

Tribochemistry of Molybdenum Disulfide



PRESENTED BY

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U.S. DEPARTMENT OF
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SURFACE INTERFACES AND MATERIALS
TRIBOLOGY LABORATORY

3 MoS₂ is a Versatile Lubricant

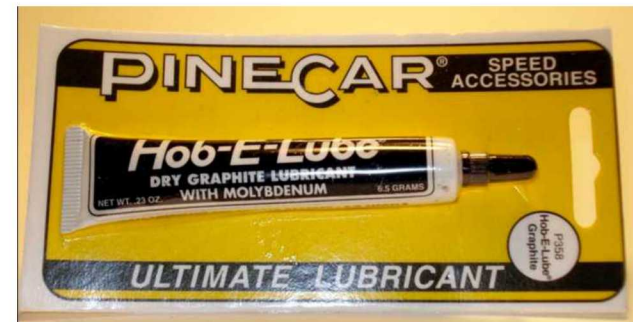


• Dry lubricant

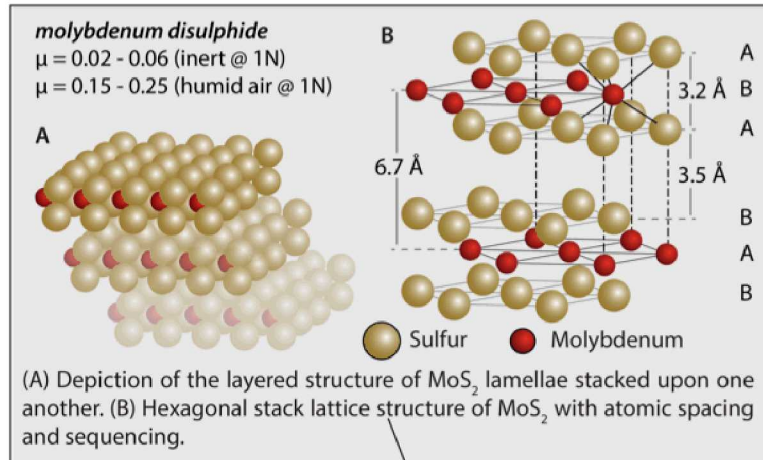
- Superlubric ($\mu < 0.01$) in dry environments
- Coaster brakes, CV joints, ski wax, bullets...
- Satellites, aircraft engines
- Self-lubricating composites with polymers

• Other uses:

- Catalysis (desulfurization, electrolysis of water)
- Memristor/memcapacitors
- Flexible circuits

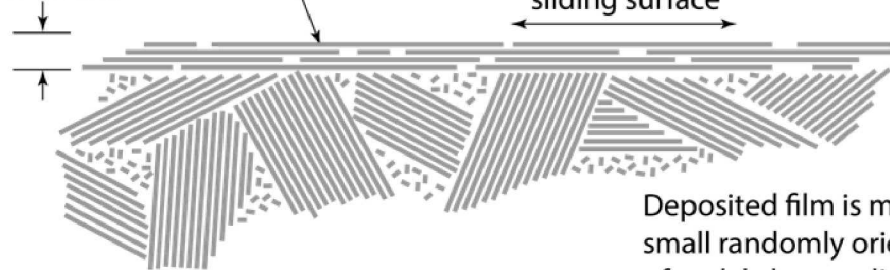


How Does it Work?

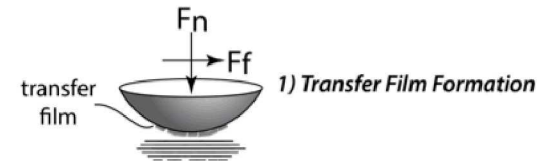


oriented surface layer
of 002 basal planes of MoS₂

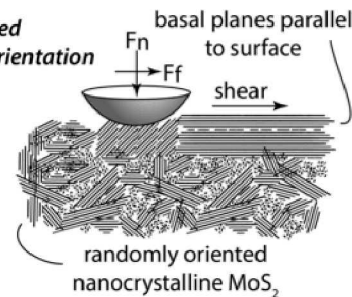
3-10 nm



Run-In Processes



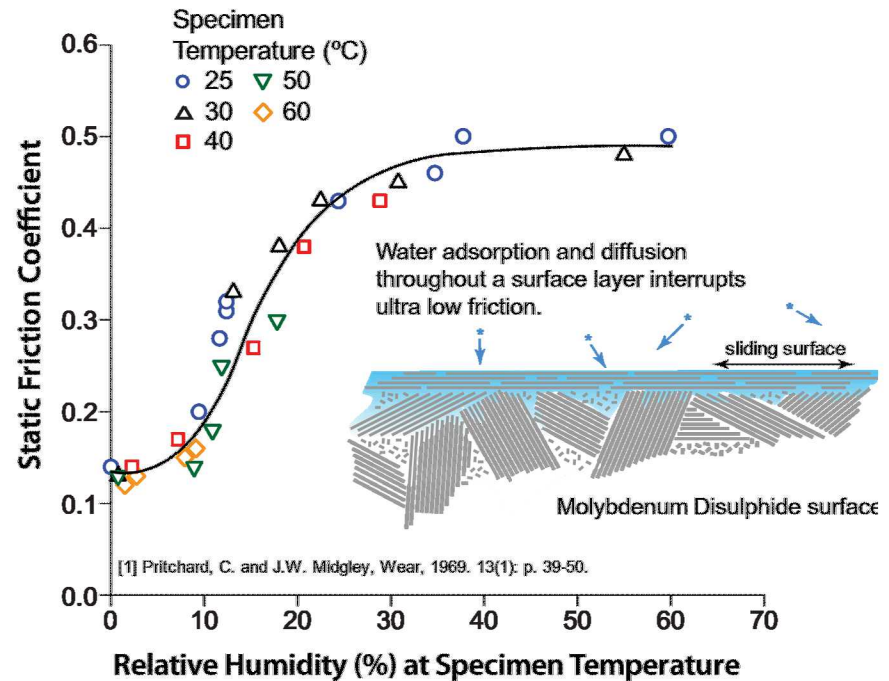
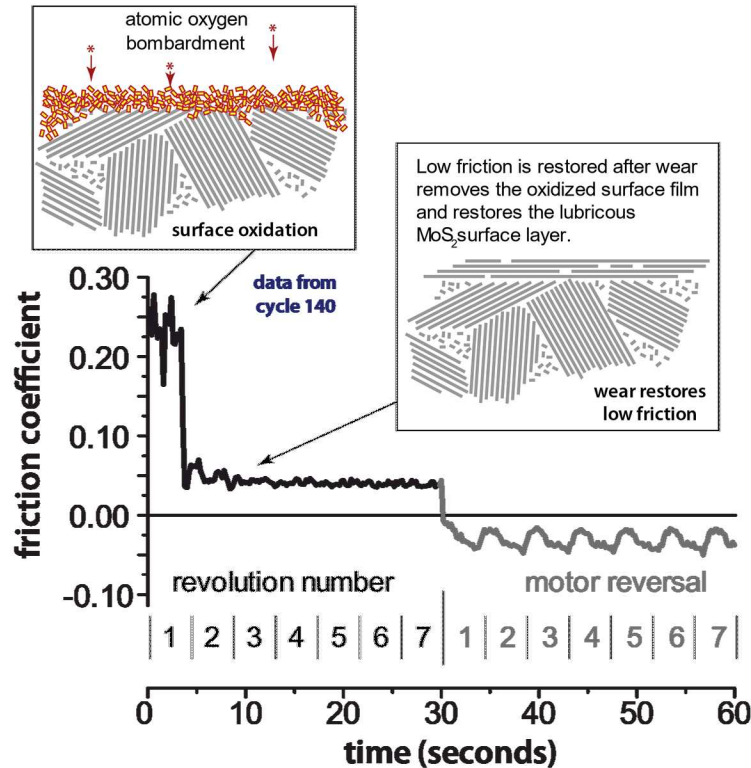
2) *Shear-induced crystallite re-orientation*



Deposited film is made of many small randomly oriented crystallites of molybdenum disulphide.

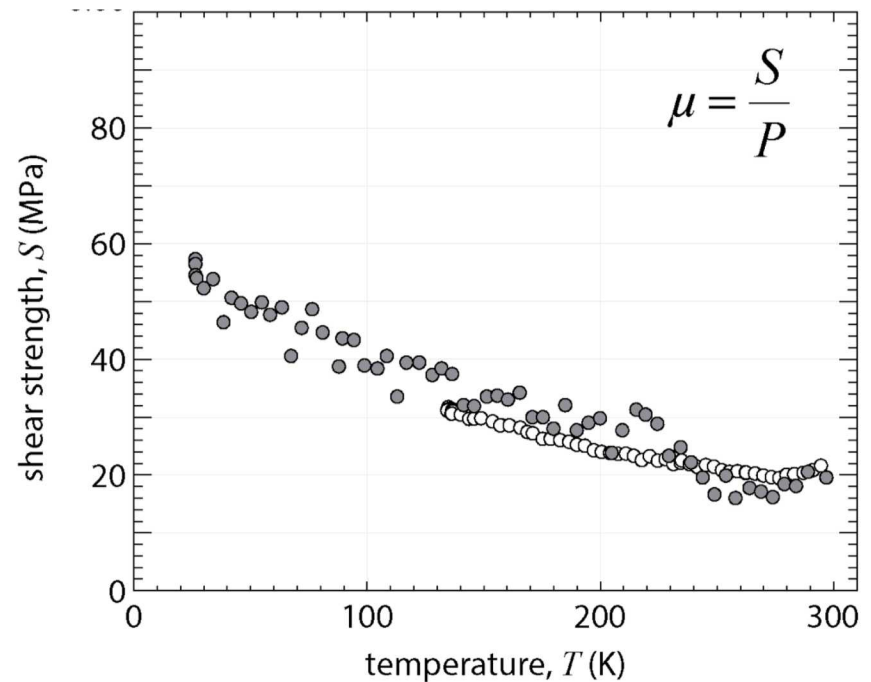
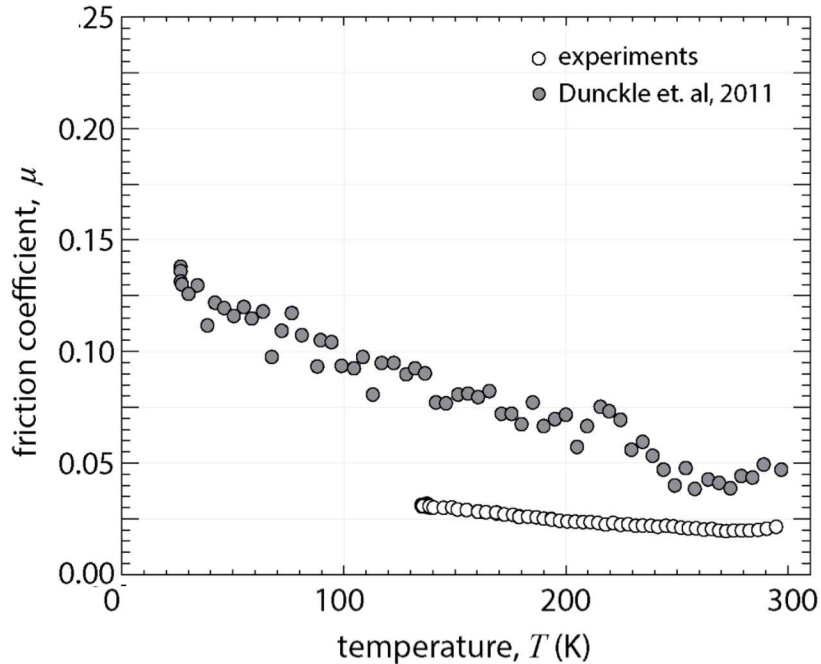
- Hexagonal structure, form thin, weakly bound lamella
- Issues: run-in and oxidation

