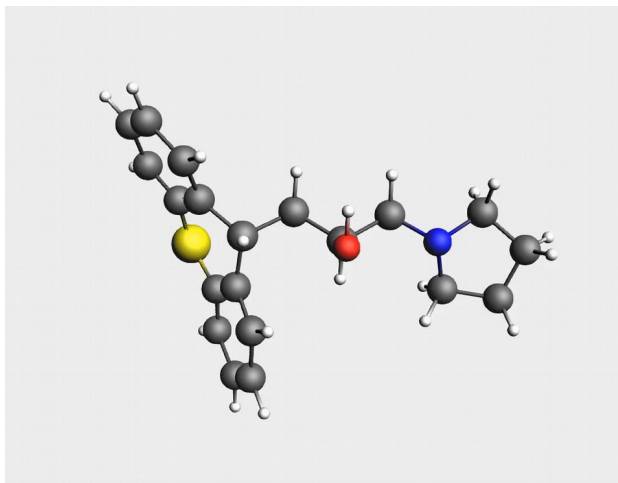




Amsterdam Modeling Suite 2023

Making Computational Chemistry Work For You



Conformational search
CREST, any engine

Matti Hellström, Product Manager

hellstrom@scm.com

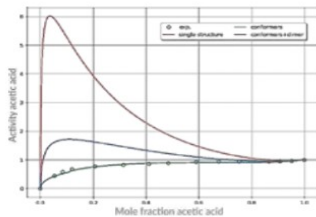


Full release notes: www.scm.com/2023

Amsterdam Modeling Suite: AMS



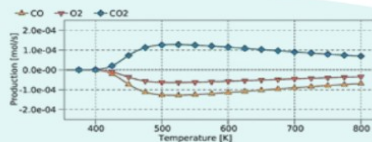
Continuum



Fluid Thermodynamics

COSMO-RS
COSMO-SAC
UNIFAC

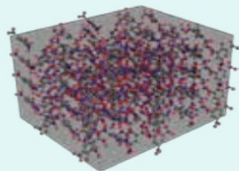
Mesoscale



Kinetics

Kinetic Monte Carlo
Microkinetics

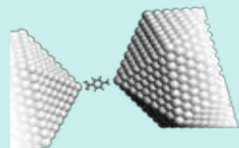
Materials



Force Fields

ReaxFF, GFN-FF
Machine Learning Potentials
Apple & P

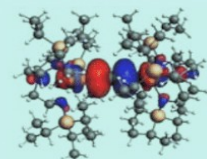
Nano



QM/MM

FDE, Hybrid Engine

Atomistic



Tight binding

GFN-xTB, DFTB

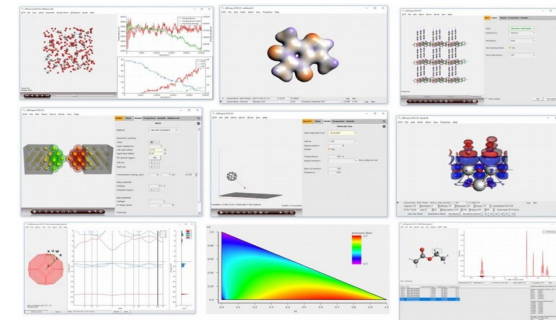
Periodic DFT

BAND, Quantum Espresso

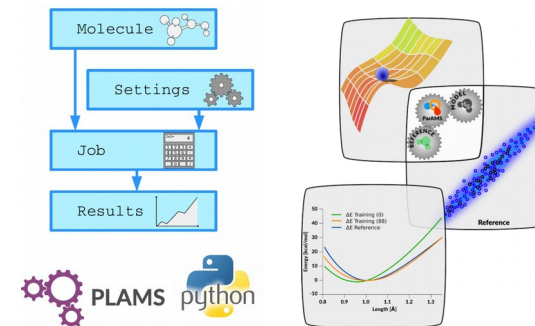
Molecular DFT

ADF

