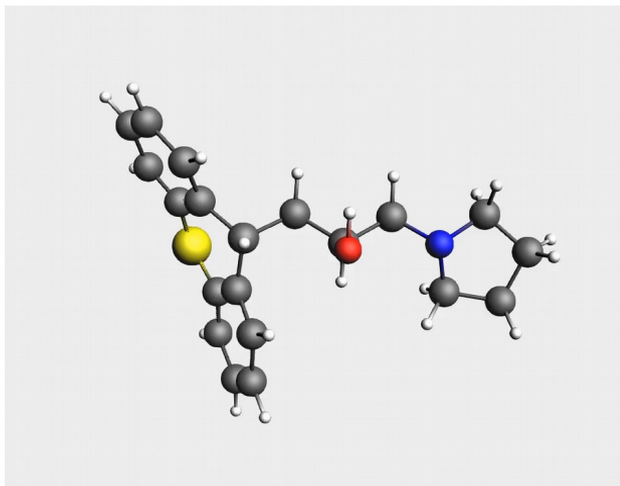




# Amsterdam Modeling Suite 2023

Making Computational Chemistry Work For You



Conformational search  
CREST, any engine

Matti Hellström, Product Manager

hellstrom@scm.com

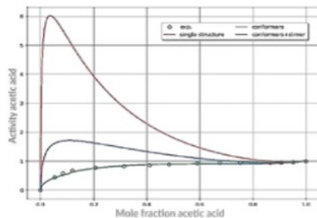


Full release notes: [www.scm.com/2023](http://www.scm.com/2023)

# Amsterdam Modeling Suite: AMS



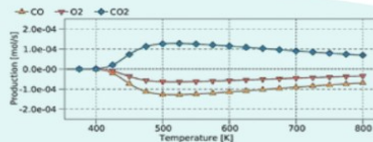
Continuum



## Fluid Thermodynamics

COSMO-RS  
COSMO-SAC  
UNIFAC

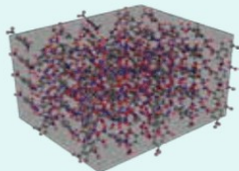
Mesoscale



## Kinetics

Kinetic Monte Carlo  
Microkinetics

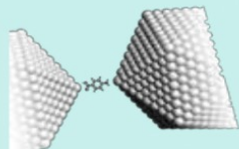
Materials



## Force Fields

ReaxFF, GFN-FF  
Machine Learning Potentials  
Apple & P

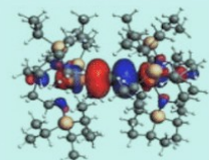
Nano



## QM/MM

FDE, Hybrid Engine

Atomistic



## Tight binding

GFN-xTB, DFTB

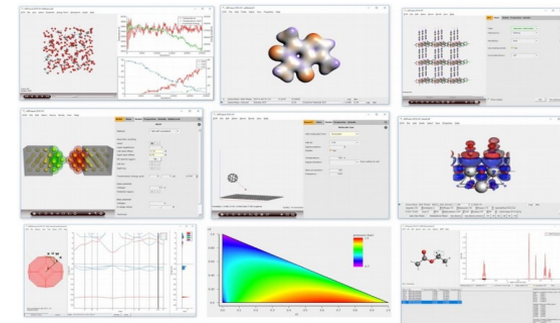
## Periodic DFT

BAND, Quantum Espresso

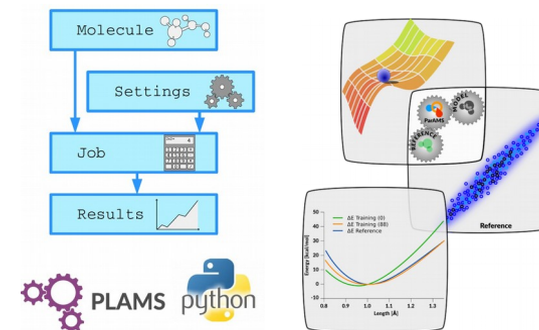
## Molecular DFT

ADF

## Graphical User Interface



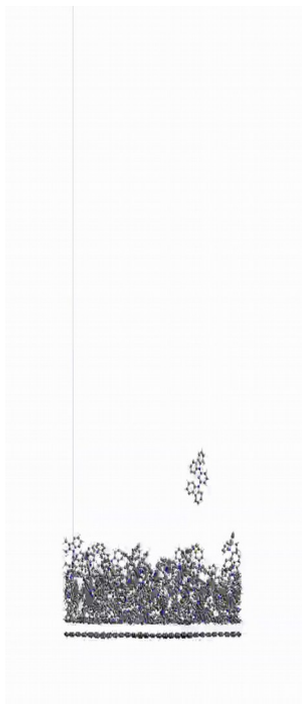
## Python scripting, workflows



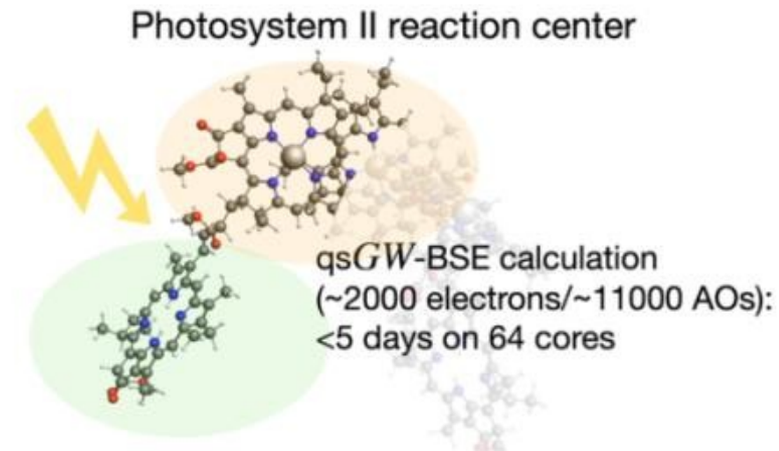
- ▶ Excitations and charge transport
- ▶ New methods/functionals in ADF
- ▶ M3GNet and ANI-2x machine learning potentials
- ▶ Conformers
- ▶ APPLE&P polarizable force field
- ▶ ACE Reaction Network
- ▶ Multiscale Modeling for Heterogeneous Catalysis
- ▶ ParAMS for ReaxFF/DFTB parametrization
- ▶ New engine: ASE calculators
- ▶ New engine: Quantum ESPRESSO (alpha)
- ▶ Python tips, PLAMS, Jupyter Lab

