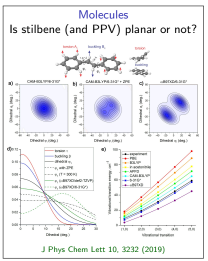
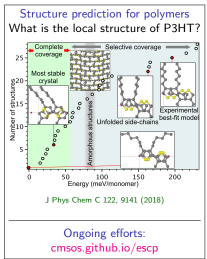
### Structural studies: slide 1 of 2

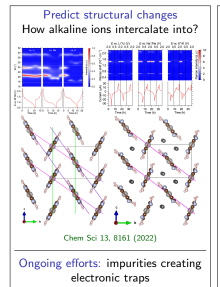
(first-principles at DFT-D accuracy, up to 1000 atoms in simulation cell)





## Structural studies: slide 2 of 2

(first-principles at DFT-D accuracy, up to 1000 atoms in simulation cell)



#### Computer-aided design of materials

- Structure refinement
  - from experiment
  - from structural models
  - from force field models
- Benchmarking methods cmsos.github.io/bmcos



JCTC 19, 8481 (2023) - 67 crystals

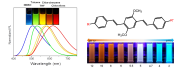
- Exploring novel architectures
  - frameworks
  - ► 2D/3D polymers
  - wiremesh Chem Mater 33, 966 (2021)
  - **nonplanar**  $\pi$ -systems
    - interdigitated herringbone

# Electronic properties of materials: slide 1 of 2

(scalable approaches  $\implies$  multiscale modeling, DFT-parametrized effective Hamiltonian)

#### Electronic/vibrational spectroscopy Commonly used to probe local structural and electronic properties

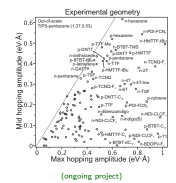
- Derive structural information
  J Phys Chem Lett 10, 3232 (2019) stilbene
  Chem Sci 13, 8161 (2022) charging
- Screen out inaccurate methods
  Chem Sci 13, 8161 (2022) by bandgap
- Understand spectral changes



ACS Appl Mater Interfaces 5, 4685 (2013) Chem Sci 6, 789 (2015)

Chem Phys 481, 133 (2016)

# Charge carrier transport How to quickly estimate mobility?

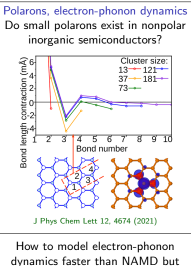


What is the origin of nonmonotonic temperature dependence of luminescence kinetics?

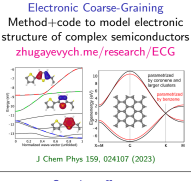
Low Temperature Physics 28, 706 (2002)

## Electronic properties of materials: slide 2 of 2

(scalable approaches  $\implies$  multiscale modeling, DFT-parametrized effective Hamiltonian)



more accurately than KMC?



#### Ongoing efforts:

- cmsos.github.io/escp
- cmsos.github.io/tbm (library of electronic prototypes)
- excited states
- add vibronic couplings