Graphical User Interface



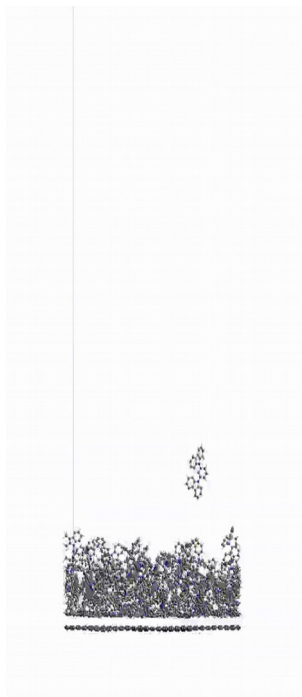
Python scripting, workflows



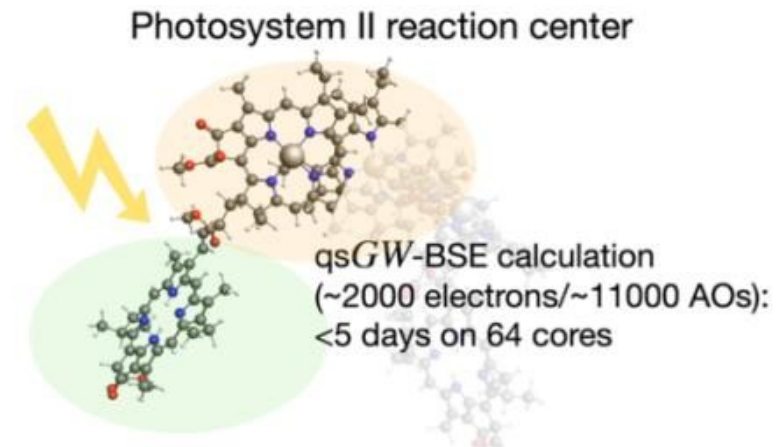
- ▶ Excitations and charge transport
- ▶ New methods/functionals in ADF
- ▶ M3GNet and ANI-2x machine learning potentials
- ▶ Conformers
- ▶ APPLE&P polarizable force field
- ▶ ACE Reaction Network
- ▶ Multiscale Modeling for Heterogeneous Catalysis
- ▶ ParAMS for ReaxFF/DFTB parametrization
- ▶ New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

Full release notes: www.scm.com/2023

- ▶ 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

▶ r²SCAN-3c

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

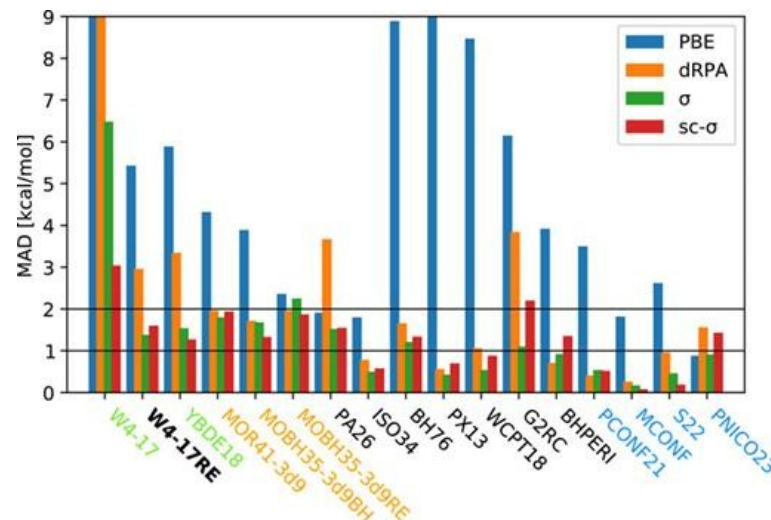


mTZ2P ✓
SR-ZORA ✓
gCP ✓
D4 ✓

T. Gasevic, J.B. Stückrath, S. Grimme, M. Bursch
Optimization of the r²SCAN-3c Composite Electronic-Structure Method for Use with Slater-Type Orbital Basis Sets
J. Phys. Chem. A 2022, 126, 23, 3826–3838

