

MULTI-DIMENSIONAL FEMTOSECOND-LASER INDUCED DYNAMICS OF CO ON METALS:

ACCOUNTING FOR ELECTRONIC FRICTION AND SURFACE MOTION WITH COMBINED MODELS

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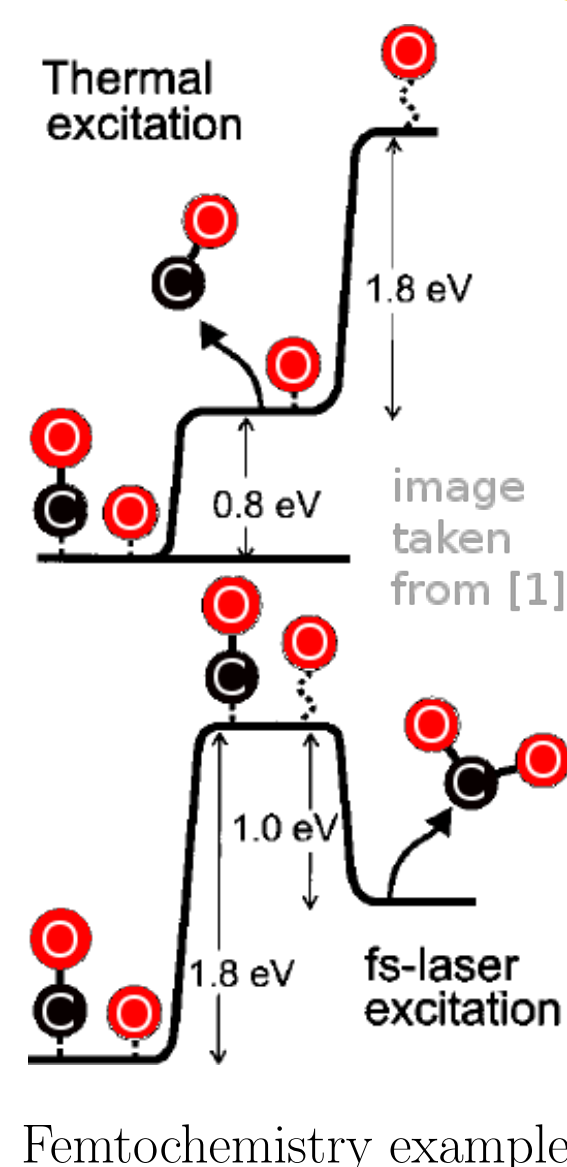
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Introduction