Environment (oxygen and water) affect friction

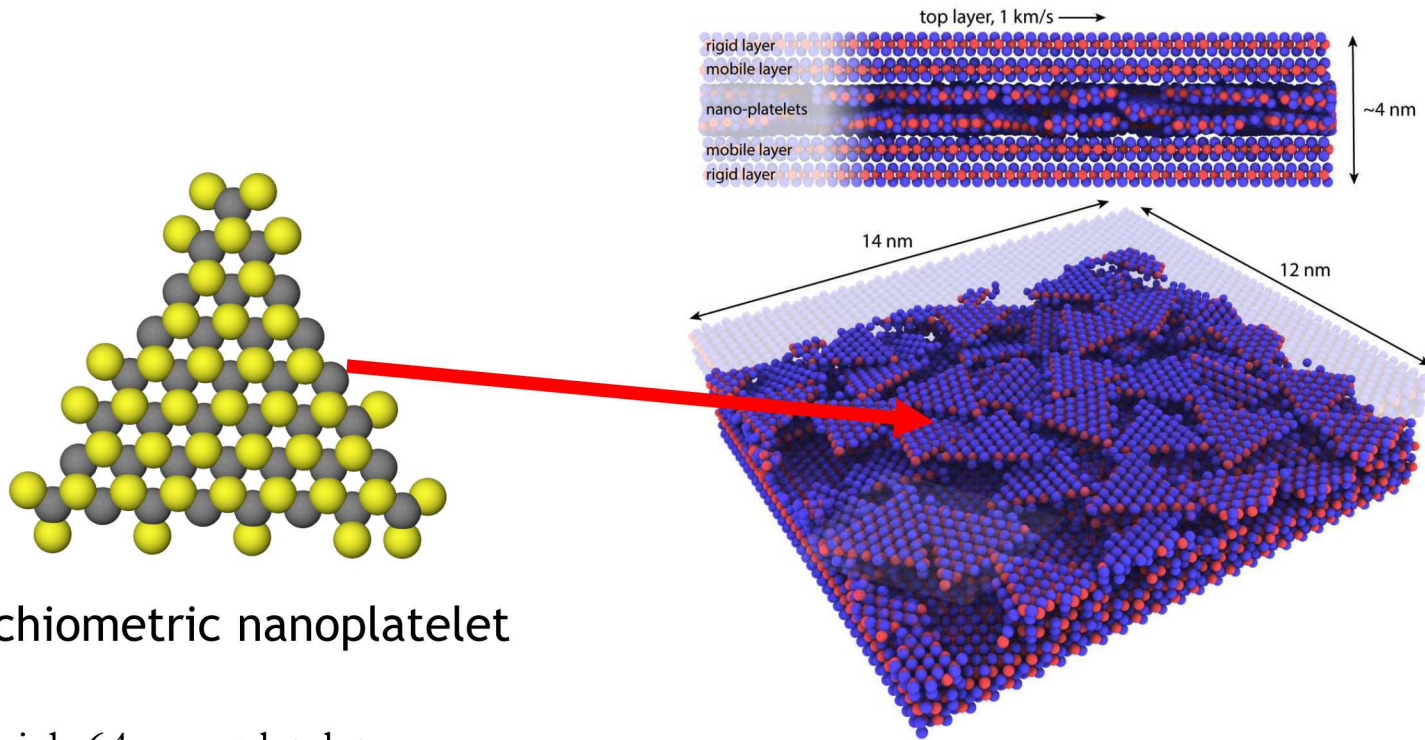


- Oxidation
 - space (AO - fast)
 - air at high temps (O_2 - fast)
 - Air at room temp (H_2O - slow)
- Water enhances static and kinetic friction
 - From environment
 - From inside the film

Start with pure MoS₂ -- Temperature Dependence

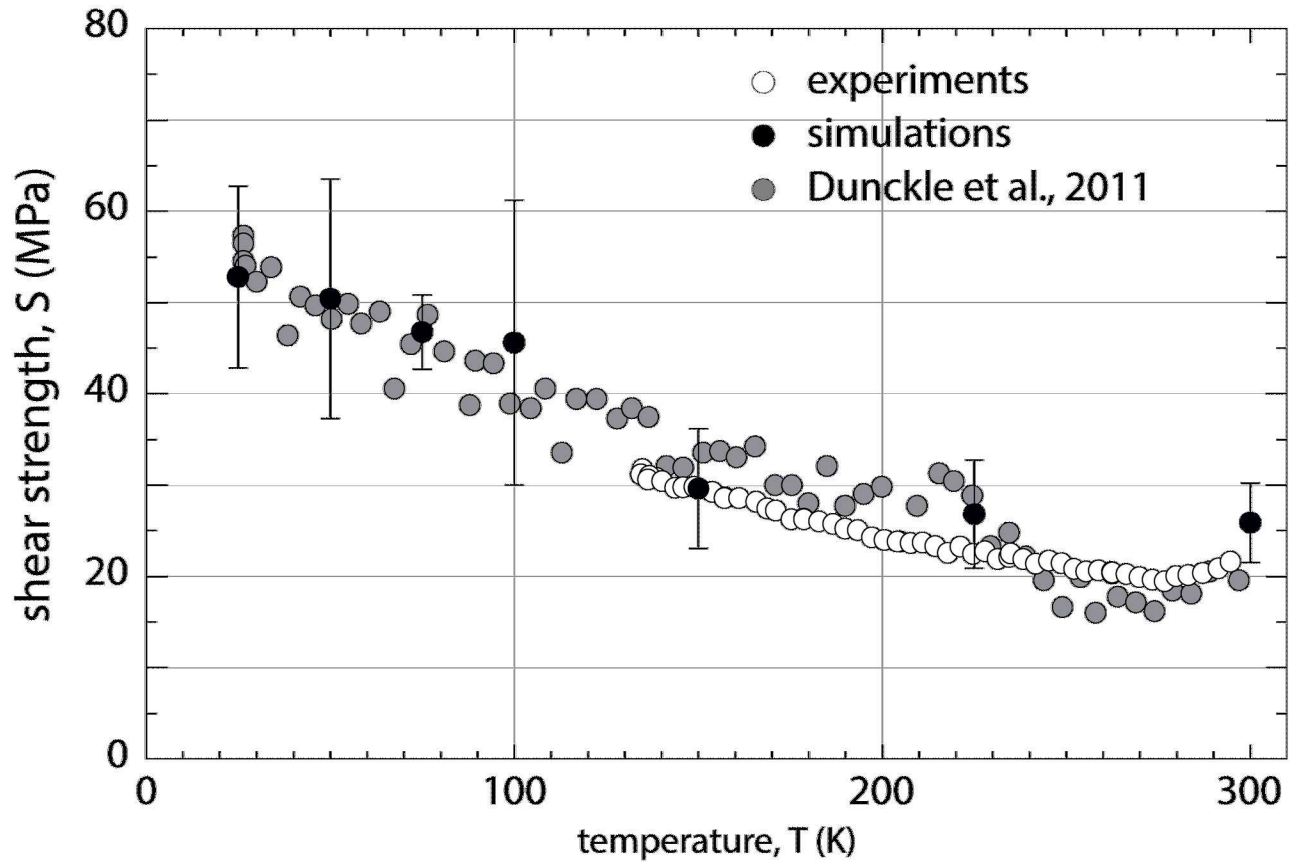


- non-Arrhenius behavior
- Singer (1990) showed contact is purely elastic
 - $\mu = S_0/P + \alpha$
 - $S \sim 25$ MPa at 300K
 - Implies sheets sliding on sheets
- Use simulations to understand the shape



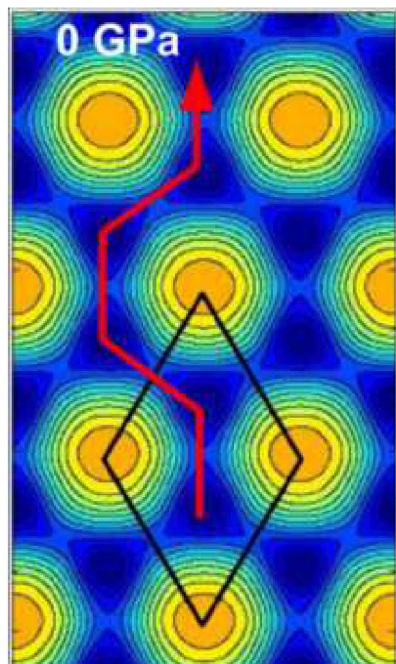
stoichiometric nanoplatelet

- Sandwich 64 nanoplatelets
 - Mobile lamella on top & bottom
 - Fixed lamella (rigid layer) to control load and speed
- ReaxFF: Vasenkov, et al., J. Appl. Phys. 2012
 - Slow technique with (reasonably) accurate chemistry
 - Lots of simulations => small & fast.

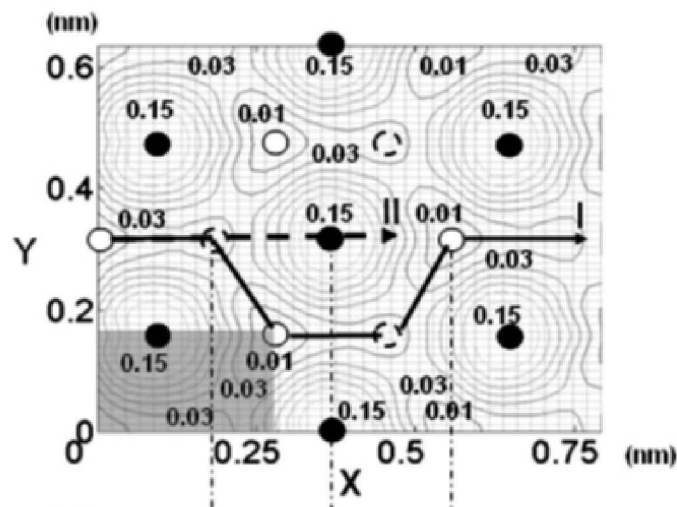


- All shear strengths collapse!
- What causes this shape?

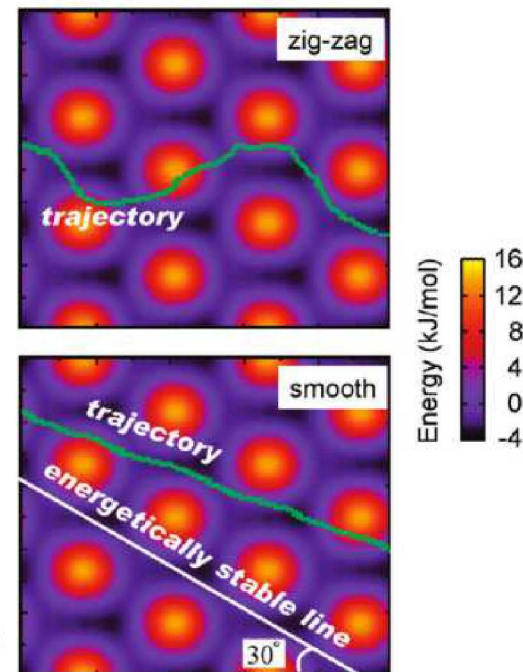
Elastic contact => Energy Barriers: Previous work



Levita et al.,
J. Phys. Chem. C 2014



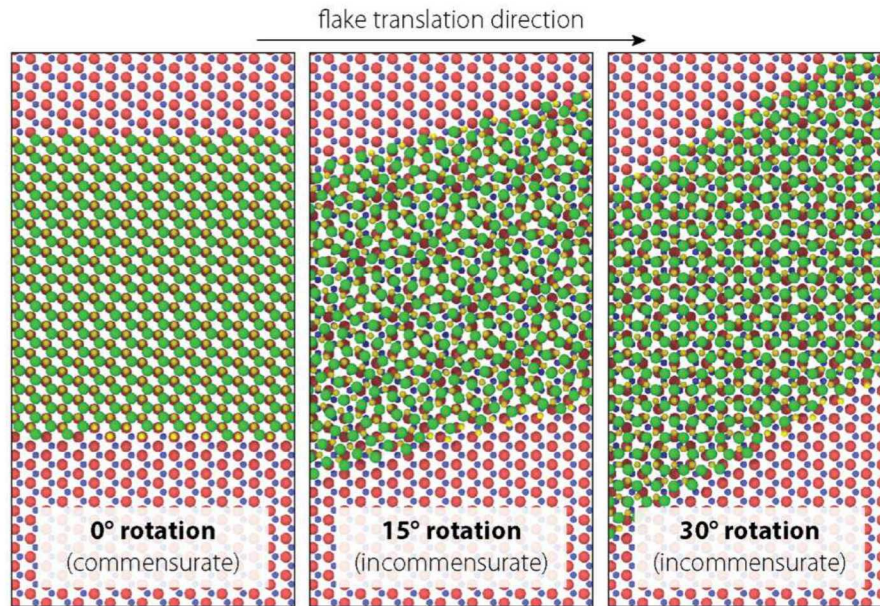
Liang et al.,
Phys. Rev. B 2008



Onodera et al.,
J. Phys. Chem. B 2010

Previous work has calculated energetic barriers to sliding,
but only for commensurate contacts