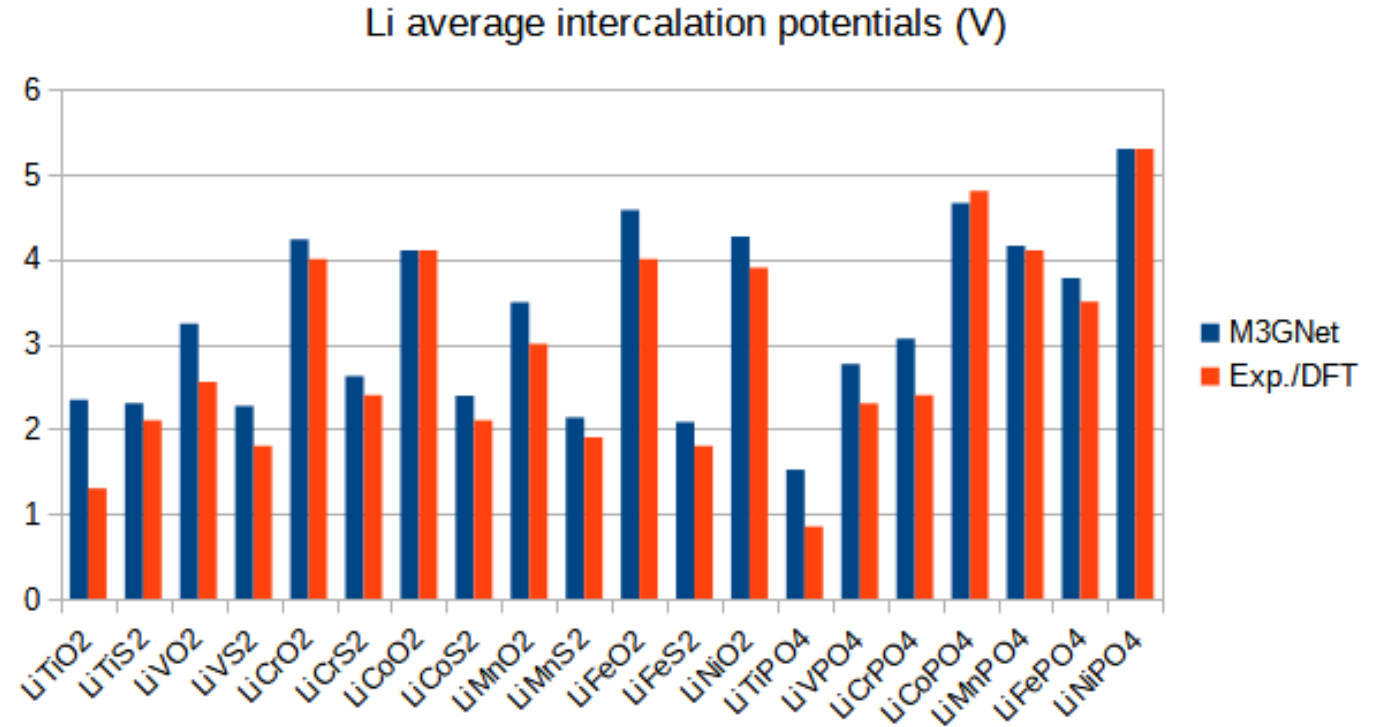
▶ Sigma functional

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



J. Erhard, S. Fauser, E. Trushin, A. Görling
Scaled σ -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>

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Article Views

4355

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21

Citations