Motivation

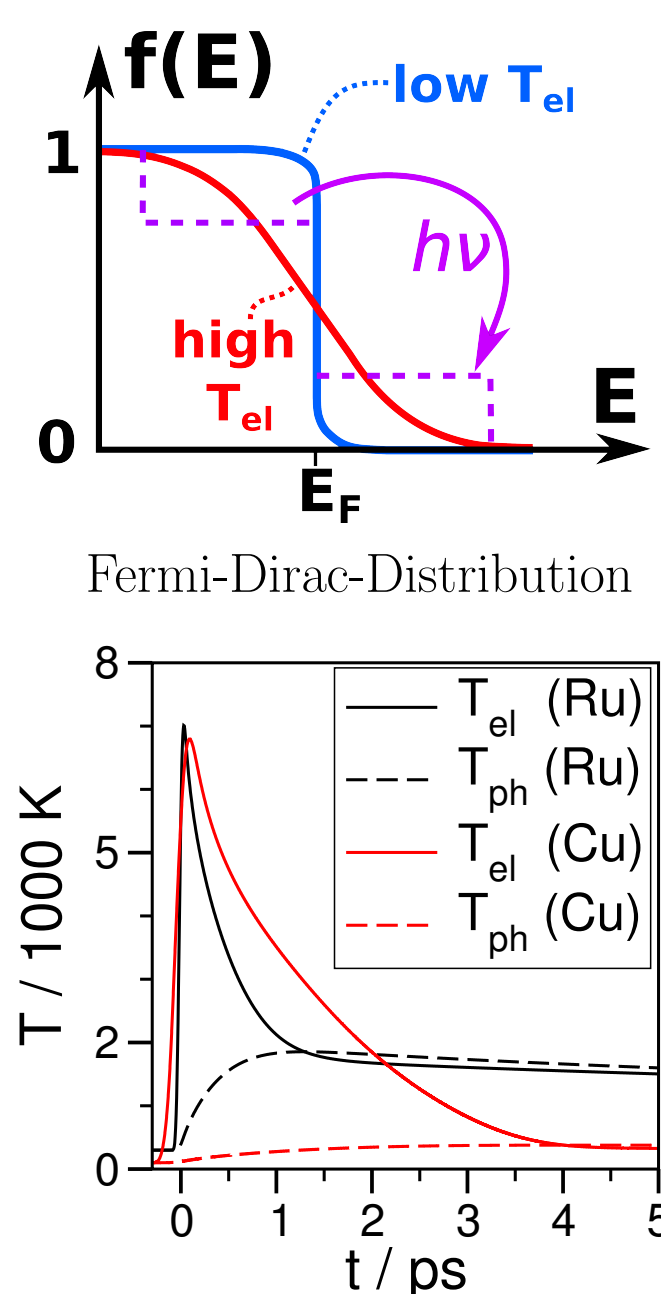
- **Aim:** gain **precise understanding** of **adsorbate bonding** on metals
⇒ Important for **Catalysis**
- **Why femtosecond(fs)-lasers?**
 - produce non-equilibrium **2-T-states**
⇒ **different than** normal heating
 - further **tool** besides STM and scattering
 - **direct** future **applications** possible
⇒ “**femtochemistry**”[1]
- **Why CO/Ru(001) and CO/Cu(100)?**
 - both are well studied **model systems**
 - recently, interesting **fs-laser experiments**[2][3]
 - also, **ab-initio** based **6-dim. potentials** available [4][5]



Femtochemistry example

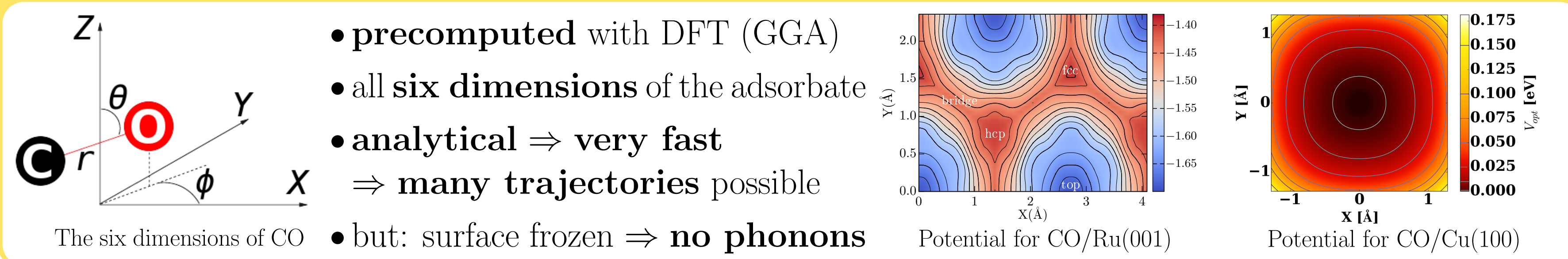
How do fs-Lasers affect Adsorbate-Metal Systems?

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- only **electrons** of metal absorb laser
 - **electron-hole pairs** thermalize fast
⇒ “**hot**” Fermi-Dirac-distribution
 - electrons transfer energy to ion lattice, via ① **electron-phonon coupling**
 - **equilibration within ps-timescale**
⇒ Thus, for few ps **two temperatures**:
 - T_{el} - electron temperature
 - T_{ph} - phonon temperature
 - both can **couple** to adsorbed **molecule**
 - low electron heat **capacity** ⇒ T_{el} higher



Models and Methods

Basis of the Dynamics: the Six-dimensional Potential Energy Surfaces (PES)[4][5]



Two-Temperature Model (2TM)[6]

