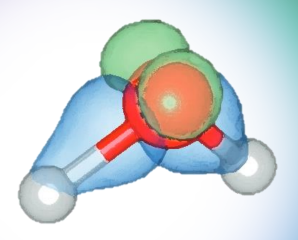
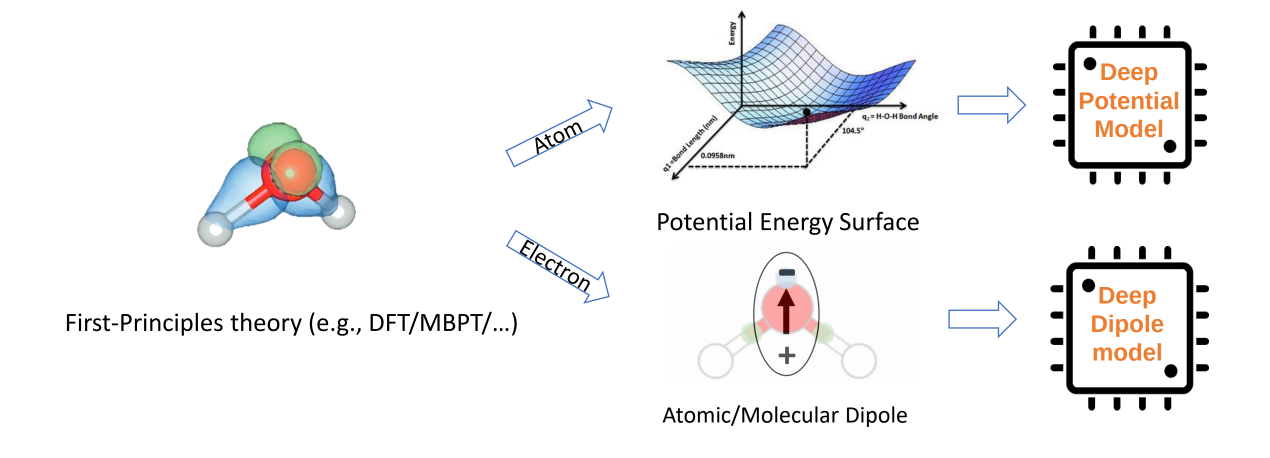
2022 Chemistry in Solution and at Interfaces (CSI) Workshop: Modeling for Molecular Simulation



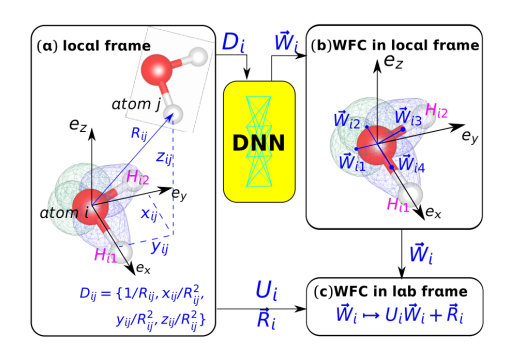
# Tutorial Session: Deep Dipole

SPEAKER: PINCHEN XIE
PACM, PRINCETON
JUNE 8, 2022

# Beyond potential energy surface



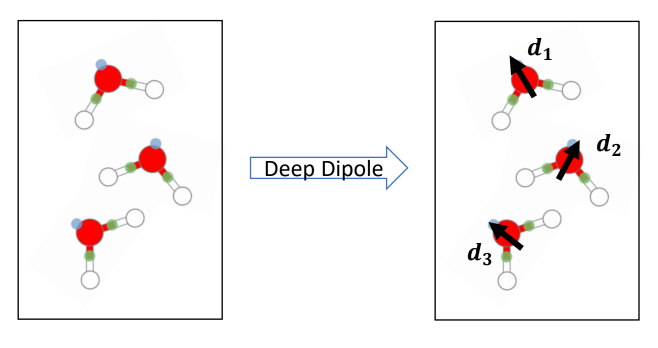
### The open box



Zhang, Linfeng, et al. "Deep neural network for Wannier function centers." *arXiv preprint arXiv:1906.11434* (2019).

Zhang, Linfeng, et al. "Deep neural network for the dielectric response of insulators." Physical Review B 102.4 (2020): 041121.