Full release notes: [www.scm.com/2023](http://www.scm.com/2023)

- ▶ OLED workflow “better and faster”
- ▶ GPU acceleration
- ▶ Ionization potentials
- ▶ Electron affinities
- ▶ Exciton energies



- ▶ Excitation energies with qsGW-BSE



Arno Förster, Lucas Visscher  
*Quasiparticle Self-Consistent GW-Bethe-Salpeter Equation  
Calculations for Large Chromophoric Systems*  
J. Chem. Theory Comput. 2022, 18, 11, 6779–6793

Also with **spin-orbit coupling** (article in progress)

# New methods/functionals in ADF

## ▶ r<sup>2</sup>SCAN-3c

- ▶ “Reliable, robust, and accurate”
- ▶ “on par with or more accurate than hybrid M06-2X-D3(0)/TZP”

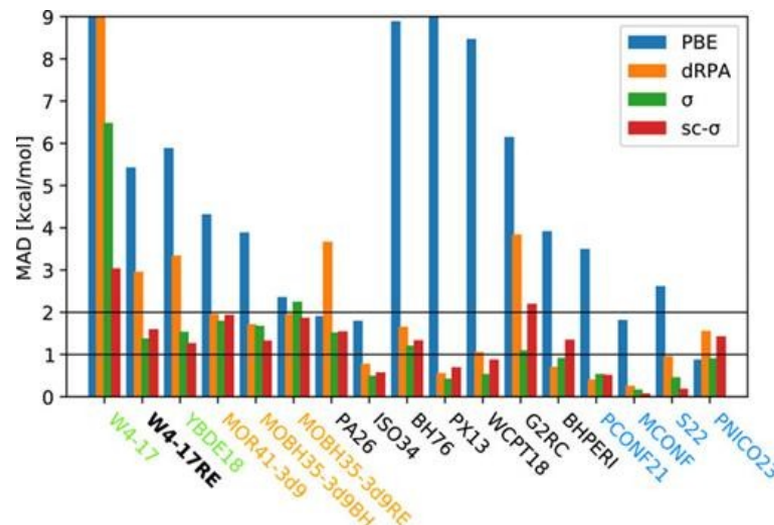


mTZ2P ✓  
SR-ZORA ✓  
gCP ✓  
D4 ✓

T. Gasevic, J.B. Stückrath, S. Grimme, M. Bursch  
*Optimization of the r<sup>2</sup>SCAN-3c Composite Electronic-Structure Method for Use with Slater-Type Orbital Basis Sets*  
J. Phys. Chem. A 2022, 126, 23, 3826–3838

## ▶ Sigma functional

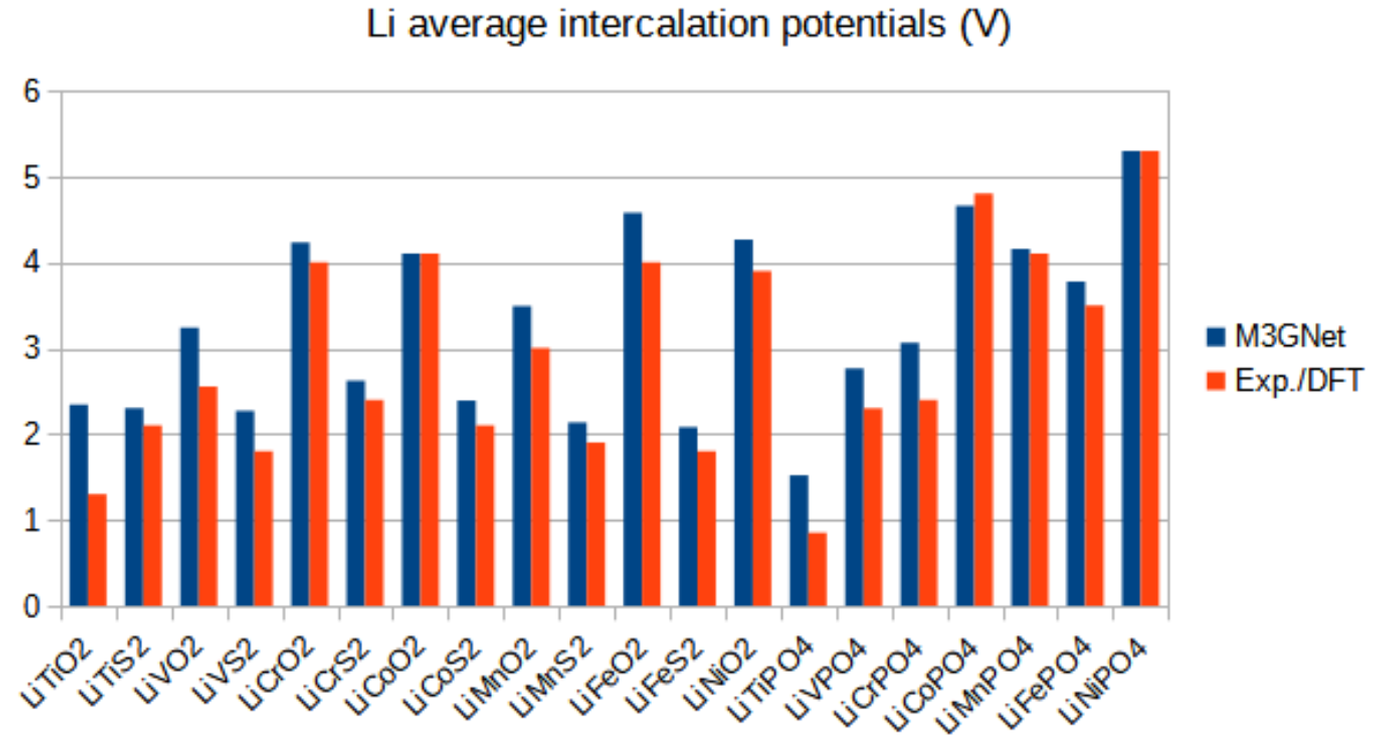
- ▶ “accuracy of 1 kcal/mol in reaction or transition state energies”