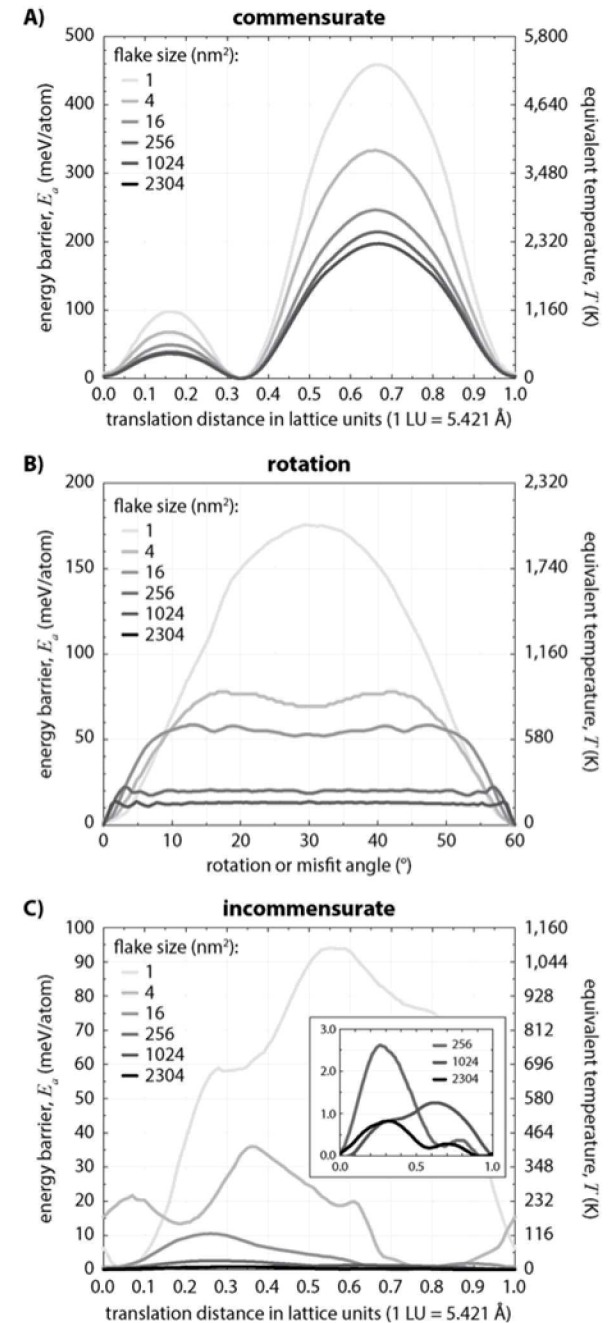
Elastic contact => Energy Barriers: Our work



commensurate
egg shell

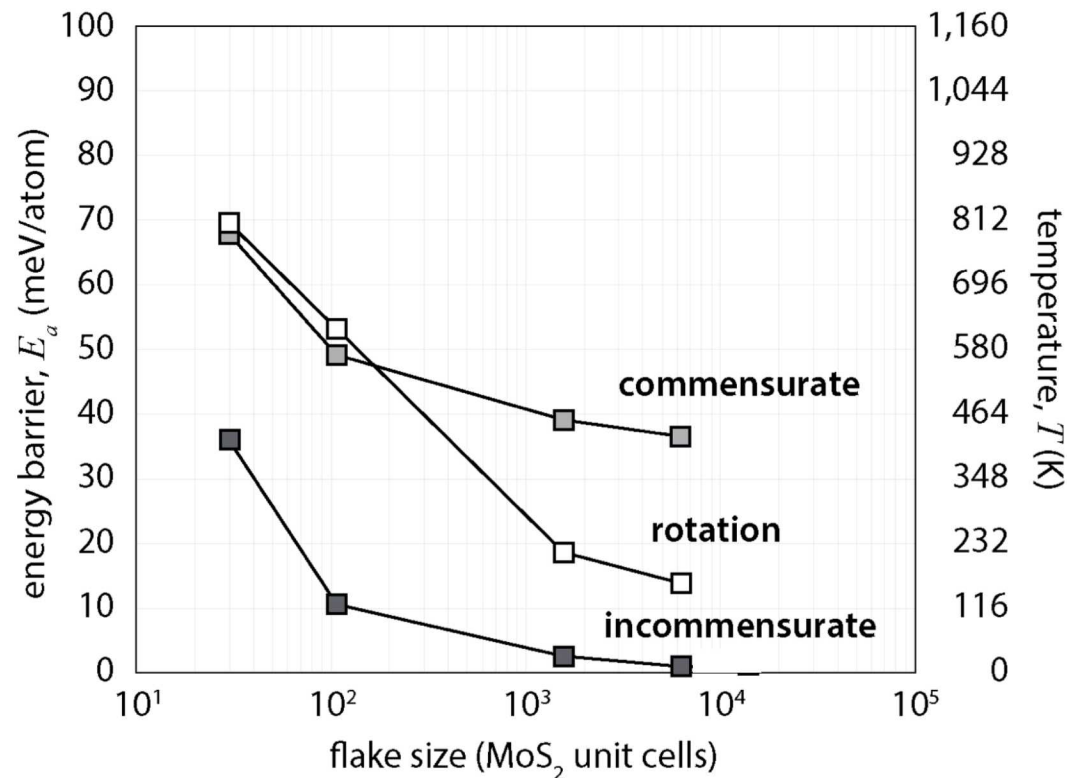


incommensurate
egg shell



Nudged elastic band calculations for barriers

Barriers converge with increasing flake size; make a toy model



Probability & Failure to cross barrier:

$$p_n \propto \exp\left(\frac{-E_n}{kT}\right)$$

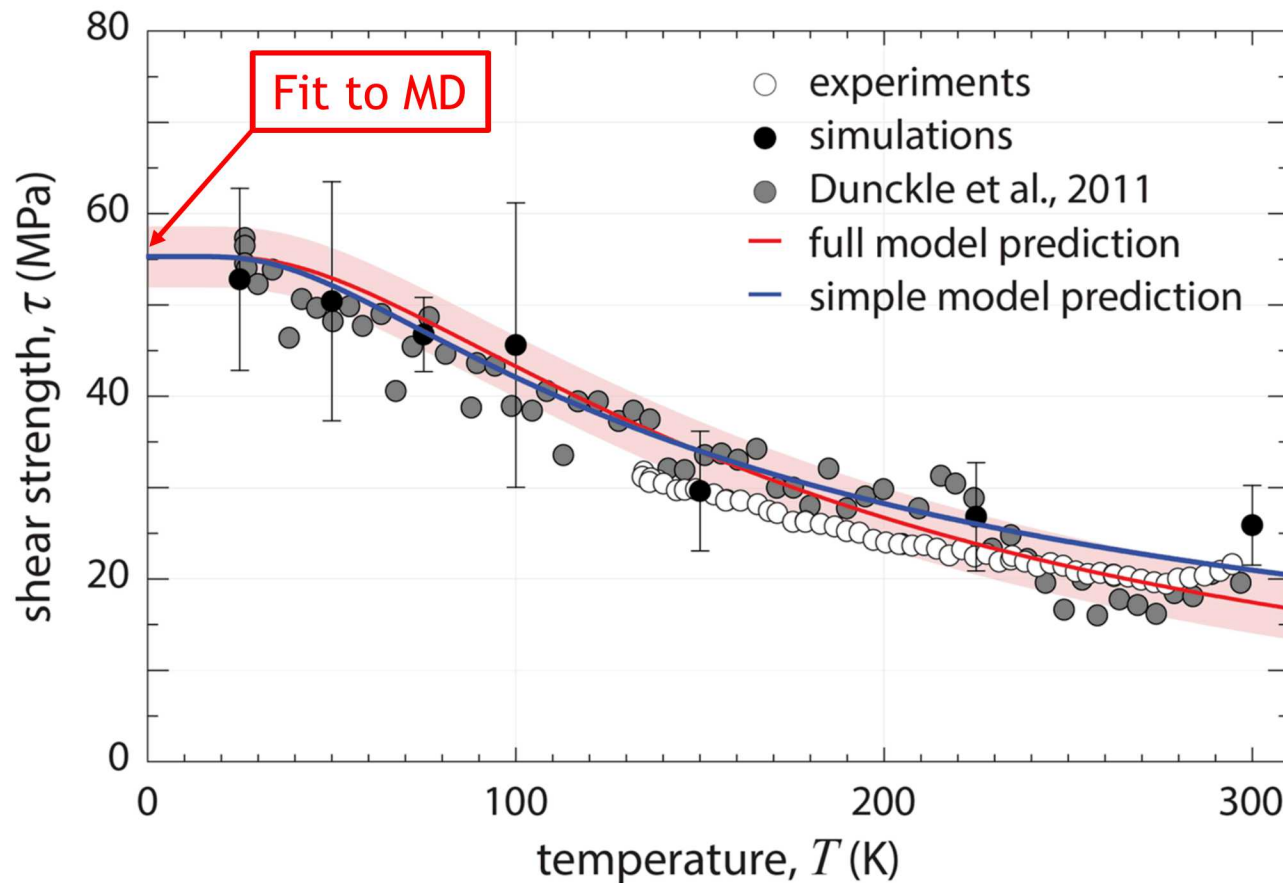
$$f_n = 1 - p_n$$

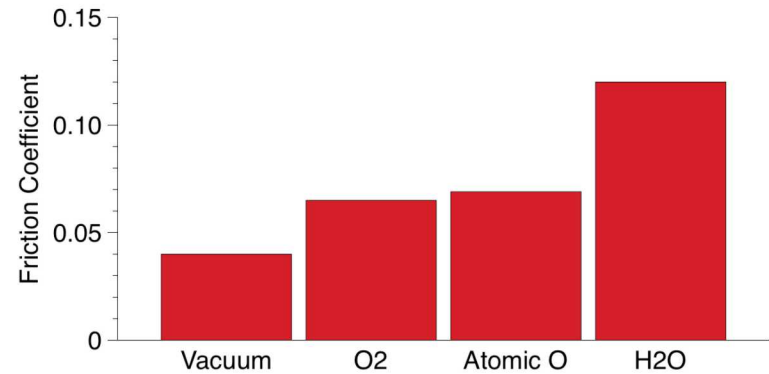
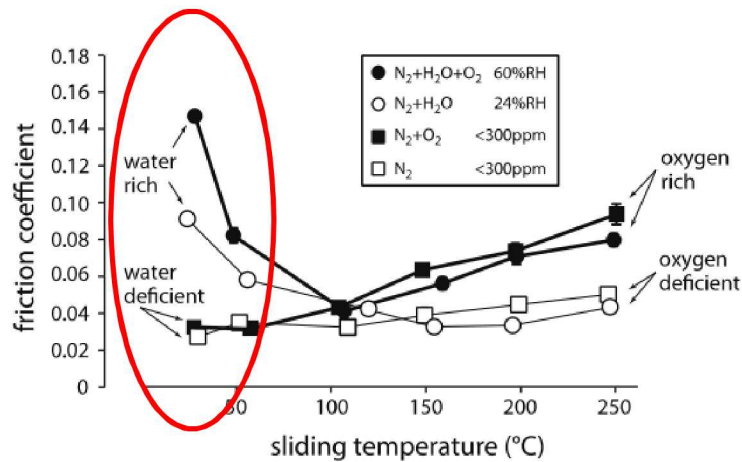
Total sliding probability & friction:

$$p_{slide} = p_r p_i + f_r p_c$$

$$f_{slide} = 1 - p_{slide} = 1 - (p_r p_i + f_r p_c)$$

$$f_{slide} = C_0 \left[1 - \exp \left(- \frac{E_r}{kT} \right) \right]$$

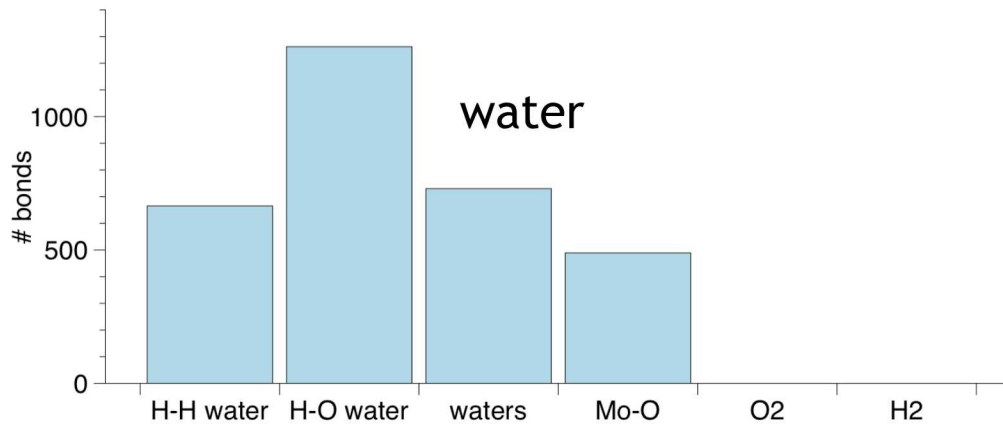
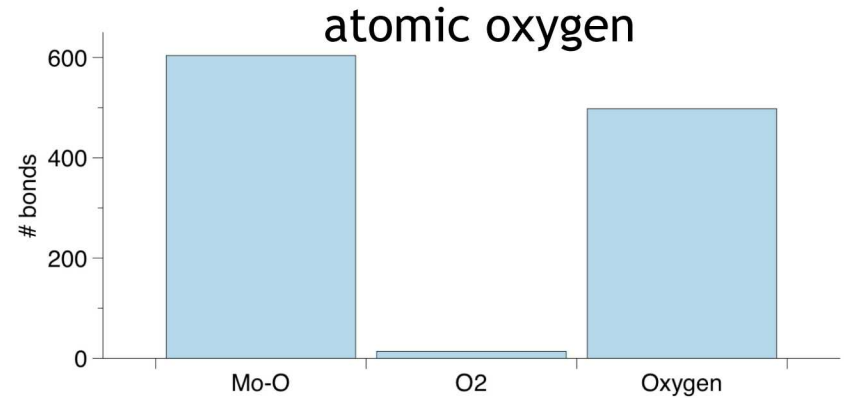
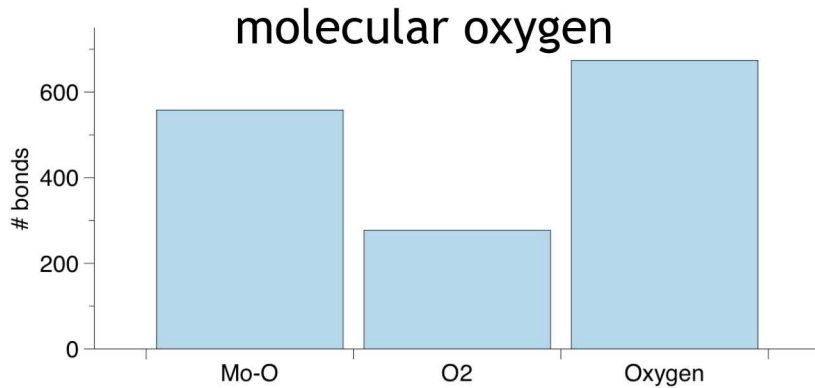




Khare and Burris, Tribol. Lett. 2013

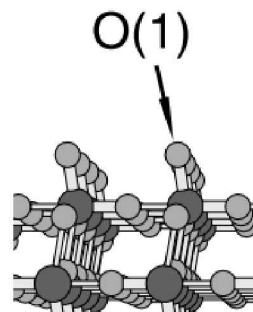
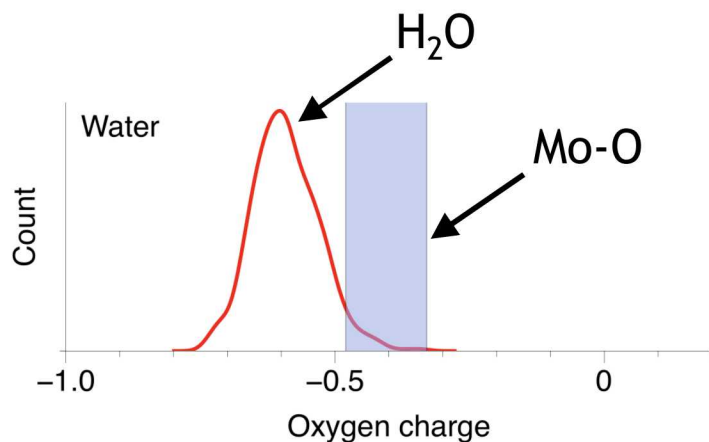
- Changes with added oxygen or water match experimental results quite well (for MD...)

Is there chemistry?

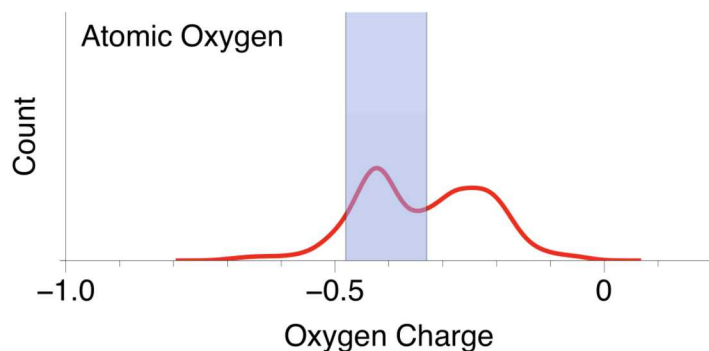
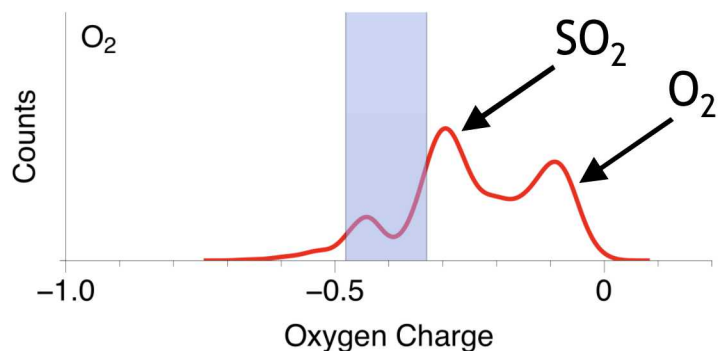


- Water does not dissociate (no O2 or H2 formed)
- Molecular O shows little dissociation (mostly in O2)
- Atomic oxygen forms little O2
- *Not much...*

Charge on Oxygens confirms chemistry

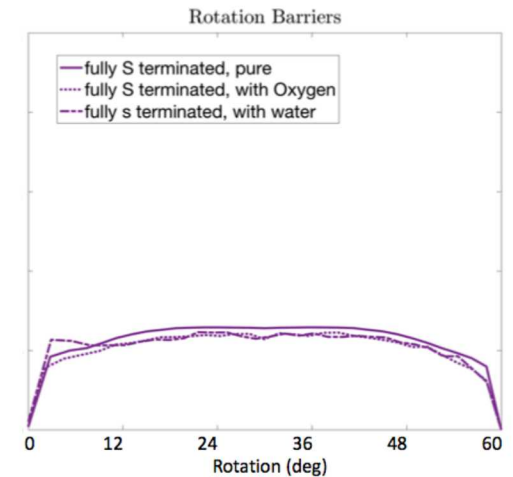
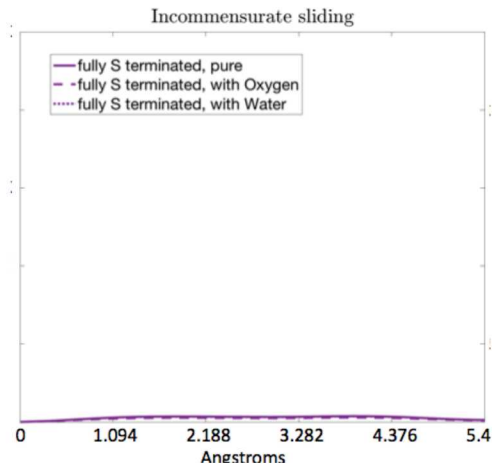
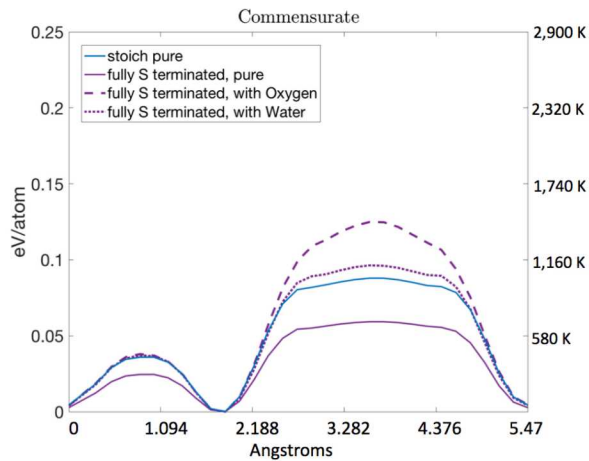


Tokarz-Sobieraj et al.
Surf. Sci. 2001



- Oxygen bonded to Mo has partial charge from -0.48 (Tokarz-Sobieraj et al. Surf. Sci. 2001) to -0.33 (Yin et al., J. Mol. Model 2001).
- Oxygen in water has partial charge from -0.6 to -0.8 (Astrand, et al., J. Phys. Chem. A 1998).
- Water shows only physisorption
- Atomic oxygen shows chemisorption
- Molecular oxygen shows slight amount of chemisorption

What happens to the energy barriers?

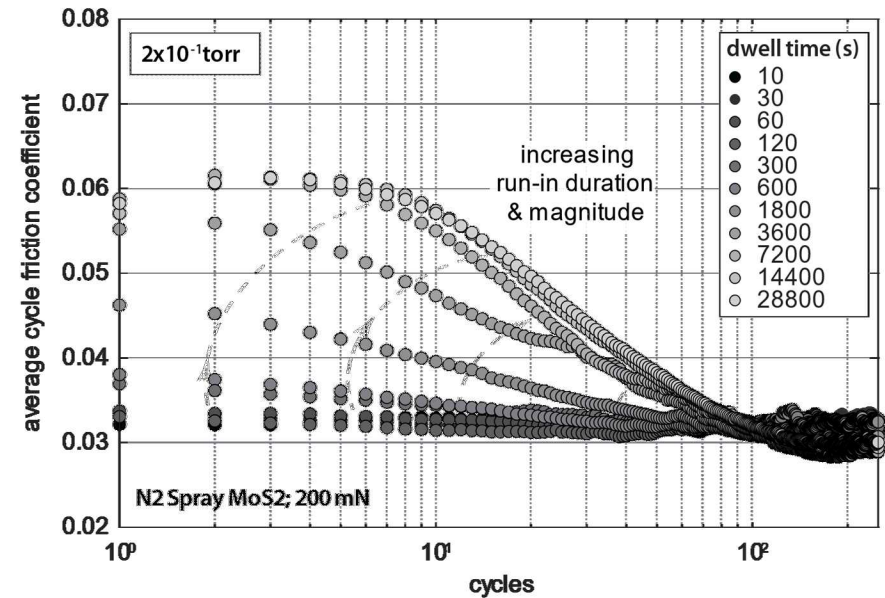
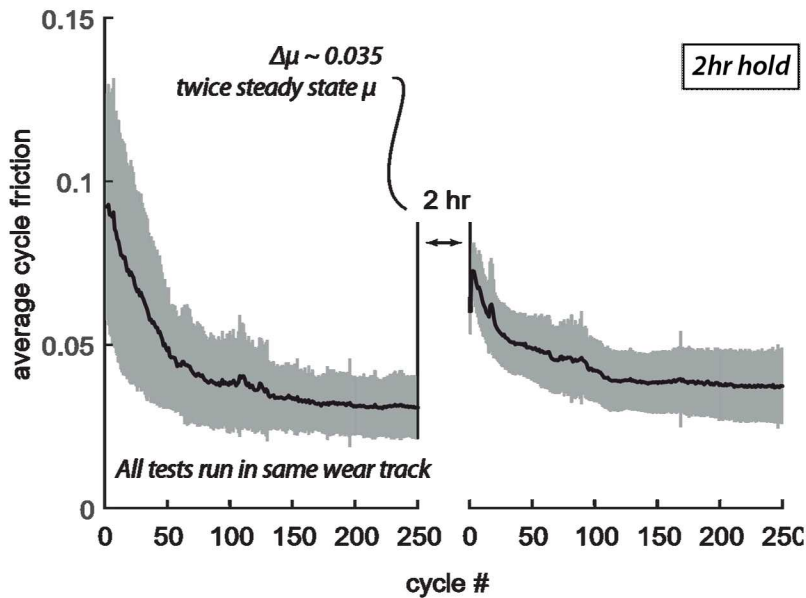


Not much...