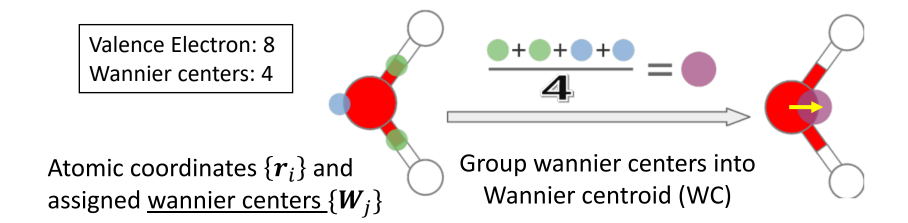
#### The black box



Atomic coordinates  $\{r_i\}$ 

The dipole moment  $\{d_j\}$ 

Example: the water molecule

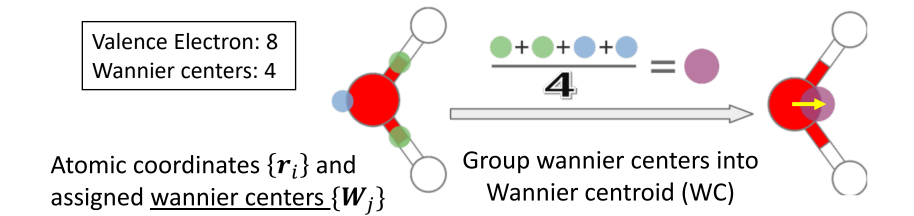


The training label

Position vector of WC w.r.t. central atom (e.g. Oxygen)

Example: the water molecule

**Issue: wannier centers not unique** (U(1) Gauge)



The training label

Position vector of WC w.r.t. central atom (e.g. Oxygen)

### Maximally localized Wannier functions (MLWF)

The General wannier function: 
$$|\mathbf{R}n\rangle = \frac{V}{(2\pi)^3} \int_{\mathrm{BZ}} d\mathbf{k} \, e^{-i\mathbf{k}\cdot\mathbf{R}} \sum_{m=1}^J U_{mn}^{(\mathbf{k})} |\psi_{m\mathbf{k}}\rangle$$
 Bloch orbitals

The MLWF: 
$$|\mathbf{R}n\rangle = \frac{V}{(2\pi)^3} \int_{\mathrm{BZ}} d\mathbf{k} \, e^{-i\mathbf{k}\cdot\mathbf{R}} \sum_{m=1}^{J} U_{mn}^{(\mathbf{k})} |\psi_{m\mathbf{k}}\rangle$$

The optimal rotation minimizing the total spanning of wannier functions

$$\Omega = \sum_{n} \left[ \langle \mathbf{0}n | r^{2} | \mathbf{0}n \rangle - \langle \mathbf{0}n | \mathbf{r} | \mathbf{0}n \rangle^{2} \right]$$
$$= \sum_{n} \left[ \langle r^{2} \rangle_{n} - \bar{\mathbf{r}}_{n}^{2} \right]$$

**Arbitrary Unitary Rotation** 

## Hands-on Exercise 1 -- 20 minutes

#### Goal:

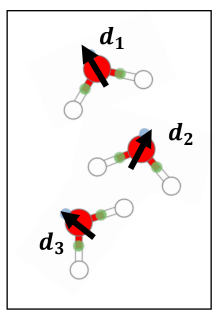
- (1) Compute maximally localized wannier functions of given water configuration with Quantum-Espresso and Wannier90
- (2) Optional: Visualize the wannier centers.

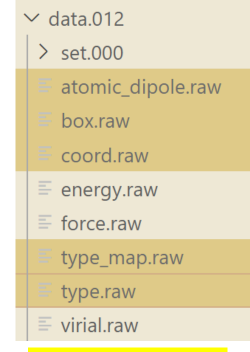
#### How:

- (1) Read and execute the shell script "submit.slurm" in workshop-july-2022/hands-on-sessions/day-2/DeepWannier/wannier90\_example/ice\_mlwf/
- (2) Get familiar with the outputs
- (3) Visualize "water\_centres.xyz" with OVITO

## Training Deep Dipole: Dataset

Required data





Optional: dipole.raw

Reference atoms for dipole moments, e.g. oxygen for H2O

Format

atomic\_dipole.raw:(#frames, #sel\_atom× 3)

Vector concatenated in the same order as its reference atom appearing in coord.raw

dipole.raw: (#frames, 3) <- the sum of atomic dipole

atomic\_dipole.npy&dipole.npy:
Same thing but in numpy format.