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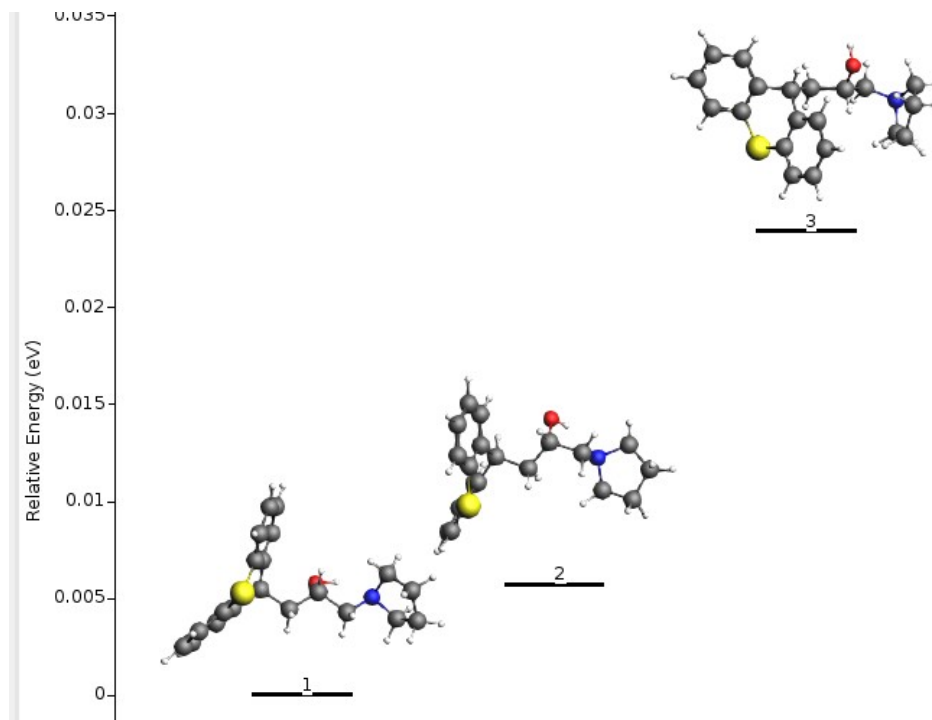
Share Add to Export



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!

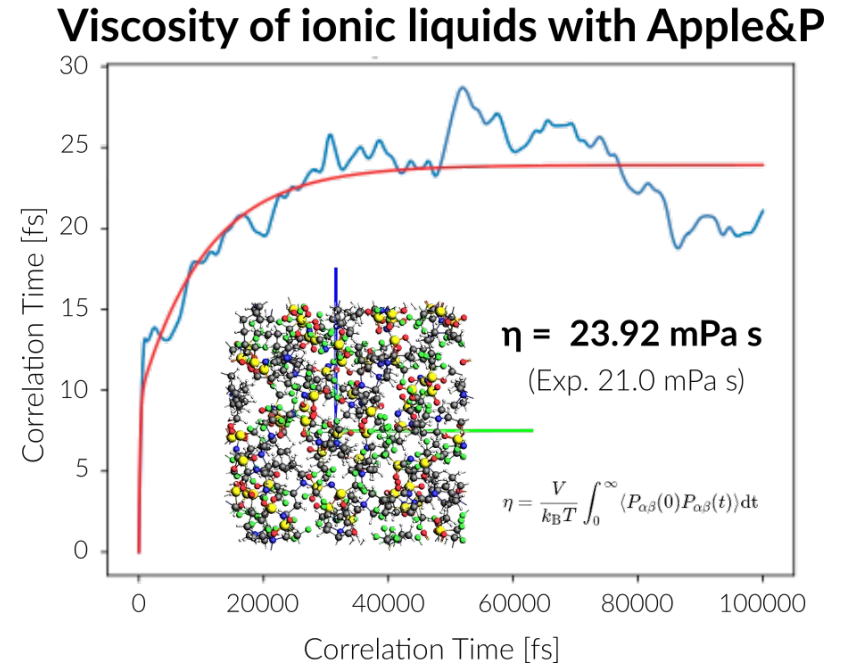
RDKit: Open-source cheminformatics; <http://www.rdkit.org>



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⁺, hydraziniums, pyrrolidiniums, ...
- ▶ Anions: hydroxide, TFSI, ...