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- simulates **interaction** of **electrons** with **phonons**
and **laser** ⇒ gives T_{el} and T_{ph} as $f(z, t)$
 - electron and phonon heat capacities C_{el} and C_{ph}
 - elec. heat conductivity κ
 - elec.-phonon coupling g
 - laser source term $S(z, t)$

$$C_{el} \frac{\partial T_{el}}{\partial t} = \frac{\partial}{\partial z} \kappa \frac{\partial T_{el}}{\partial z} - g(T_{el} - T_{ph}) + S(z, t),$$

$$C_{ph} \frac{\partial T_{ph}}{\partial t} = g(T_{el} - T_{ph}).$$

Electronic Friction: Langevin Dynamics[7] and Local Density Friction Approximation (LDFA)[8]

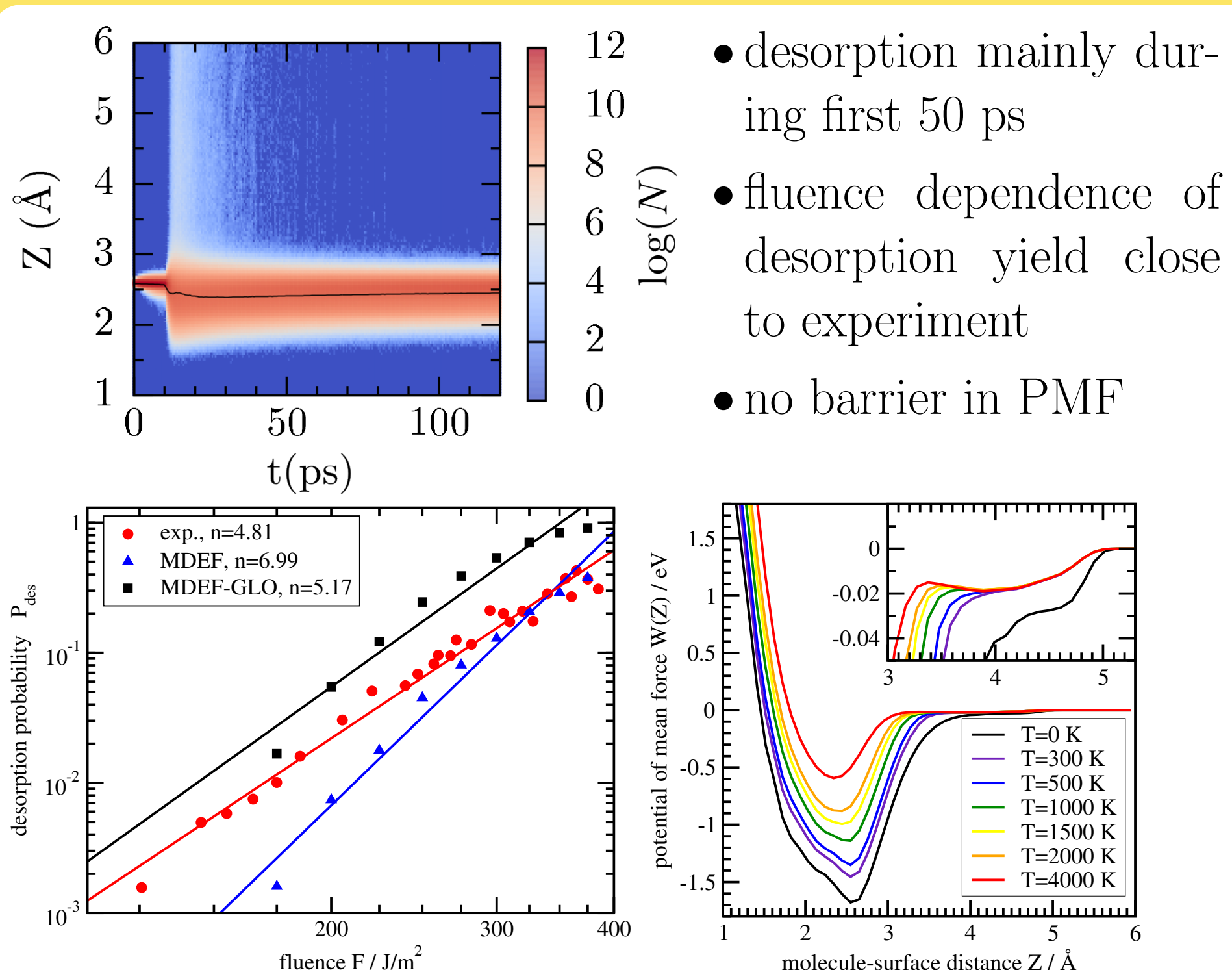
-
- the **Langevin** equation of motion, a **stochastic** differential equation:
- $$m_k \frac{d^2 \underline{r}_k}{dt^2} = \underbrace{-\nabla_k V(\underline{r}_1, \underline{r}_2)}_{\text{Force on Atom } k} - \underbrace{\eta_{el,k}(\underline{r}_k)}_{\text{Force due to PES}} \frac{d \underline{r}_k}{dt} + \underbrace{\underline{R}_{el,k}(t)}_{\text{Friction force slows movement}} + \underbrace{\underline{R}_{el,k}(t)}_{\text{Random force from e-h pairs}}$$
- describes **movement** of CO and **interaction** with **electron-hole pairs** (friction and excitation)
 - **Local Density Friction Approx.** (LDFA): *ab-initio* model that gives **friction coefficients** $\eta_{el,k}$
 - Atom k embedded in **free electron gas** with density of bare surface at current position \underline{r}_k
 - **Random forces** $\underline{R}_{el,k}$: white noise, **dependent** on both $\eta_{el,k}$ (from LDFA) and T_{el} (from 2TM)