# Training Deep Dipole: parameters

#### Difference between training Deep Potential & Deep Dipole

	fitting_net	loss
Energy model	"fitting_net": {	"loss": {     "start_pref_e": 0.01,     "limit_pref_e": 1,     "start_pref_f": 100,     "limit_pref_f": 1,     "start_pref_v": 0,     "limit_pref_v": 0, },
Dipole model	"fitting_net": {	"loss": {  "type": "tensor",  "pref": 1.0, ← O.0 if there is no did on the control of the contr

Types of atom that carries a dipole.

## Hands-on Exercise 2 -- 10 minutes

#### Goal:

(1) Train a deep dipole with provided dataset: workshop-july-2022/hands-on-sessions/day-2/DeepWannier/data

#### How:

- (1) Example can be found in workshop-july-2022/hands-on-sessions/day-2/DeepWannier/train\_dipole\_model/
- (2) Before training, examine dataset and verify that the number of columns in "atomic\_dipole.raw" is compatible with "sel\_type" and the number of water molecules.
- (3) Try submit a training task. Check logs.

## Application of Deep Dipole

Postprocessing (Today)

After a regular DPMD simulation. Process the dumped trajectory with deep dipole model. You can compute dipole distribution / polarization/ dielectric response .....

Enhanced sampling / Electric field-driven MD

Pair Lammps up with PLUMED and use deep dipole model as a collective variable.

## Hands-on Exercise 3 -- 20 minutes

#### Goal:

(1) Run a DPMD for liquid water with provided Deep Potential model. Then analyzed the dumped trajectory with Deep Dipole model.

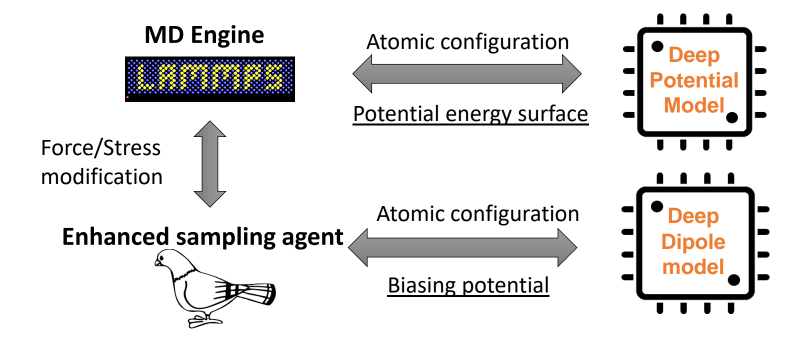
#### How:

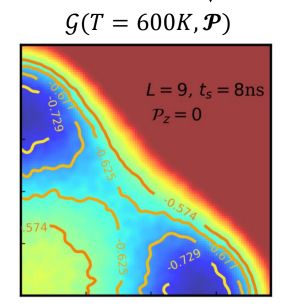
- (1) Example for running DPMP can be found in workshop-july-2022/hands-on-sessions/day-2/DeepWannier/liquid\_dipole/
- (2) Compute and plot the molecule dipole moments distribution with Python. Example script:

workshop-july-2022/hands-on-sessions/day-2/DeepWannier/dipole\_distribution.py

# Enhanced sampling

**PLUMED** 





A cross section of the 3D free energy surface for the polarization of PbTiO3.
Computed by metadynamics.

Reference: Xie, P., Chen, Y. and Car, R., 2022. Ab initio multi-scale modeling of ferroelectrics:

The case of PbTiO3. arXiv preprint arXiv:2205.11839.

PLUMED plugin: <a href="https://github.com/y1xiaoc/deepmd-plumed">https://github.com/y1xiaoc/deepmd-plumed</a> (not MPI-ready yet)