J. Erhard, S. Fauser, E. Trushin, A. Görling  
*Scaled  $\sigma$ -functionals for the Kohn-Sham correlation energy with scaling functions from the homogeneous electron gas*  
J. Chem. Phys. 157, 114105 (2022)



- ▶ Equilibrium structures
- ▶ Formation energies
- ▶ Phase stabilities
- ▶ Lattice optimizations
- ▶ ML: Do not blindly trust results!



**M3GNet-UP-2022**  
Materials  
*Phase stabilities*  
*Reaction energies*  
H-Pu

## A universal graph deep learning interatomic potential for the periodic table

[Chi Chen](#)  & [Shyue Ping Ong](#) 

*Nature Computational Science* **2**, 718–728 (2022) | [Cite this article](#)

1662 Accesses | 3 Citations | 180 Altmetric | [Metrics](#)

**ANI-2x**  
Drug-like molecules  
*Conformers*  
*Reaction energies*  
H, C, O, N, F, S, Cl

## Extending the Applicability of the ANI Deep Learning Molecular Potential to Sulfur and Halogens

Christian Devereux, Justin S. Smith\*, Kate K. Huddleston, Kipton Barros, Roman Zubatyuk, Olexandr Isayev\*, and Adrian E. Roitberg\*

 **Cite this:** *J. Chem. Theory Comput.* 2020, 16, 7, 4192–4202

Publication Date: June 16, 2020 

<https://doi.org/10.1021/acs.jctc.0c00121>

Copyright © 2020 American Chemical Society

[RIGHTS & PERMISSIONS](#)

Article Views

4355

Altmetric

21

Citations

80

[LEARN ABOUT THESE METRICS](#)

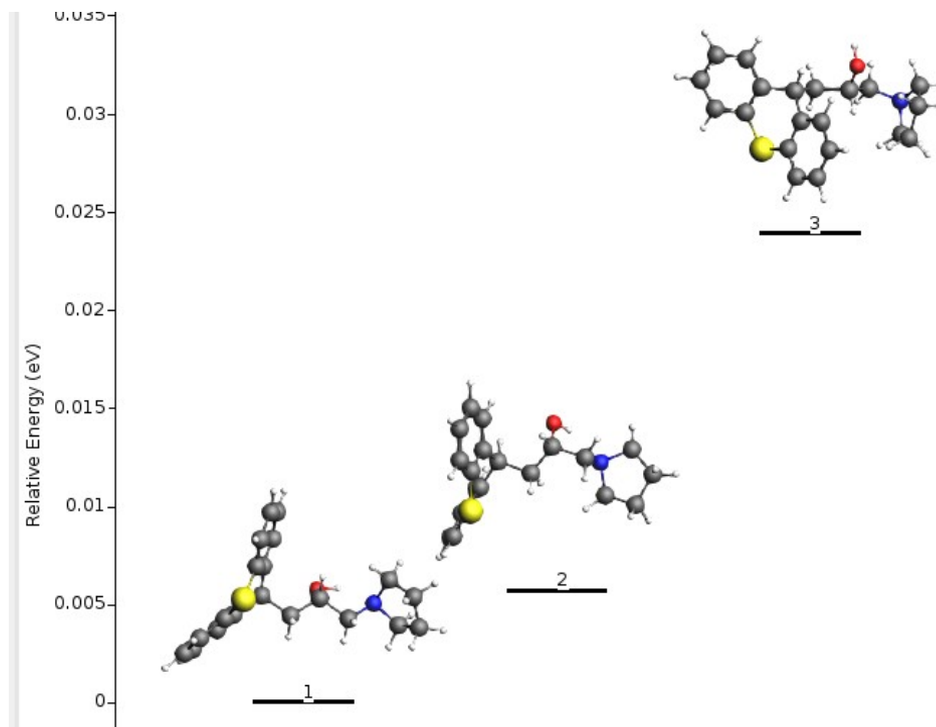
Share Add to Export



# Conformers

- ▶ Generate with Rdkit (fast) or CREST (thorough)
- ▶ Use ANI-2x, GFN-FF, DFTB, ADF, or any other AMS methods to generate, optimize, and score conformers
- ▶ Calculate Boltzmann-averaged properties, for example IR spectra!

