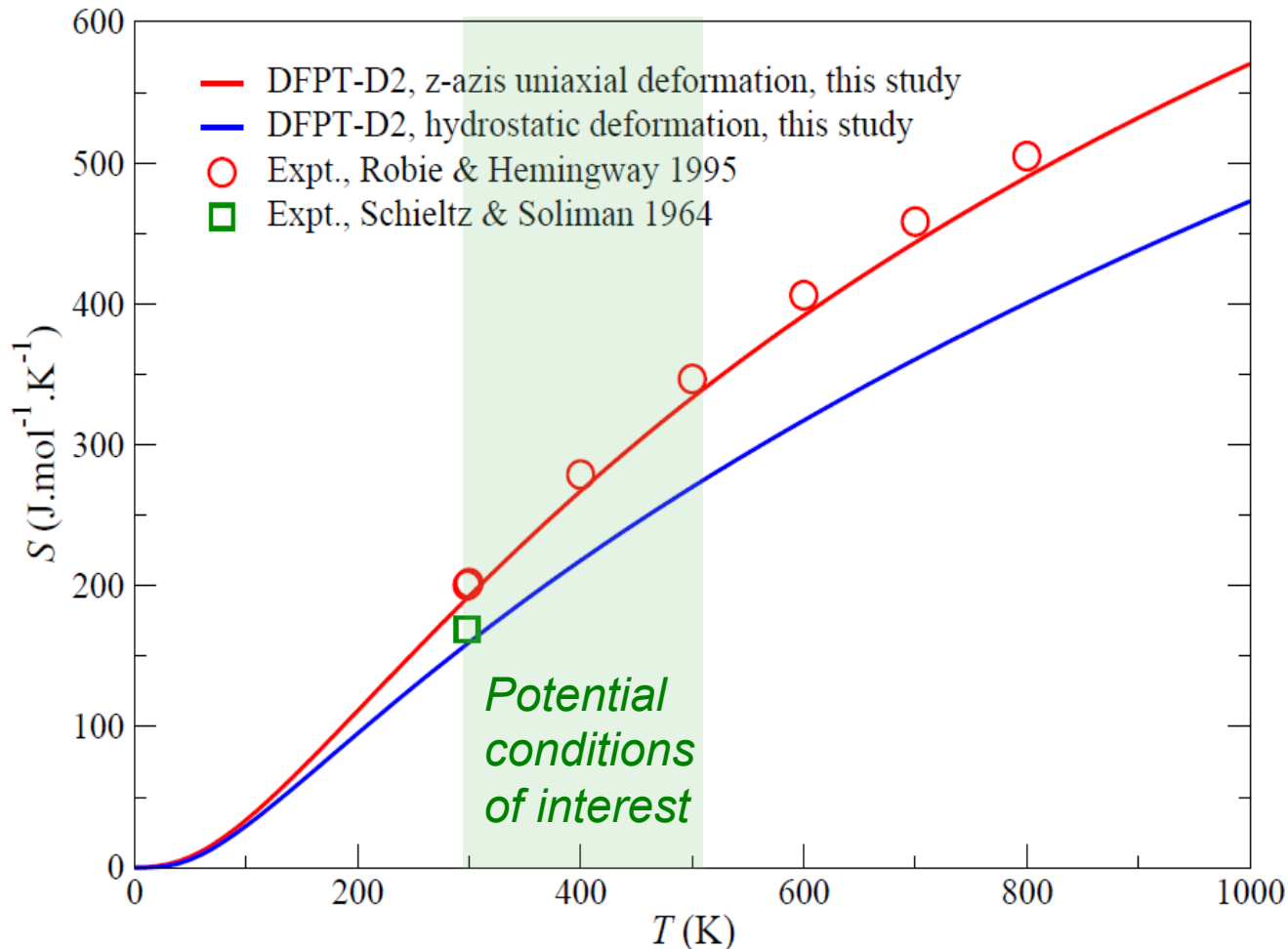
- **Expt.:**  $B_0 = 44\text{-}56$  GPa
- **DFT, hydrostatic:**  $B_0 = 54$  GPa
- **DFT, uniaxial:**  $B_0 = 116$  GPa