So why does the friction go up?

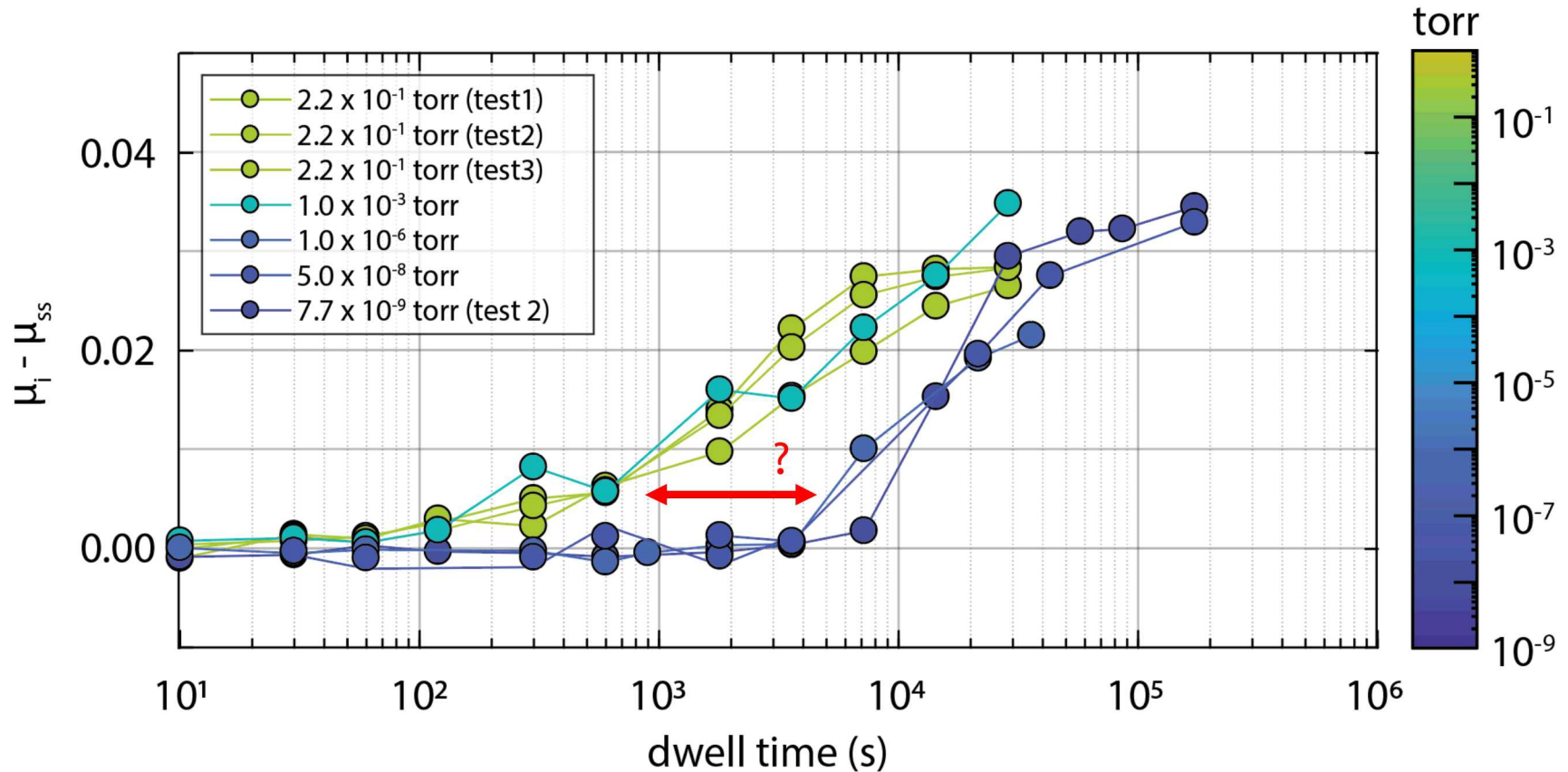
Run-in and re-run-in

- Recipe for success: run film in to steady state... and watch friction increase upon return



- Increase in initial friction increases with time in between; run-in duration also affected
- Also seen in vacuum

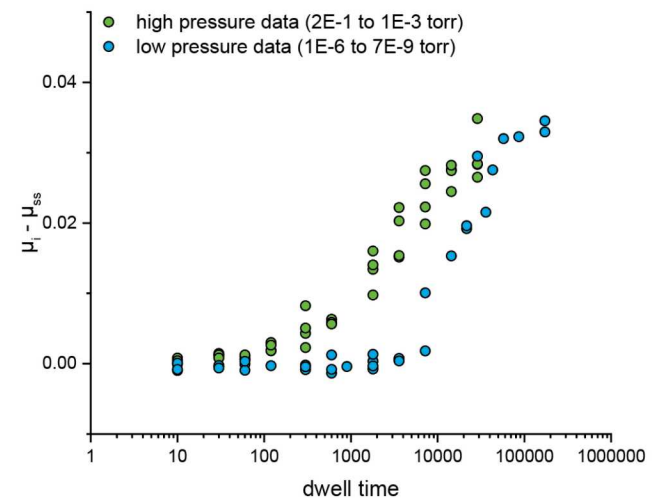
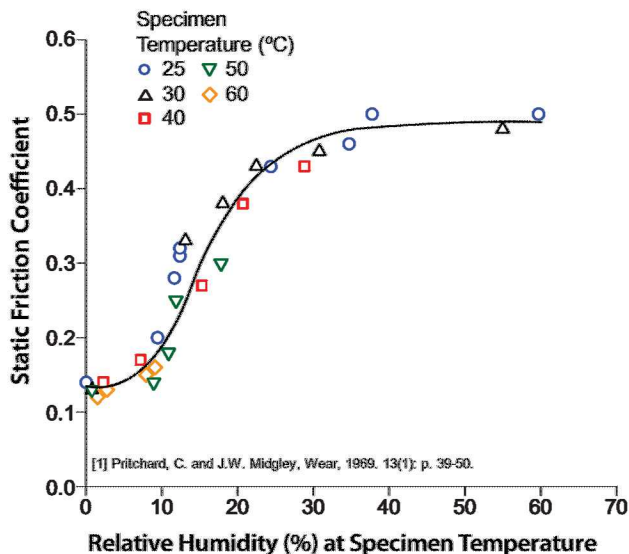
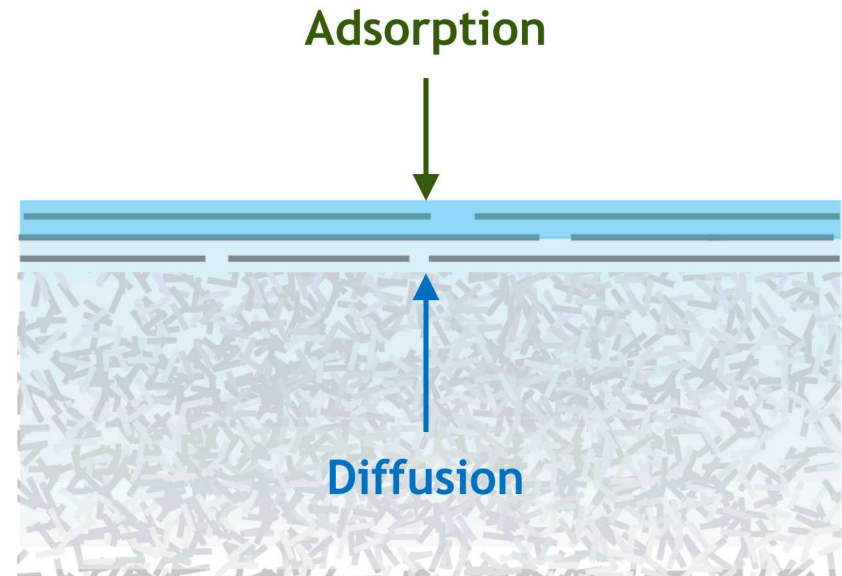
Pressure Matters too!



- difference between previous steady state and returning initial friction
- Stop time from 2×10^{-1} to 7×10^{-9} torr
- Low and high pressures are different

Environmental factors change shear strength

- **Simple theory:** adsorption and diffusion
- Not a new idea:
 - Johnston & Moore 1964
 - Pritchard & Midgeley, 1969
 - Colbert, Ph.D. thesis 2012

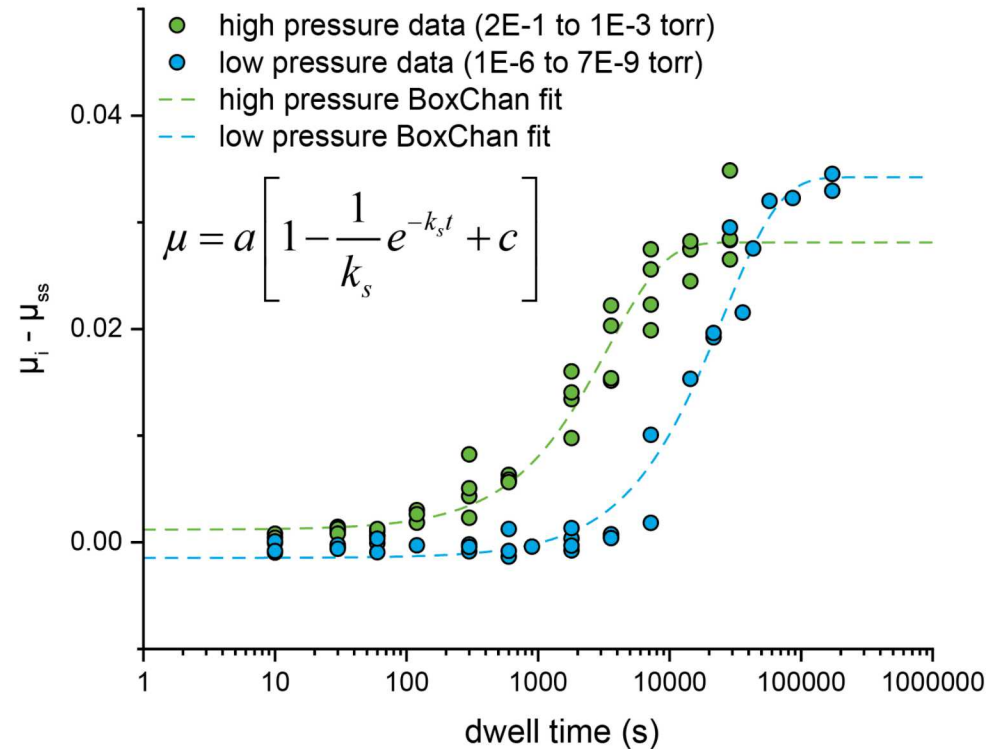


Simple Coverage Model

- simple fractional coverage model
- coverage θ depends on available sites
- two variables:
 - k = rate of arrival
 - s = sticking coefficient
 - $k = k_1 + k_2$, $s = s_1 + s_2$

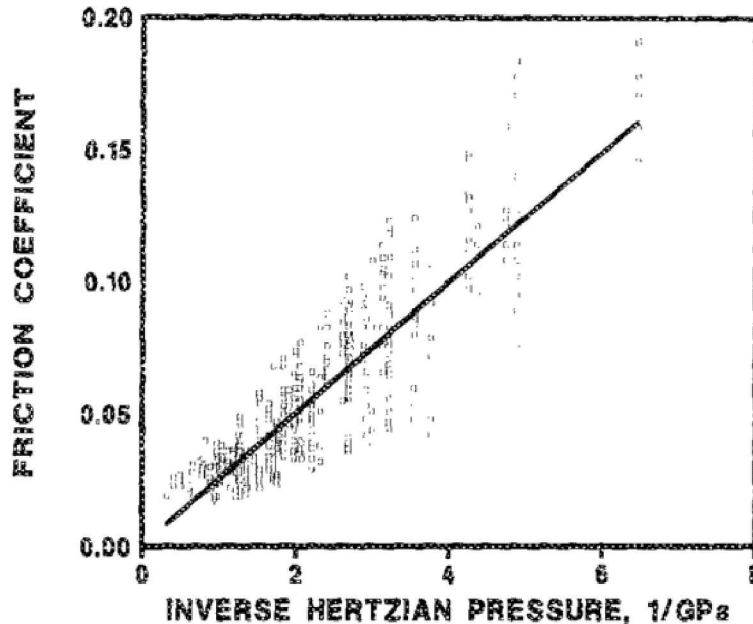
$$\frac{d\theta}{dt} = k \cdot s(1 - \theta)$$

$$\theta = 1 - \frac{1}{k \cdot s} e^{-k \cdot s \cdot t}$$



- MoS₂ shows purely elastic contact
- Shear is predominantly due to inter-lamellar interactions
- Simple model predicts temperature dependence
- No chemistry with water, little with molecular O, lots with atomic O
- Environment hinders formation of large sheets
- Run-in and re-run-in strongly affected by water
 - Adsorption from vapor (at high and low pressures)
 - Diffusion from bulk (low pressure only)
 - Baking out helps!