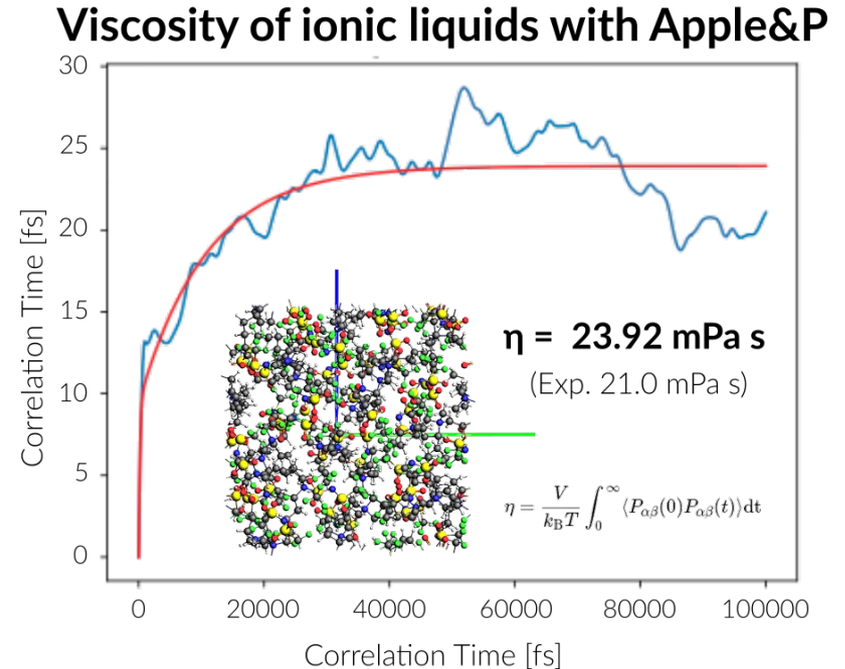
RDKit: Open-source cheminformatics; <http://www.rdkit.org>



P. Pracht, F. Bohle, S. Grimme.  
*Automated exploration of the low-energy chemical space with fast quantum chemical methods*  
Phys. Chem. Chem. Phys. 2020, 22(14):7169-7192

- ▶ Collaboration with Wasatch Molecular Inc.
- ▶ Non-reactive
- ▶ Automatic atom typing
- ▶ Organic molecules: amides, esters, ...
- ▶ Solvents: nitriles, carbonates, sulfones, ...
- ▶ Cations: Li<sup>+</sup>, hydraziniums, pyrrolidiniums, ...
- ▶ Anions: hydroxide, TFSI, ...

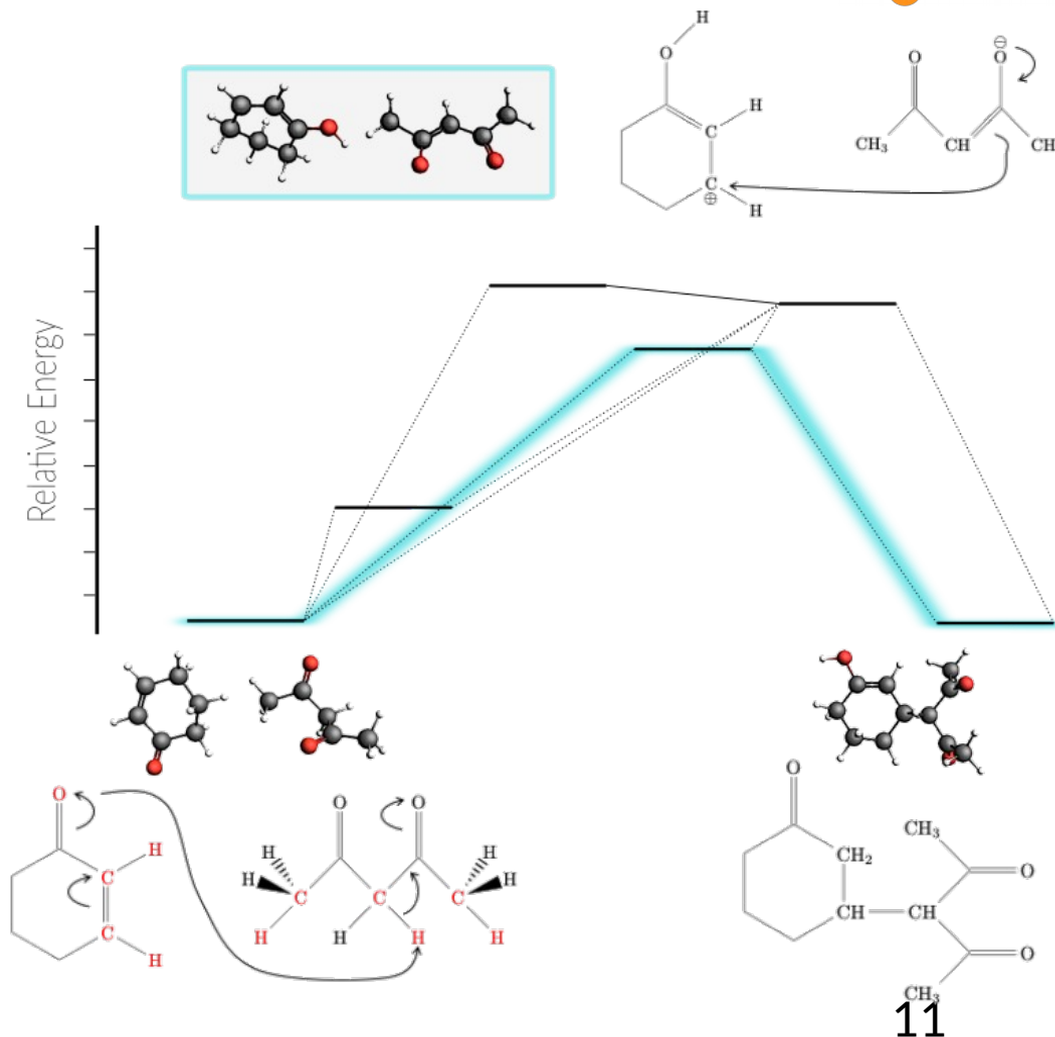
O. Borodin, J. Phys. Chem. B 113, 11463–11478 (2009)  
*and many subsequent publications*