→ **Strong impact of the system anisotropy on the mechanical-elastic properties of the clays.**

$P$ : pressure;  $V_0$ : ref. volume;  
 $V$ : deformed volume.

$$B_0 = -V \left( \frac{\partial P}{\partial V} \right)_{P=0} \quad B'_0 = \left( \frac{\partial B}{\partial P} \right)_{P=0}$$

# Clays: Structure & Thermal-Mechanical Properties of Kaolinite ( $\text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4$ )



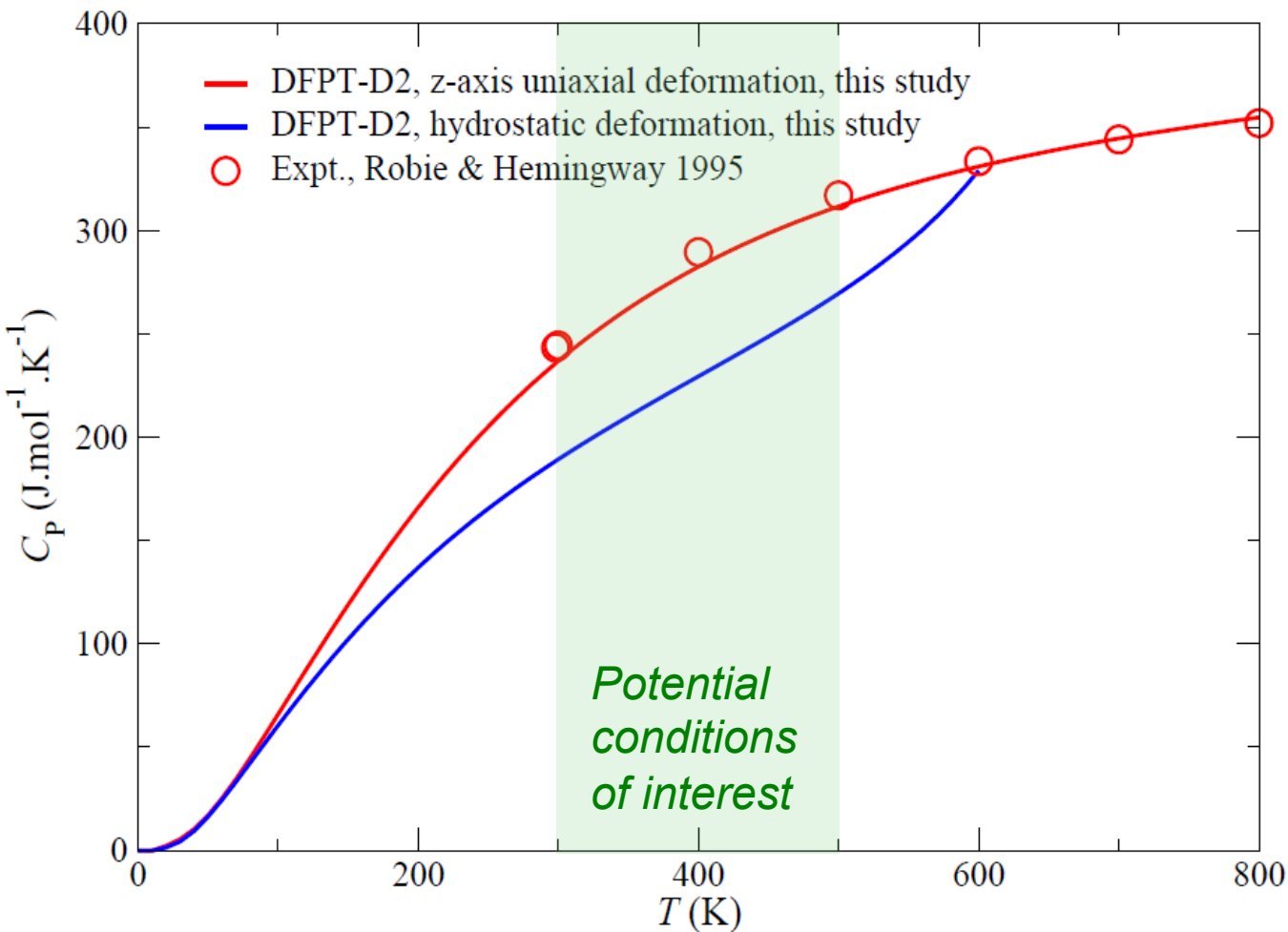
## Isochoric calculations:

- Calculated entropy using **uniaxial deformation** of the unit cell is in excellent agreement with experiment.
- Entropy computed using **hydrostatic deformation** always underestimates data derived from calorimetry.

→ **Uniaxial deformation of the cell maximizes the entropy compared to hydrostatic deformation.**

→ **Thermodynamic driving force for the formation of layered structures.**

# Clays: Structure & Thermal-Mechanical Properties of Kaolinite ( $\text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4$ )



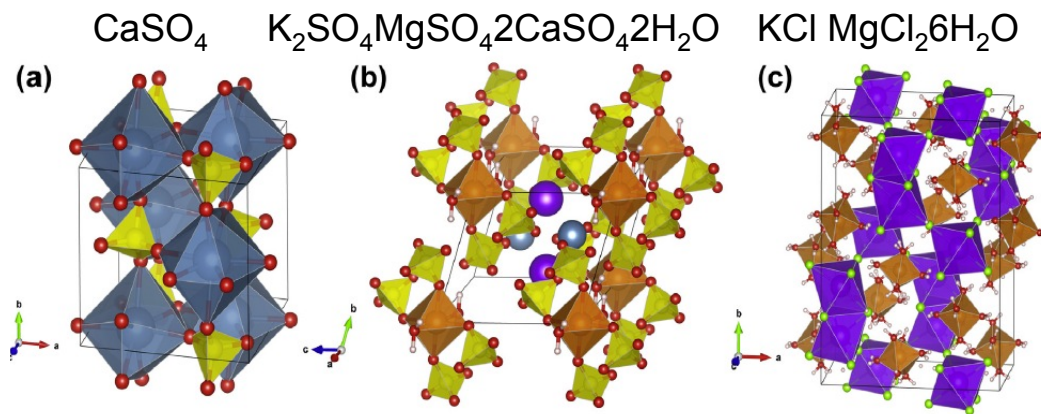
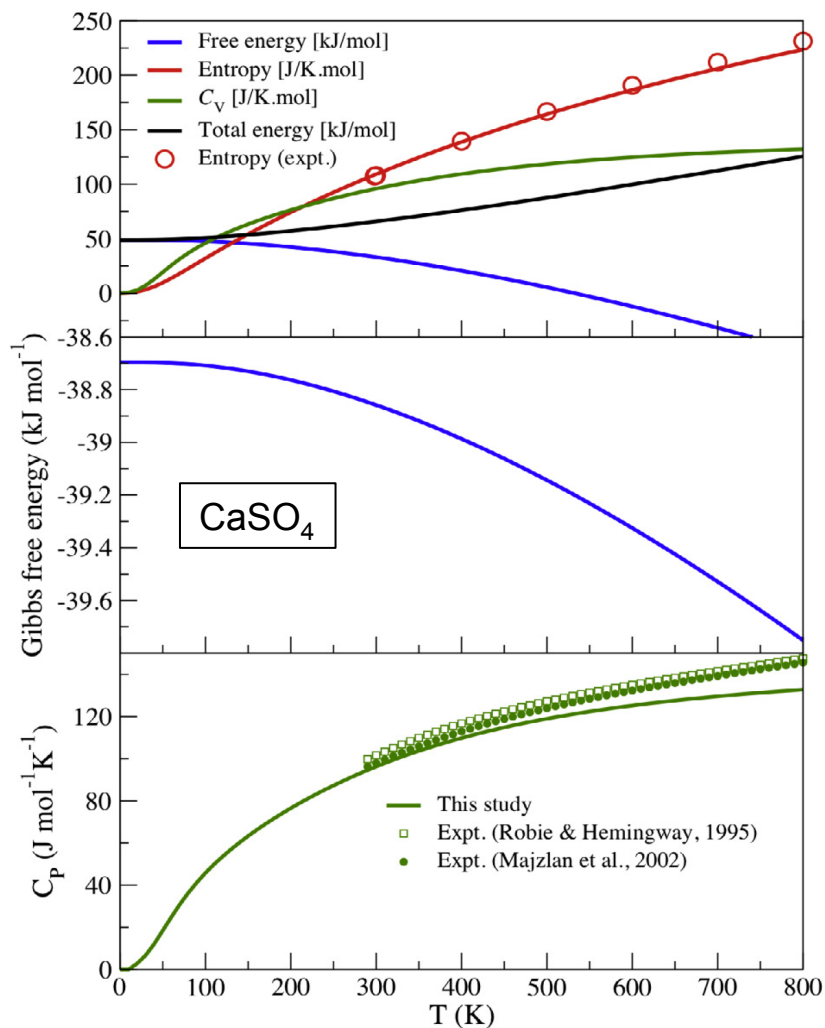
## Isobaric calculations:

- Calculations carried out within the quasi harmonic approximation (QHA), at a fixed pressure of 1 bar.
- Computed isobaric heat capacity using **uniaxial deformation** of the cell reproduces calorimetric data
- $C_p$  calculated with **hydrostatic deformation** underestimates calorimetric data for temperatures < 600 K.



# Used Fuel Disposition

## R&D Highlight on Crystalline Salts: Anhydrite, Polyhalite and Carnallite

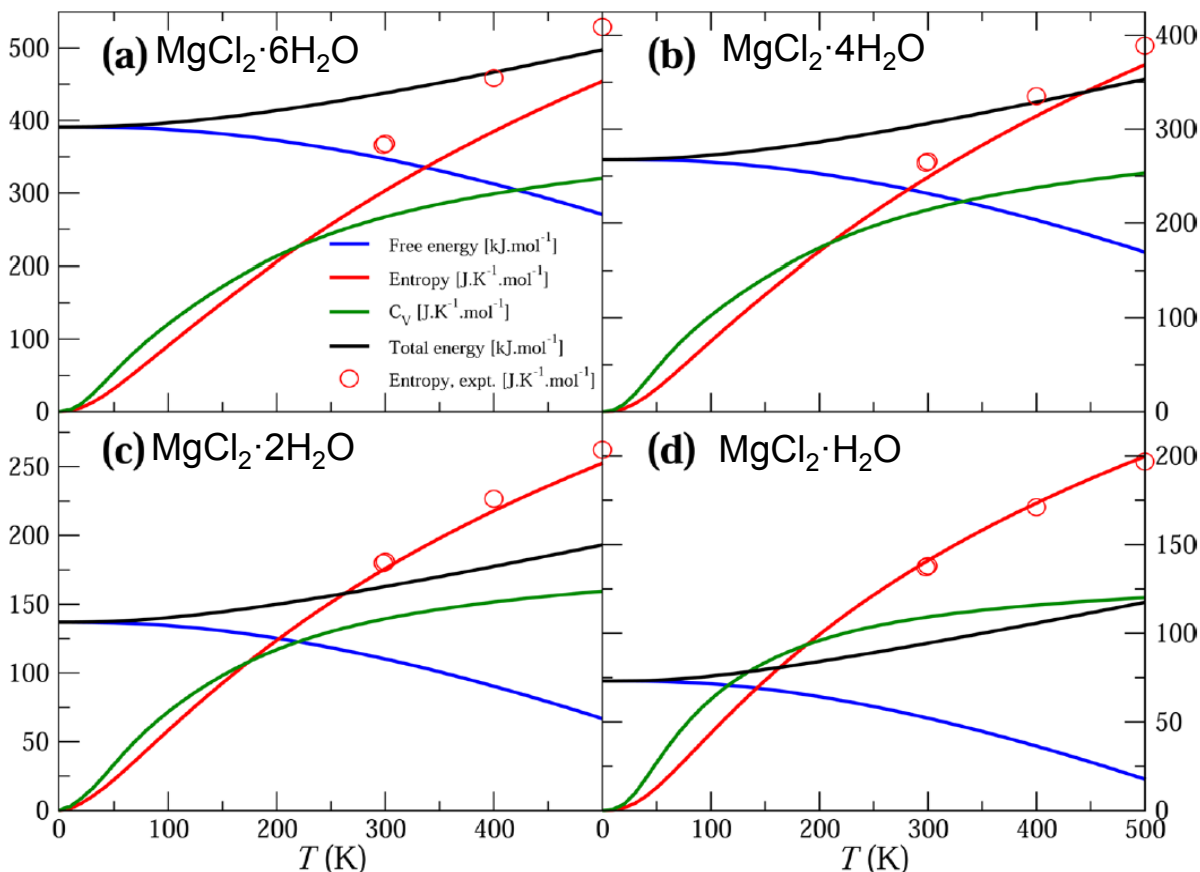