Singer, et al., Appl. Phys. Lett. 1990



• Singer's explanation:

- $\mu = S/P$

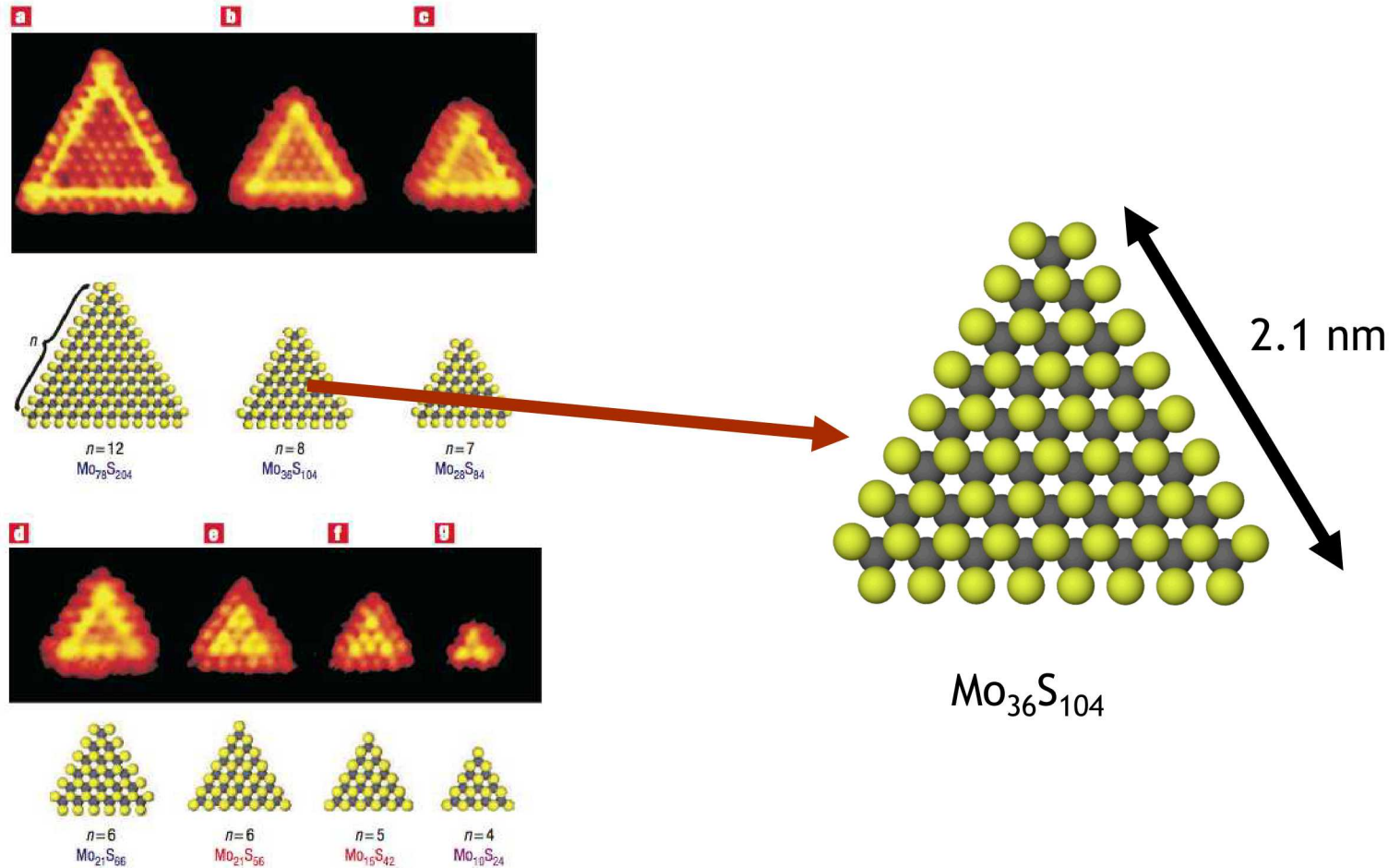
- Expand $S = S_0 + \alpha P$

- $\mu = S_0/P + \alpha$

- $\mu = S_0 \pi (3R/4E)^{2/3} L^{-1/3} + \alpha$

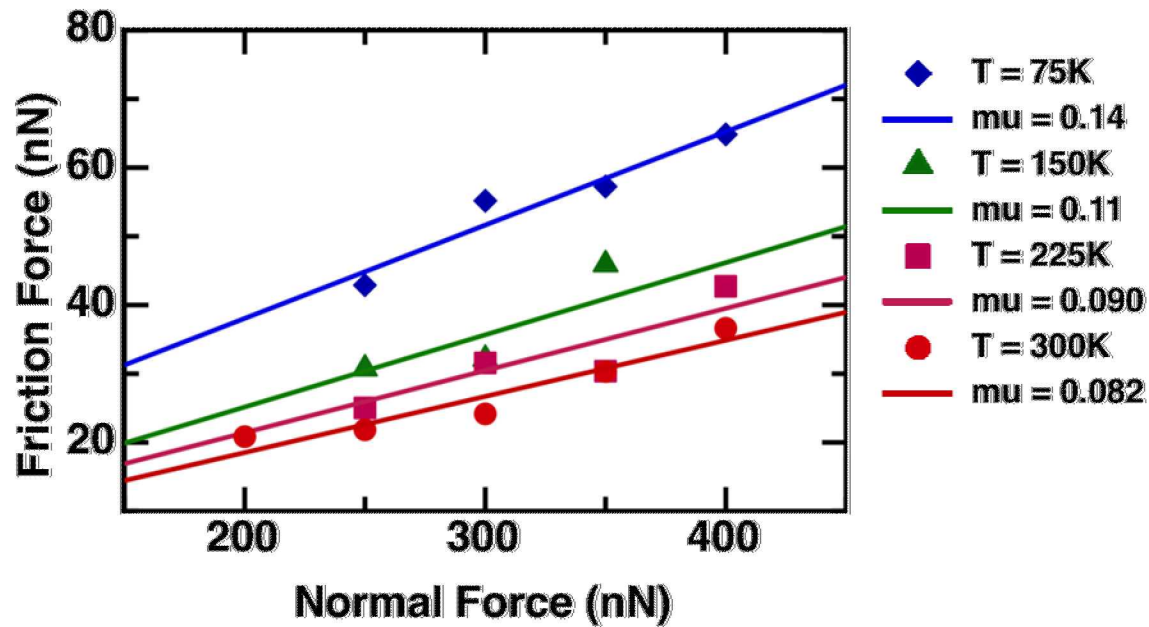
- $S_0 = 25 \text{ MPa}$

• Contact is purely elastic => sheets sliding over sheets

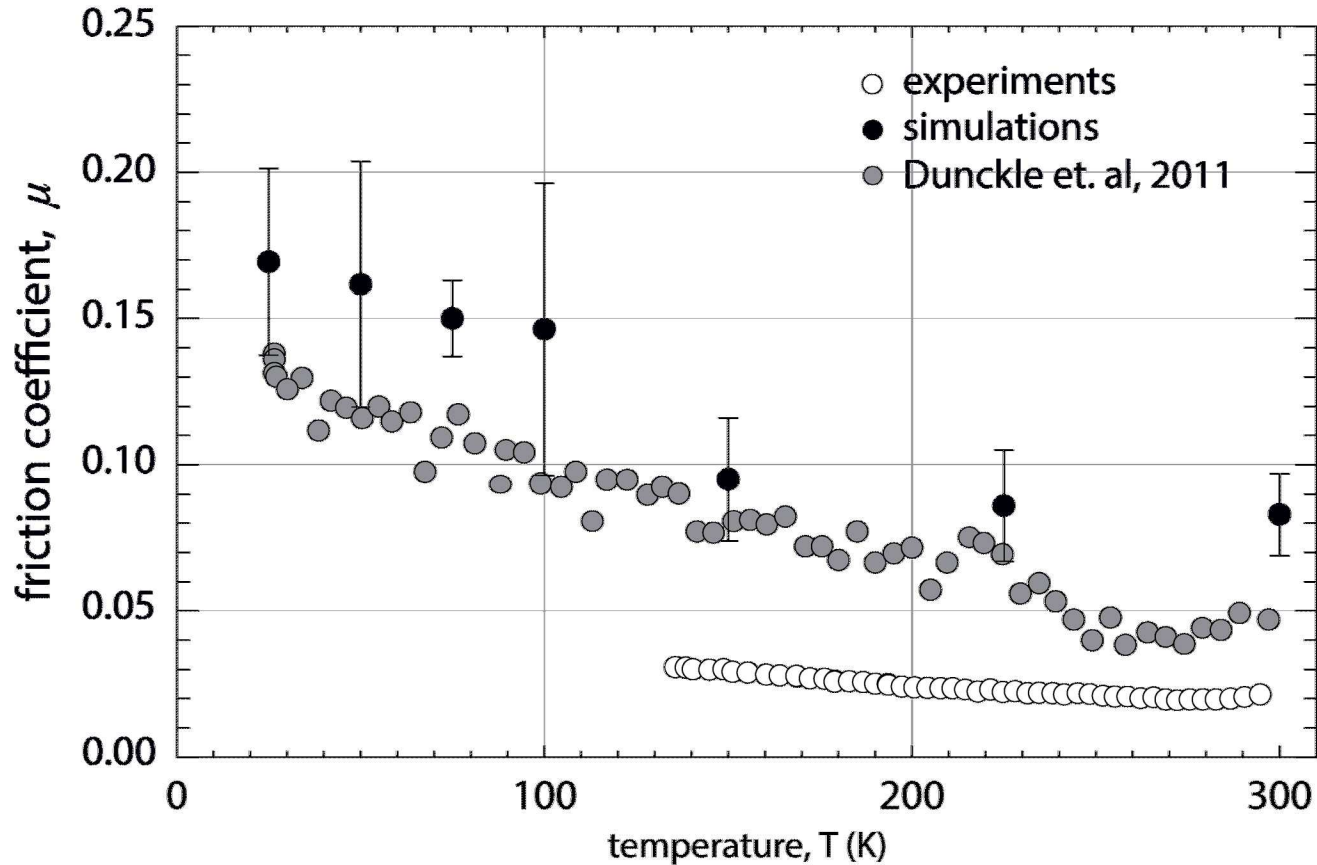


Lauritsen et al., Nature Nanotech. 2007

- Start with nanoplatelets
- Defect free platelets are non-stoichiometric

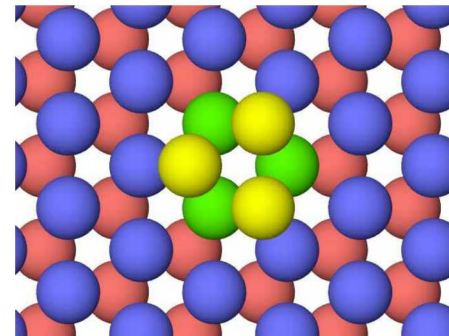
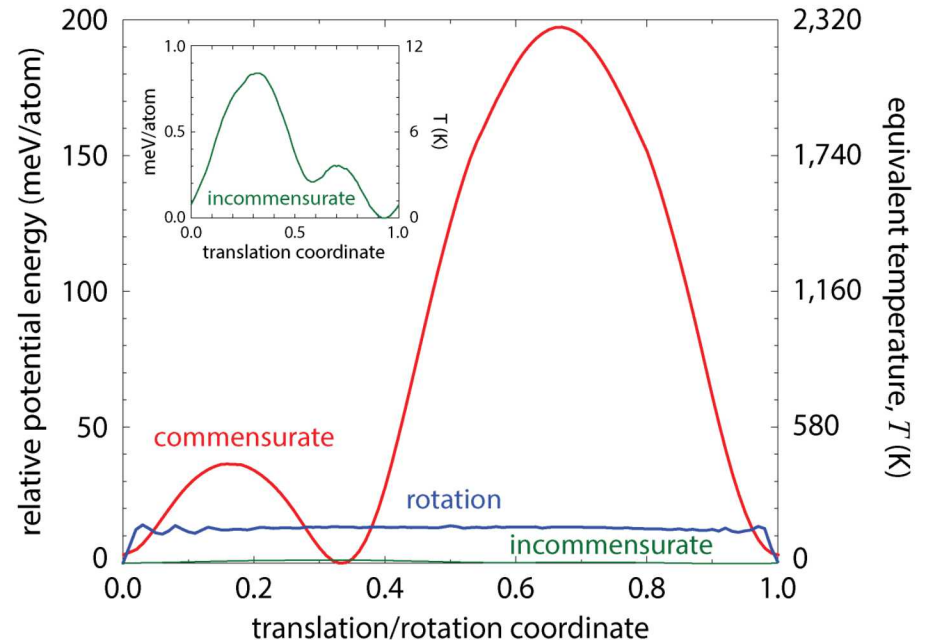
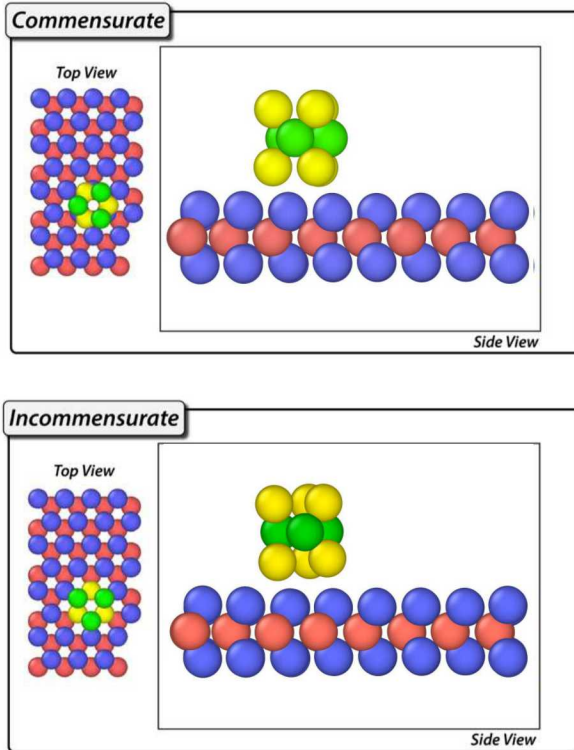


- Six loads at each temperature
- $\mu = dF_f / dF_n$ gives friction coefficient
- Contact conditions $\Rightarrow A = \beta F_N$, can use to calculate shear stress



- MD has more defects, expect higher μ
- Functional form is the same

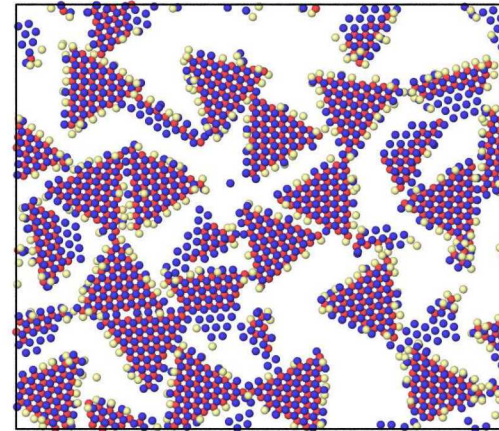