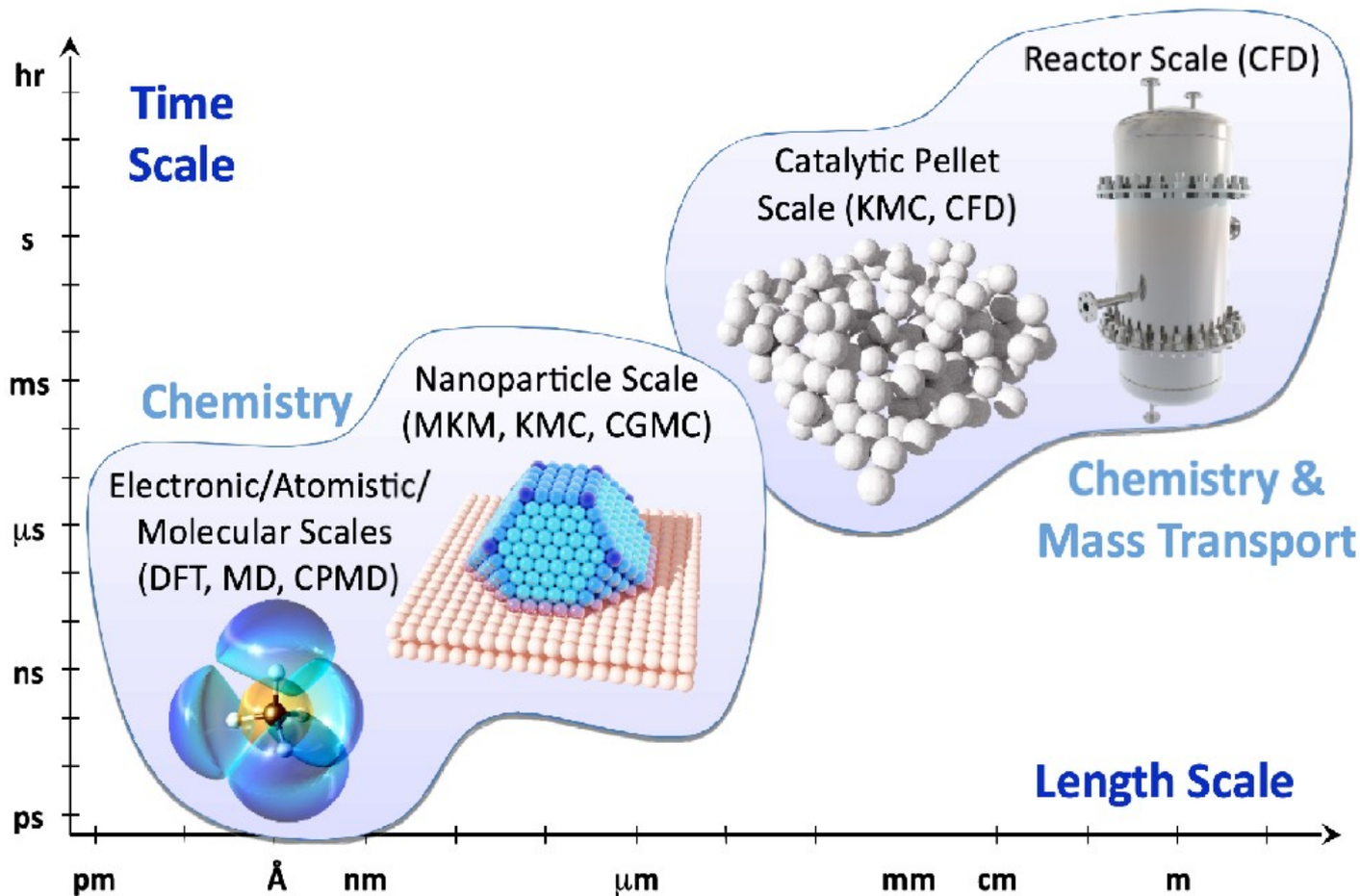
# ACE Reaction Network

- ▶ Quickly generate and minimize reaction networks
- ▶ Find reaction mechanisms

Y. Kim, J.W. Kim, Z. Kim, W.Y. Kim,  
*Efficient Prediction of Reaction Paths through  
Molecular Graph and Reaction Network Analysis*  
Chem. Sci. 2018, 9, 825-835.



# Multiscale Modeling for Heterogeneous Catalysis (1/2)



More information:  
[www.reaxpro.eu](http://www.reaxpro.eu)