**Paper selected as Cover Page:**

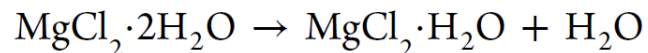
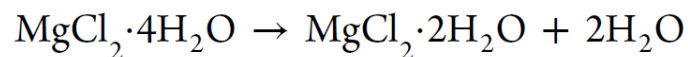
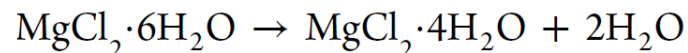
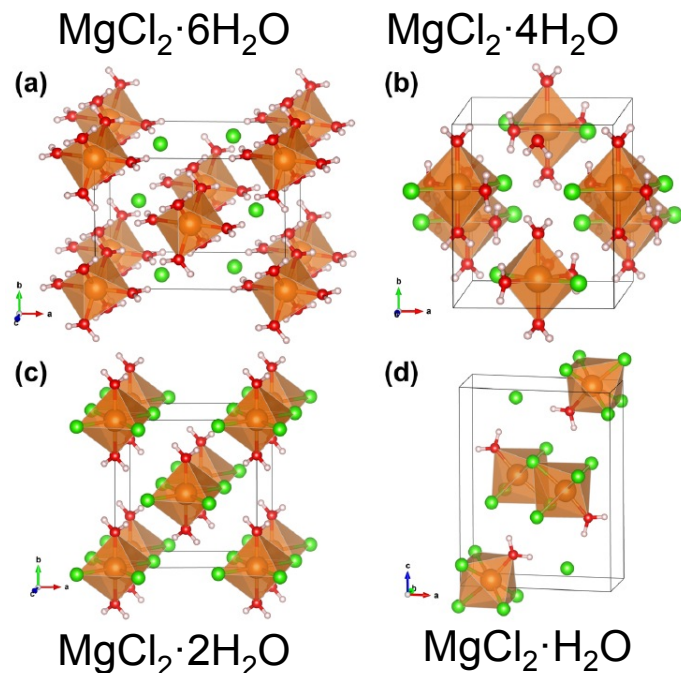
Weck P.F., Kim E., Jove-Colon C.F., Sassani D.C., "First-Principles Study of Anhydrite, Polyhalite and Carnallite", *Chemical Physics Letters* **594**, 1-5 (2014)