Inclusion of Phonons: Generalized Langevin Oscillator(GLO)-model[9]

-
- influence of **phonons** effectivly modeled (**augments frozen surface**)
 - **entire surface** understood as **3D oscillator** (coords. \underline{r}_s , mass 1 atom)
 - **coupling** to molecule **via shifting**: $V_{GLO}(\underline{r}_C, \underline{r}_O; \underline{r}_s) = V(\underline{r}_C - \underline{r}_s, \underline{r}_O - \underline{r}_s)$
 - additionally coupled to **ghost oscillator** \underline{r}_g , **models** influence of **bulk**
 - ghost oscillator is subject to **friction** η_{ph} and **random forces** $\underline{R}_{ph}(T_{ph})$

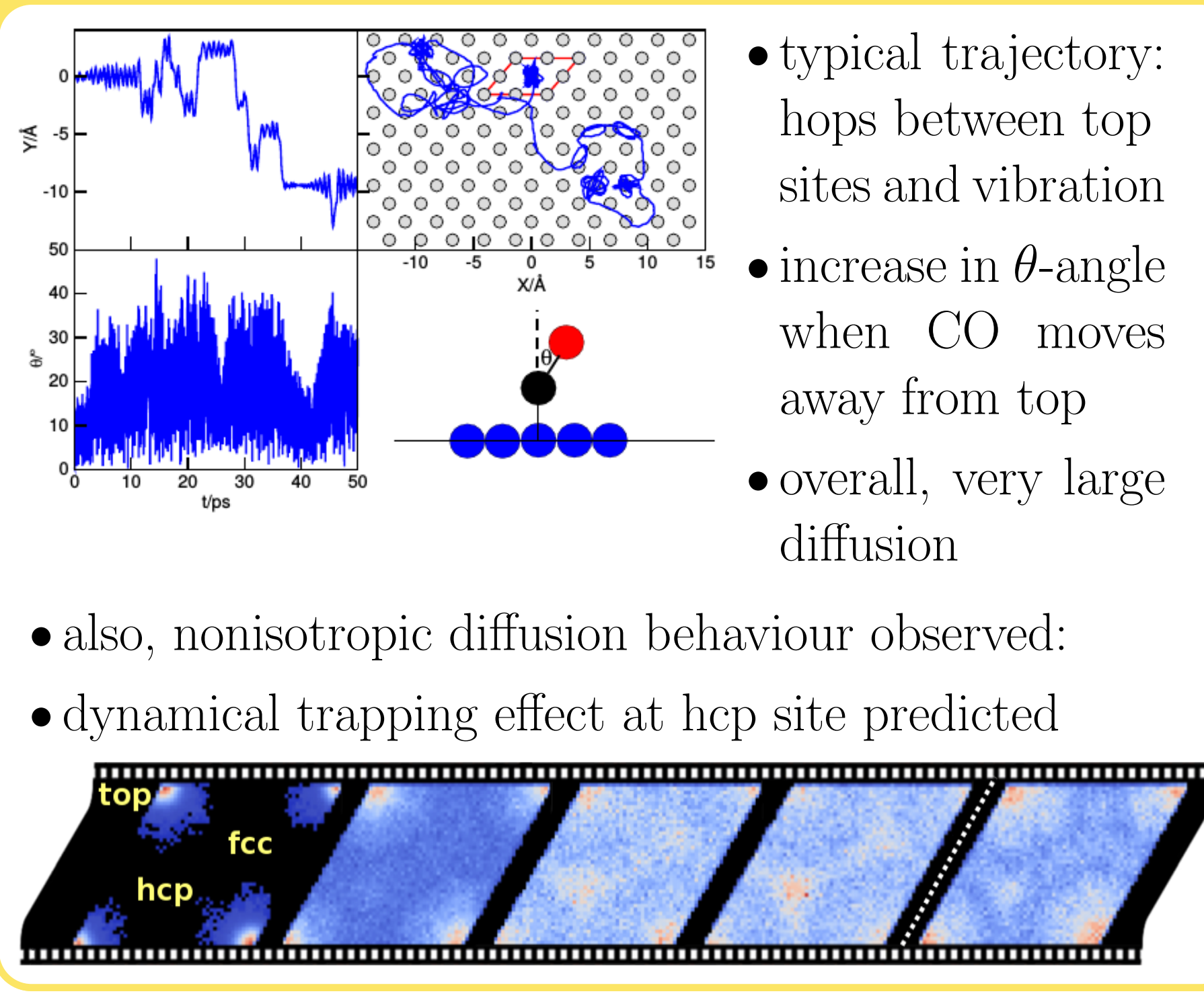
Results

Desorption (Data for Ru)



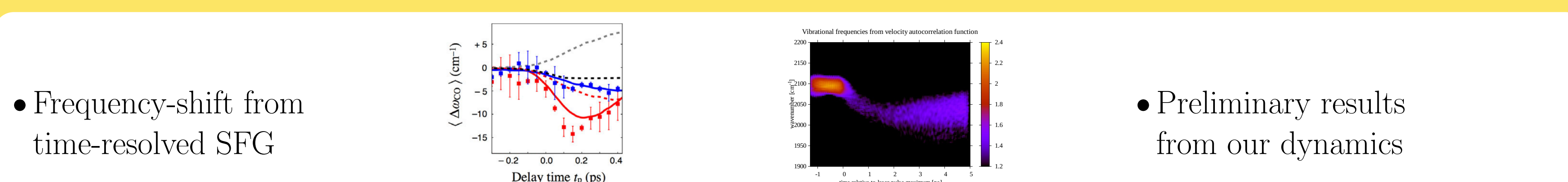
- desorption mainly during first 50 ps
- fluence dependence of desorption yield close to experiment
- no barrier in PMF

Diffusion (Data for Ru, but Cu similar)



- typical trajectory: hops between top sites and vibration
- increase in θ -angle when CO moves away from top
- overall, very large diffusion
- also, nonisotropic diffusion behaviour observed:
- dynamical trapping effect at hcp site predicted

Vibrations (Data for Cu)



- Frequency-shift from time-resolved SFG

- Preliminary results from our dynamics

Conclusions

- 6D Langevin dynamics of CO on Ru and Cu
- based on first principles, no “free” parameters
- accounting for (via LDFA) electronic friction, hot electron excitation and (via GLO) substrate motion
- allows for detailed time- and space-resolved insights
- no physisorbed state, molecules desorb directly

Outlook

- better electronic friction ($\eta(T_{el})$) and beyond LDFA
⇒ Long term goal: use tensorial friction (exact)
- non-equilibrium lattice model (NLM) instead 2TM
- simulate other coverages (bigger super cells)
- simulate bigger systems (CO + H; hydrocarbons)
- include interaction between adsorbate molecules

References

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