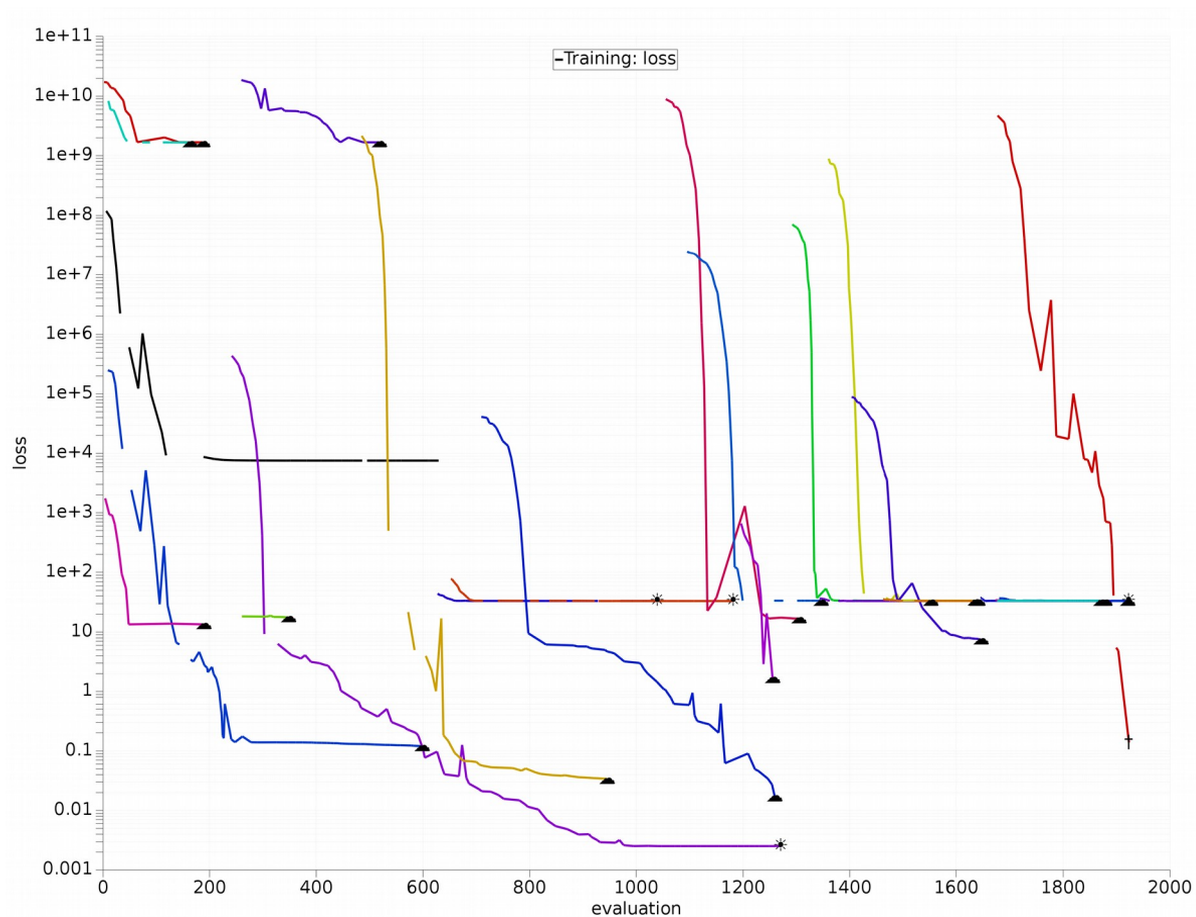


# ParAMS for ReaxFF and DFTB parametrizations (1/2)

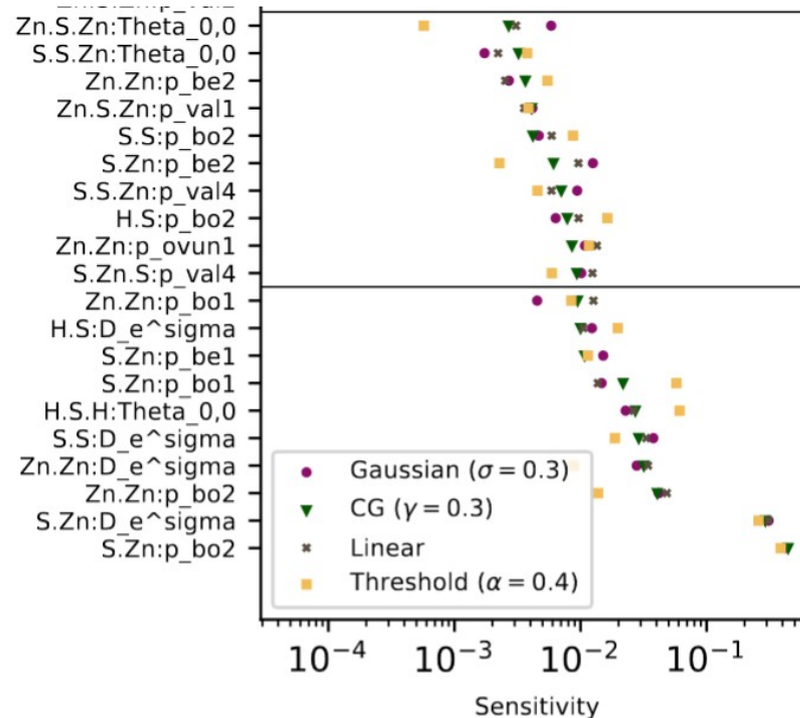
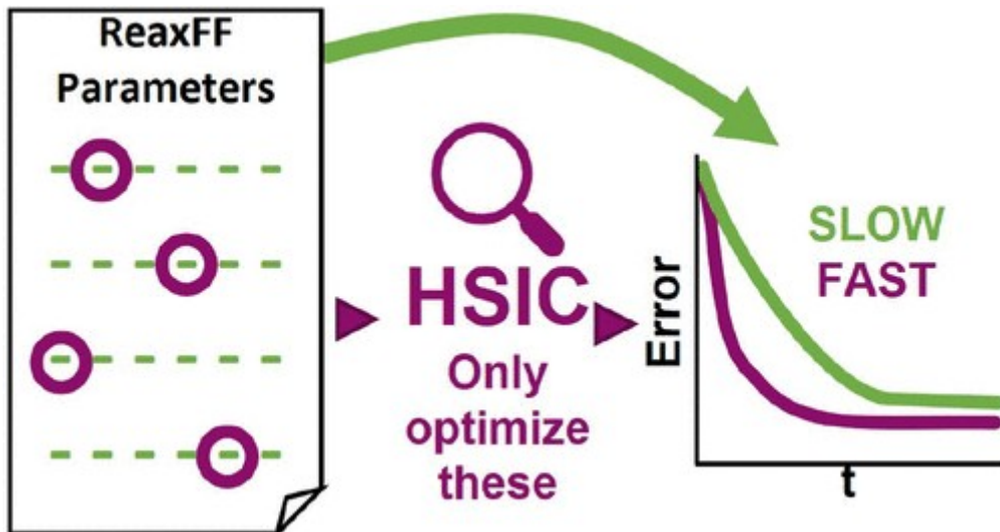