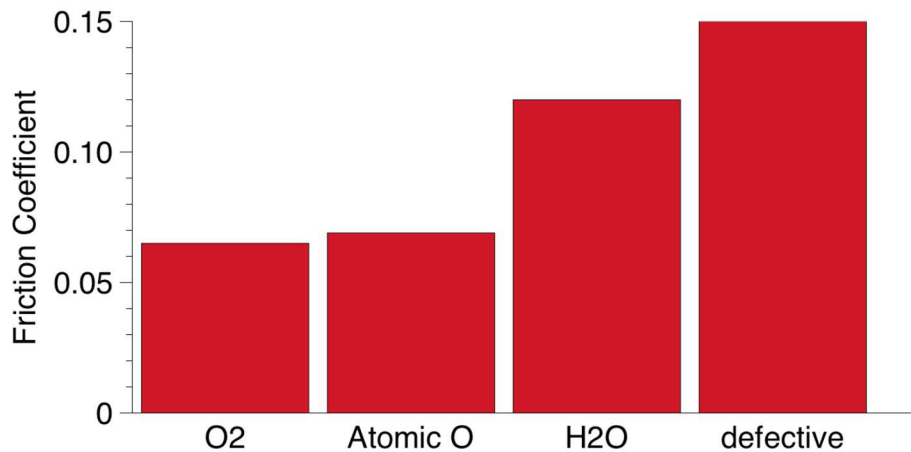
Commensurate vs. Incommensurate Sliding



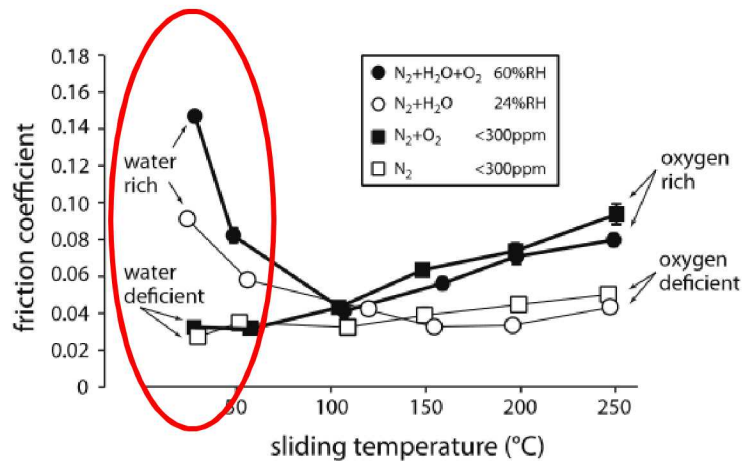
rotation

- Commensurate barrier ~ 300 K
- Incommensurate barrier ~ 10 K
- Rotation barrier ~ 150 K

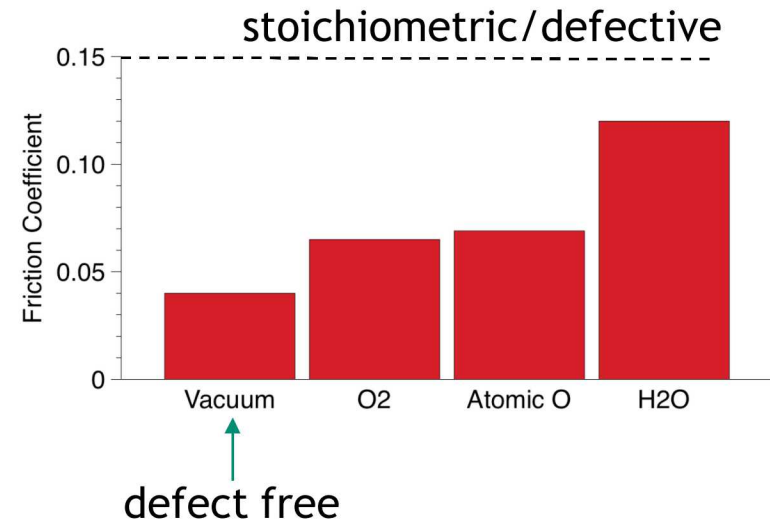
What happens with oxygen and water?



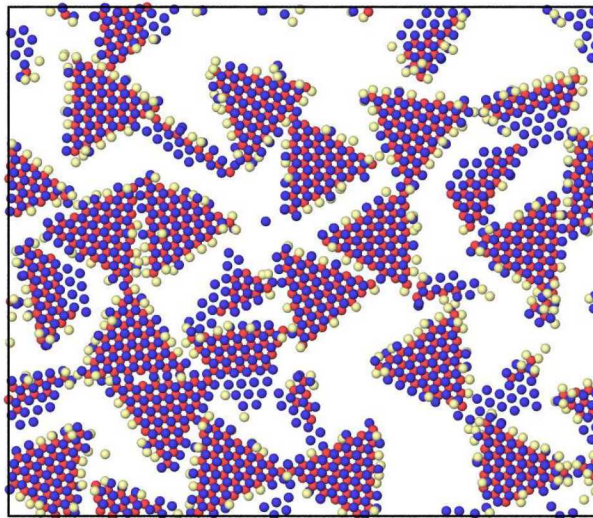
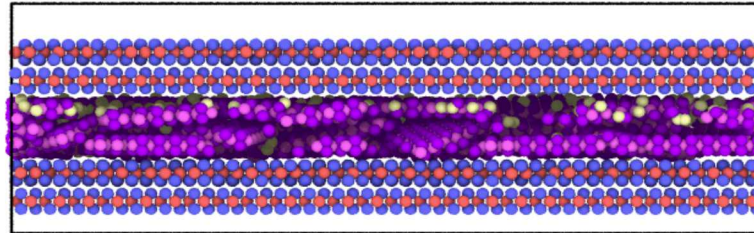
- Friction goes down?
- This is unfair...
 - Water and oxygen passivate defect sites
 - Need to do this in the pure system, too
 - Look at non-stoichiometric (i.e. defect-free) nanoplatelets



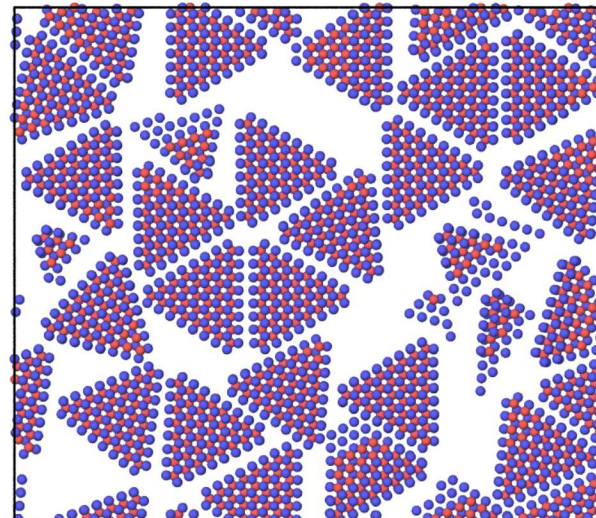
Khare and Burris, Tribol. Lett. 2013



- Changes with added oxygen or water match experimental results

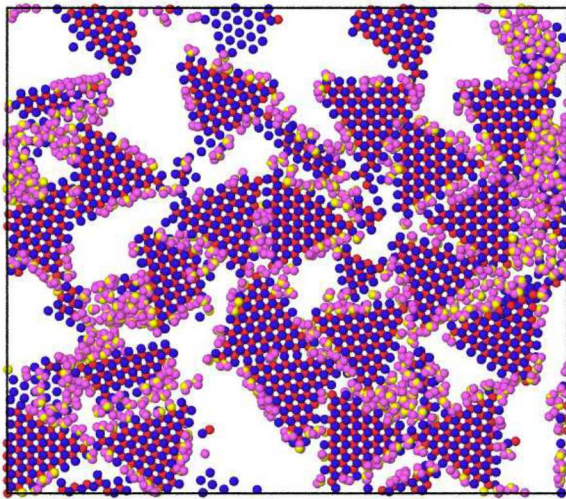


oxygen passivated

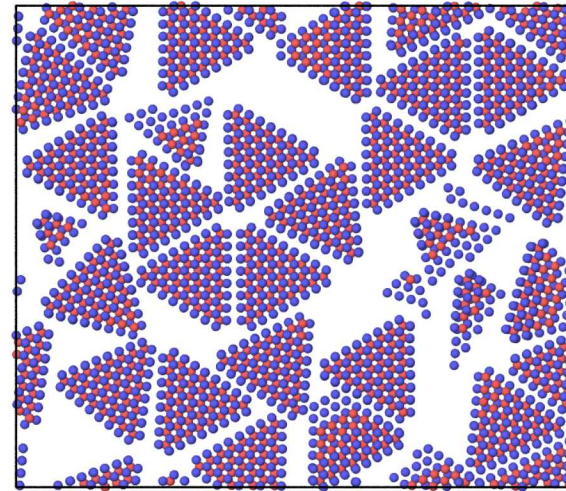


defect free

- Oxygen bonds to defect sites & prevents formation of larger sheets
- Molecular oxygen looks very similar