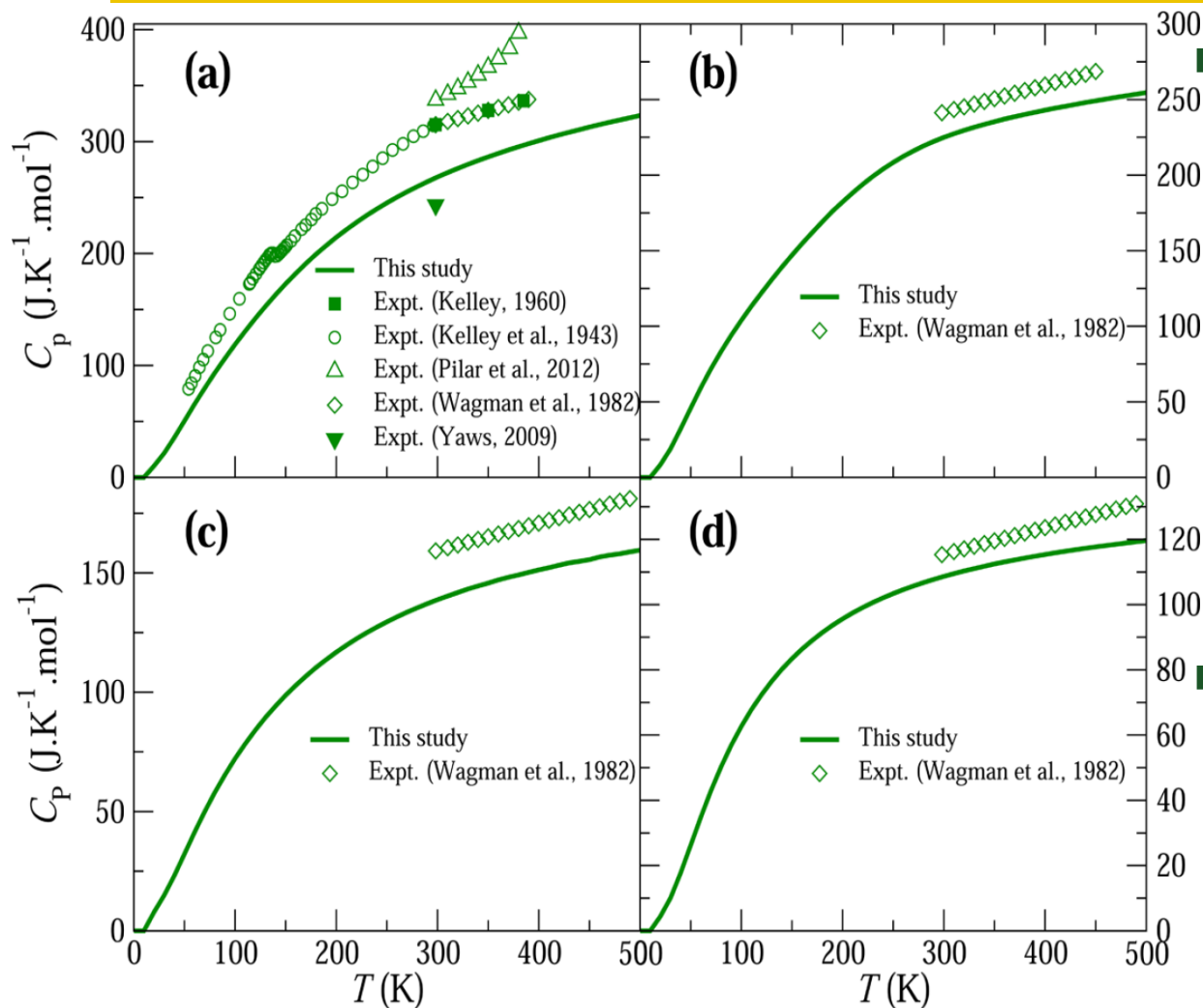
# R&D Highlight on Crystalline Salts: Bischofite and its Dehydrated Phases



(Experimental data for the entropy: FactSage  
FACTPS database.)



# R&D Highlight on Crystalline Salts: Bischofite and its Dehydrated Phases



## Thermal properties:

- Isochoric and isobaric properties were calculated for bischofite and its dehydrated phases.
- Good agreement between computed molar latent heat  $Q_P = 60.8 \text{ kJ mol}^{-1}$  for the  $\text{MgCl}_2\cdot 6\text{H}_2\text{O}$  to  $\text{MgCl}_2\cdot 4\text{H}_2\text{O}$  transformation and expt. ( $Q_P = 68.2 \text{ kJ mol}^{-1}$ ).

## FY14/15 outlook:

Extend structural and thermal-mechanical calculations within the framework of DFT/DFPT to other NS minerals (clays, salts, granite...) and EBS materials.