- ▶ Run and control  
multiple optimizers  
in parallel

Freitas Gustavo, M., Verstraelen, T.  
*GloMPO (Globally Managed Parallel Optimization):  
A tool for expensive, black-box optimizations,  
application to ReaxFF reparameterizations.*  
*J. Cheminform.* 2022, 14, 7.



► Parameter sensitivity analysis

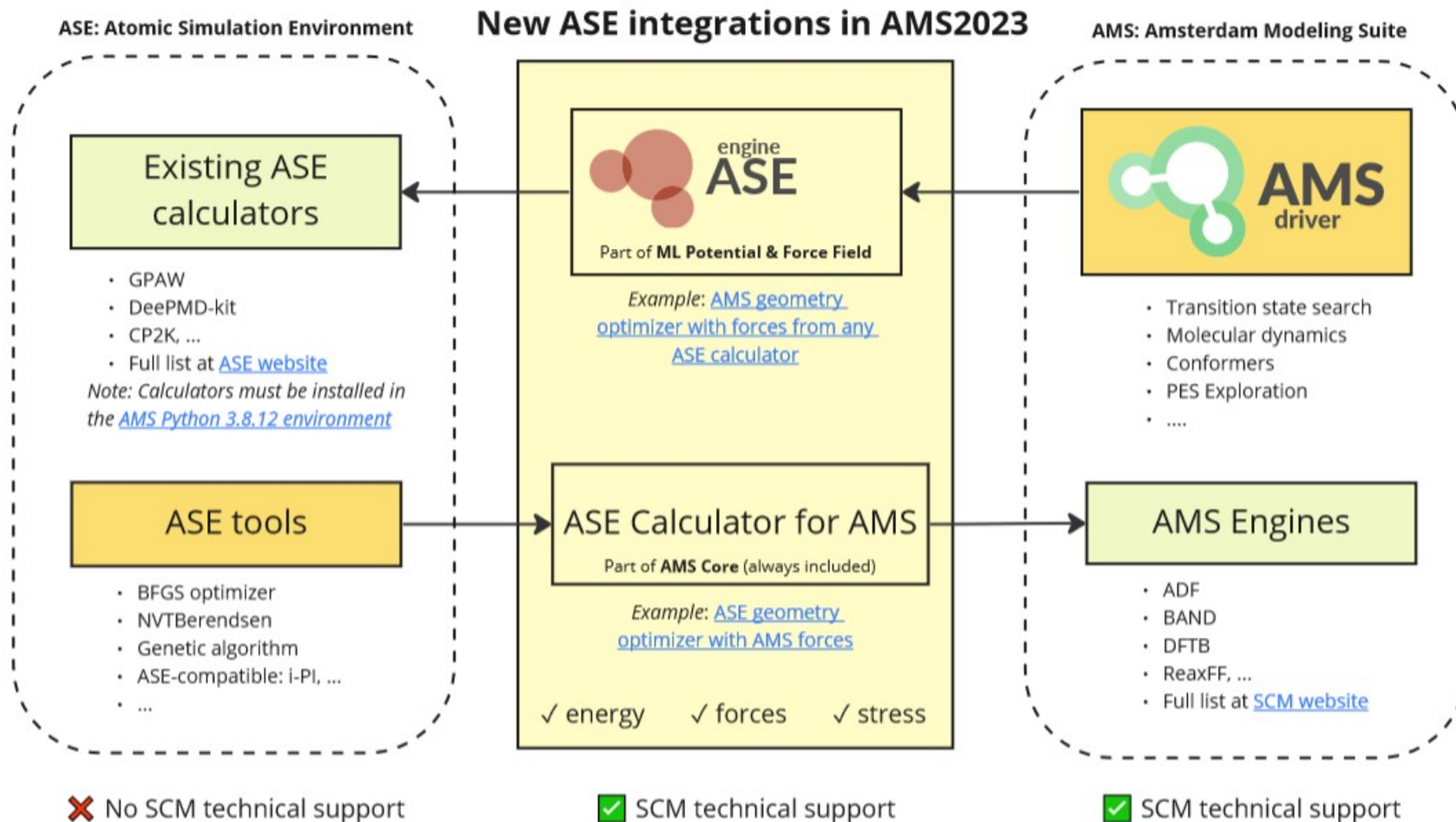


M.F. Gustavo, M. Hellström, T Verstraelen.