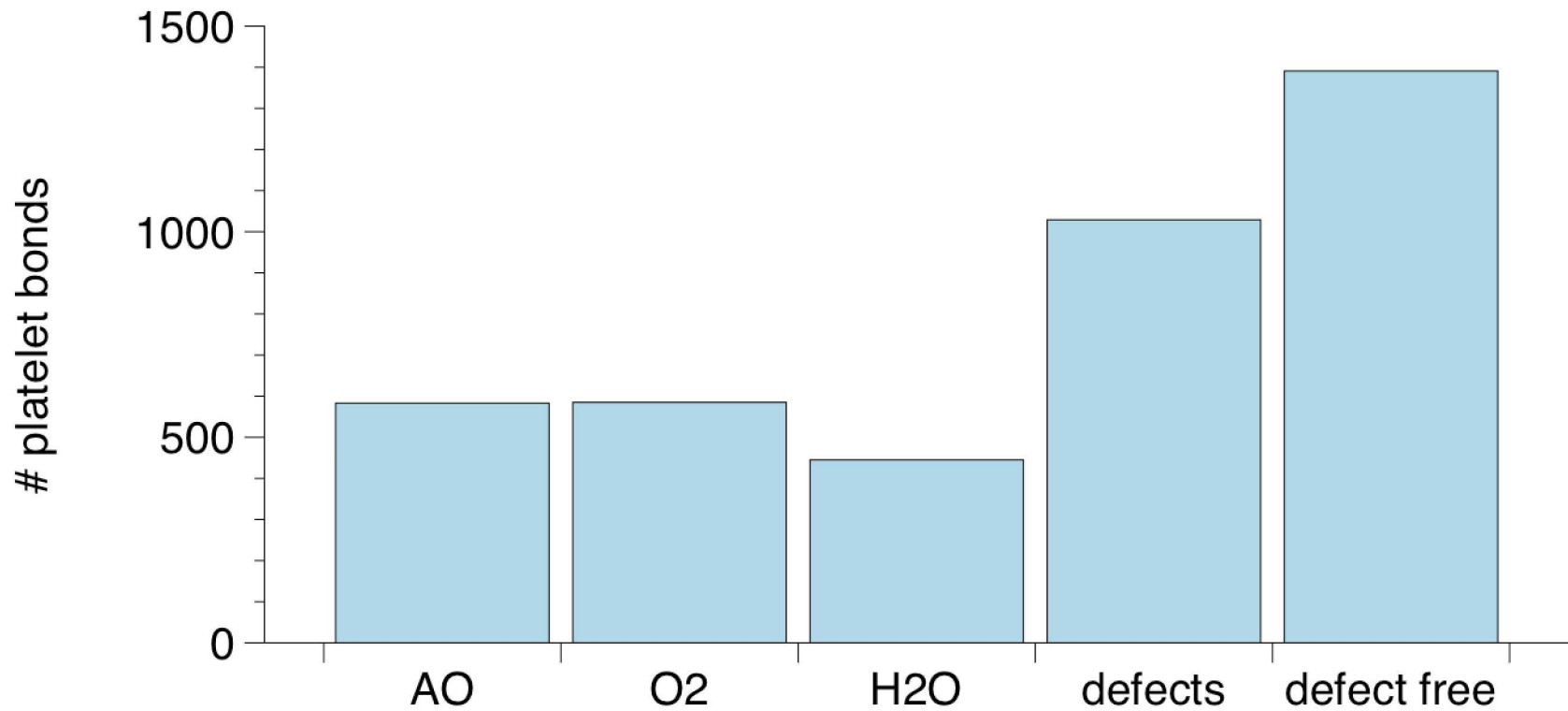
water passivated



defect free

- Water also bonds to defect sites & prevents formation of larger sheets
- Water aggregates with itself more than oxygen does

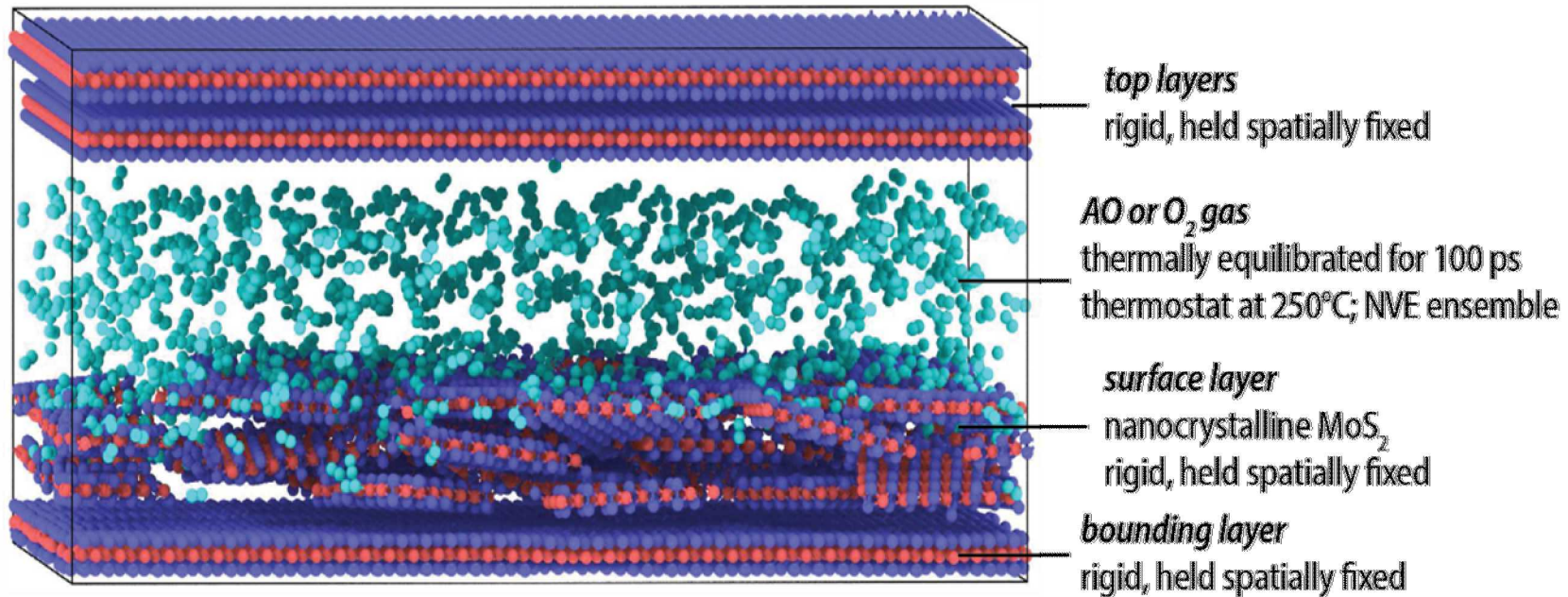
Counts of inter-platelet bonds confirm



Environmental species interrupt formation of larger flakes

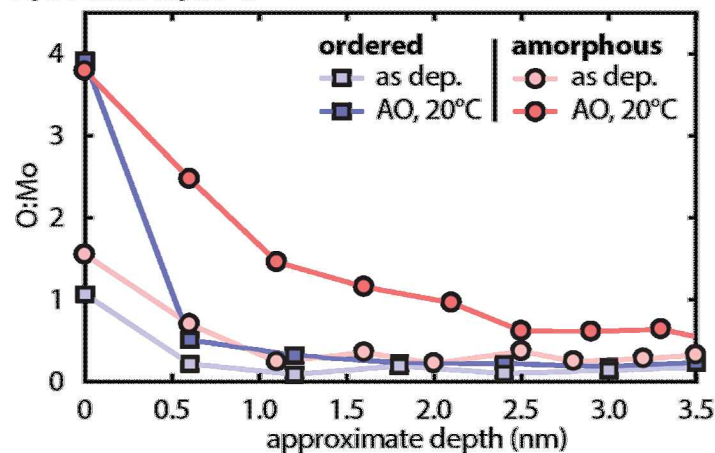
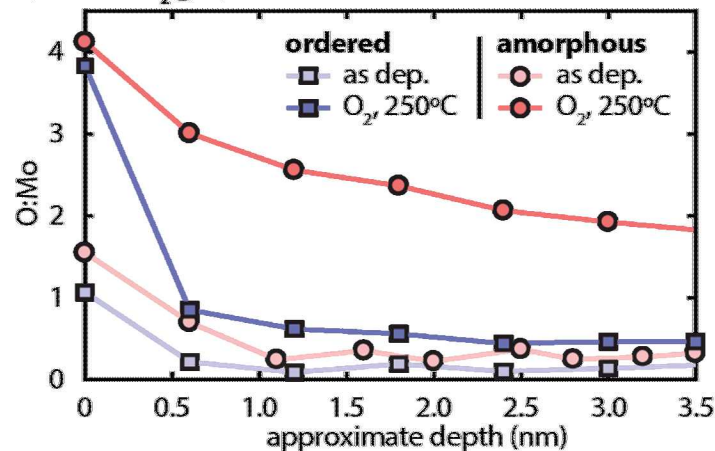
- MoS₂ shows purely elastic contact
- Shear is predominantly due to inter-lamellar interactions
- MD calculates correct shear strengths as a function of temperature
- Developed simple model based on probabilities:
 - Energy barriers determine the shear strength
 - Rotate, and slide incommensurately
 - Fail to rotate and slide commensurately
- Incommensurate sliding is the most important – can neglect commensurate
- Simple model predicts temperature dependence

What about chemistry?

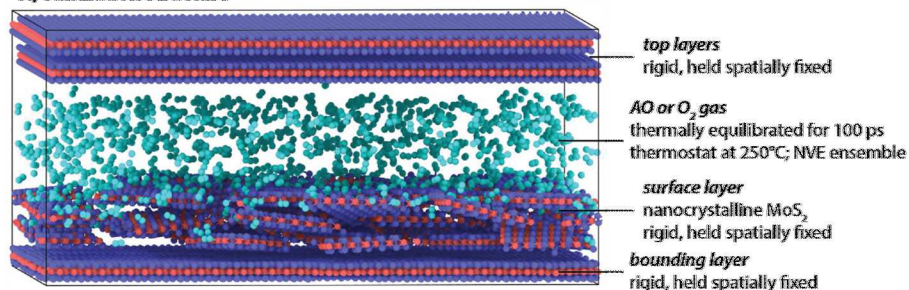


- Take systems that have “run-in” (i.e. reached steady-state shearing)
- Remove top layers
- Apply O₂, AO or H₂O at 100 atm
- Replace top layers

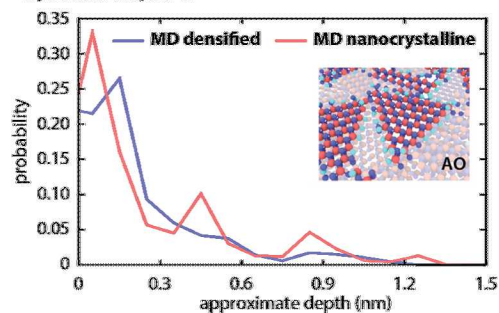
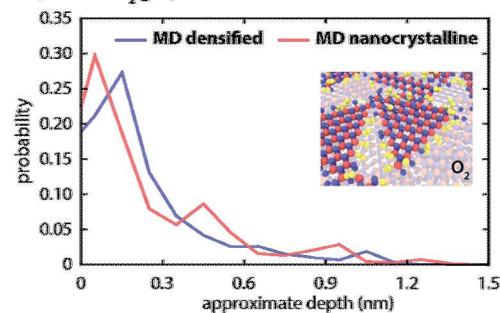
A) 30 min AO, 20°C

B) 30 min O₂ gas, 250°C

A) Simulation structure



B) 30 min AO, 20°C

C) 30 min O₂ gas, 250°C

Curry, et al, ACS Appl. Mater. Interfaces, 2017

MD accurately represents oxygen depth profiles as seen in LEIS experiments