*Sensitivity Analysis for ReaxFF Reparametrization Using the Hilbert-Schmidt Independence Criterion*

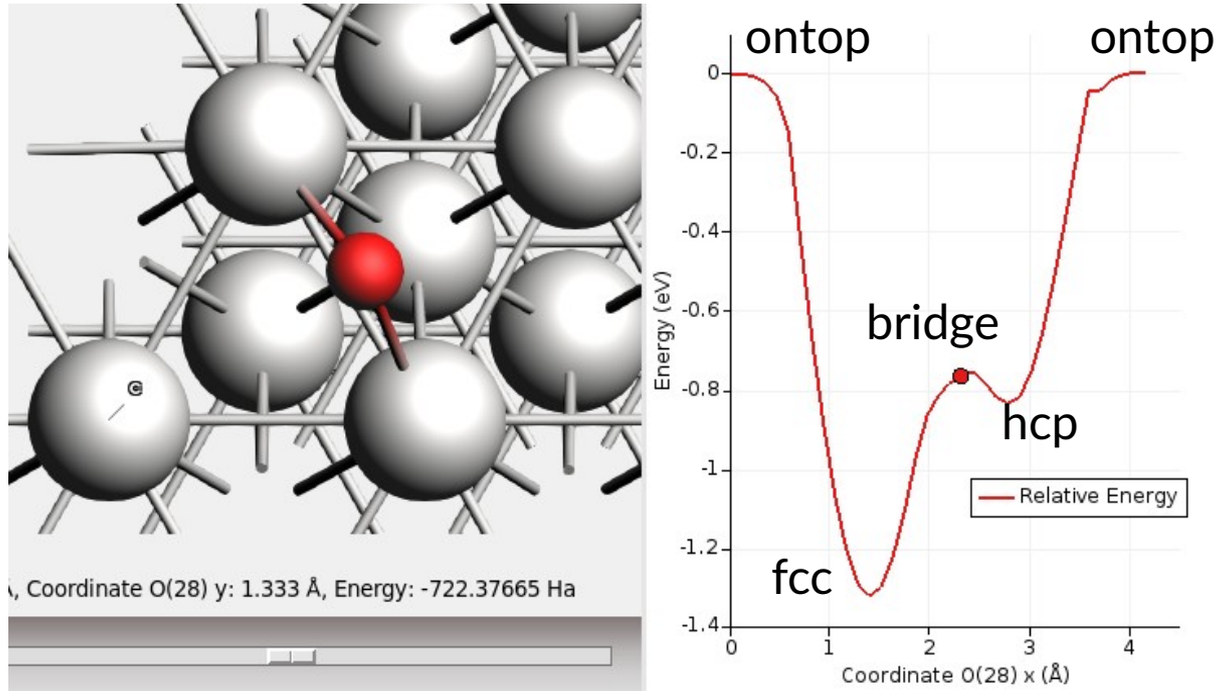
J. Chem. Theory Comput. 2023, 19, 9, 2557-2573

# Atomic Simulation Environment (ASE) interface to/from AMS



- ▶ Expand QE functionality to all AMS Driver tasks
  - ▶ Transition state search
  - ▶ PES Scan
  - ▶ ...
- ▶ For people who prefer QE. The above is also possible with BAND!

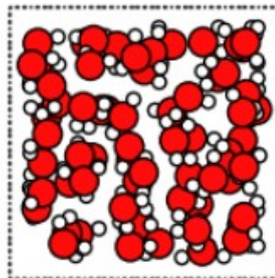
Example PES Scan: O atom diffusion on Pt(111)



- ▶ Install Jupyter Lab with package manager
- ▶ Build, show, and preoptimize structures
- ▶ Run simulations
- ▶ Plot results or launch AMSmovie

```
print('pure liquid with explicit number of molecules and exact density')  
out = packmol(water, n_molecules=64, density=1.0)  
printsummary(out)  
out.write('water-3.xyz')  
show(out)
```

```
pure liquid with explicit number of molecules and exact density  
192 atoms, density = 1.000 g/cm^3, box = 12.417, 12.417, 12.417, formula = H128O64
```



# Licensing for AMS2023



Functionality	License
OLED workflow	ADF + DFTB + Workflows
qsGW-BSE, r2SCAN-3c, sigma functional	ADF
M3GNet, ANI-2x	MLPot&FF
Conformers	engine
APPLE&P	MLPot&FF + special
ACE Reaction	Workflows (+engine)
Zacros	Workflows + special

Functionality	License
ParAMS	Workflows + engine
ASE Engine	MLPot&FF
ASE Calculator for AMS	engine
Quantum ESPRESSO (AMS Engine)	BAND
Quantum ESPRESSO (standalone)	any
Jupyter Lab, PLAMS	any

# Academic collaborators, partners, and contributors for AMS2023



- ▶ Woo Youn Kim (Korea Advanced Institute of Science & Technology): ACE-Reaction
  - ▶ Christine Aikens (Kansas State University): TD-DFT+TB gradients
  - ▶ Artur Michalak (Jagiellonian University): NOCV functionality
  - ▶ Mauro Stener (Trieste University): POLTDDFT
  - ▶ Jochen Autschbach (University at Buffalo): CD with spin-orbit
  - ▶ Laurent Joubert (Rouen Normandy University): QTAIM additions
  - ▶ Martin Kaupp (Berlin Institute of Technology): 3D-RISM entropy
  - ▶ Shyue Ping Ong (University of California San Diego): M3GNet
  - ▶ Kai Leonhard (RWTH Aachen), ChemTraYzer
  - ▶ Toon Verstraelen (Ghent University), ParAMS
  - ▶ Stefan Grimme (Bonn University): r2SCAN-3c, CREST
  - ▶ Arno Förster, Lucas Visscher (VU Amsterdam): BSE
  - ▶ Hannes Jónsson (University of Iceland), EON
  - ▶ Michail Stamatakis (University College London), Zacros
  - ▶ Mauro Bracconi, Matteo Maestri (Polytechnic University of Milan): CatalyticFOAM
- ... and many more!**



# The End

- ▶ Thank you for your attention!
- ▶ Release notes: [www.scm.com/2023](http://www.scm.com/2023)
- ▶ Free trial: [www.scm.com/free-trial](http://www.scm.com/free-trial)
- ▶ YouTube: webinars, tip of the week, ...:  
[www.youtube.com/c/AmsterdamDensityFunctional](http://www.youtube.com/c/AmsterdamDensityFunctional)
- ▶ Questions?