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

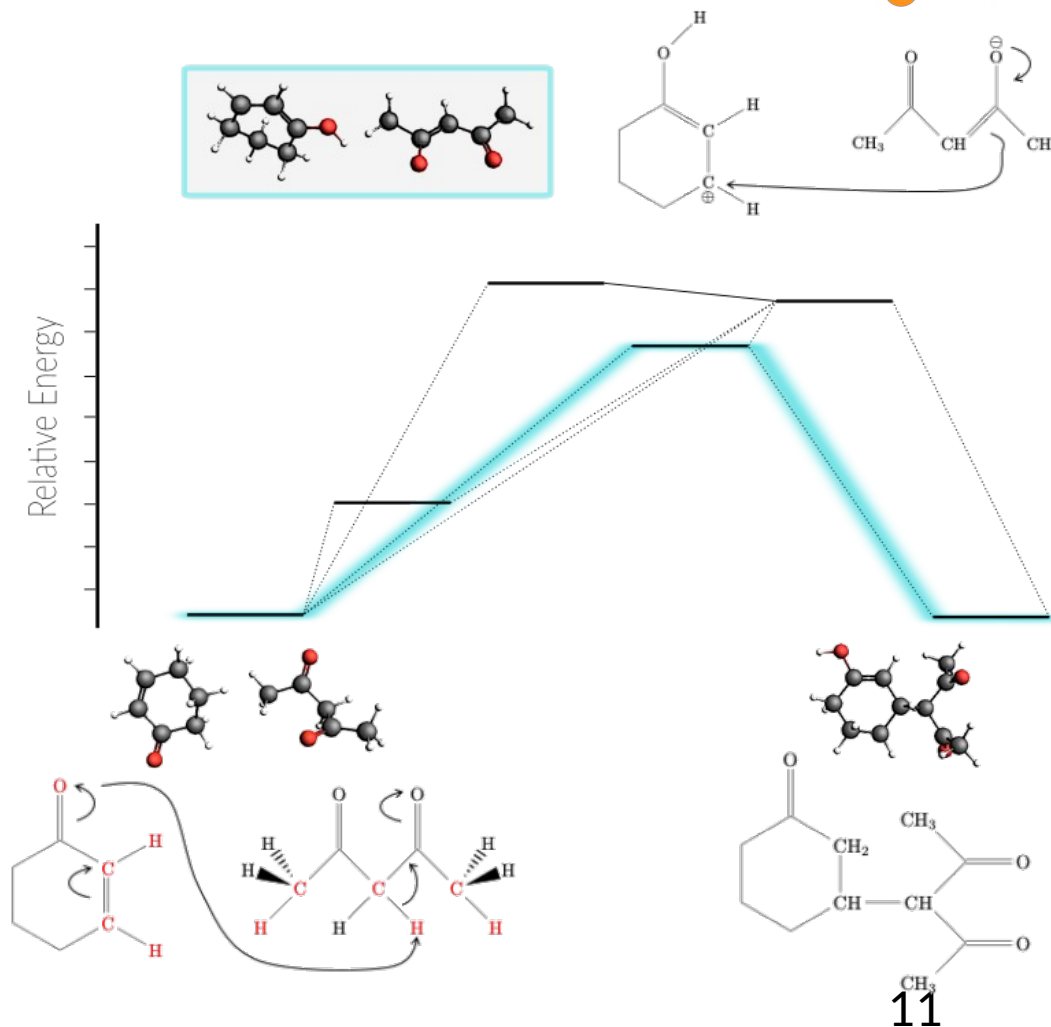


ACE Reaction Network

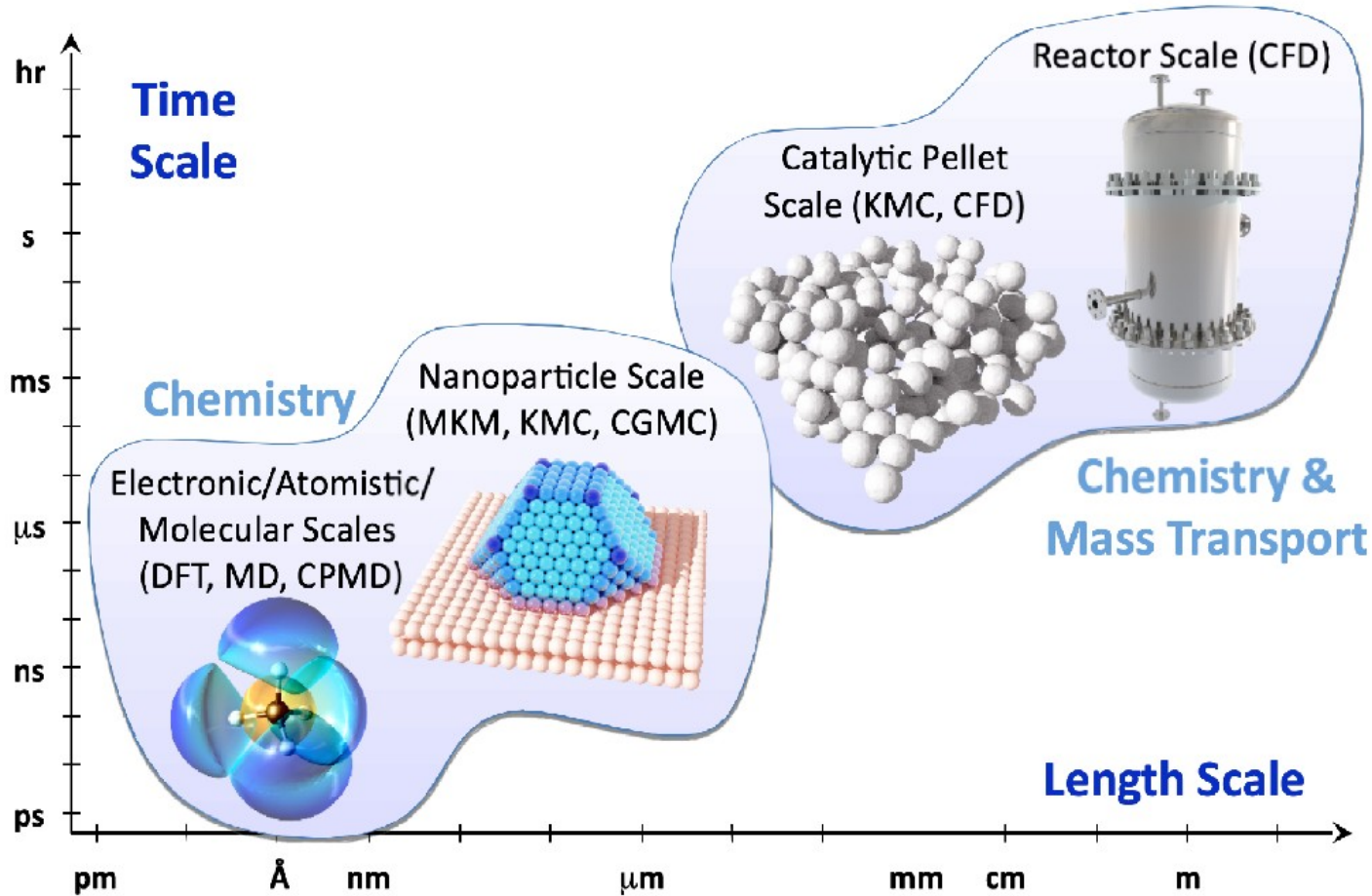


- ▶ Quickly generate and minimize reaction networks
- ▶ Find reaction mechanisms

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.

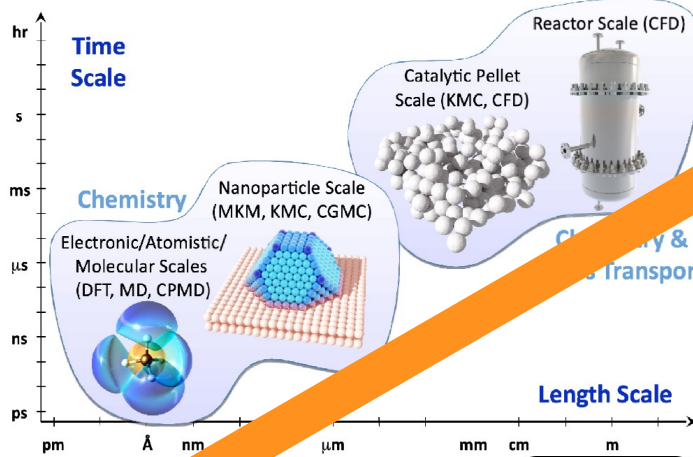


Multiscale Modeling for Heterogeneous Catalysis (1/2)

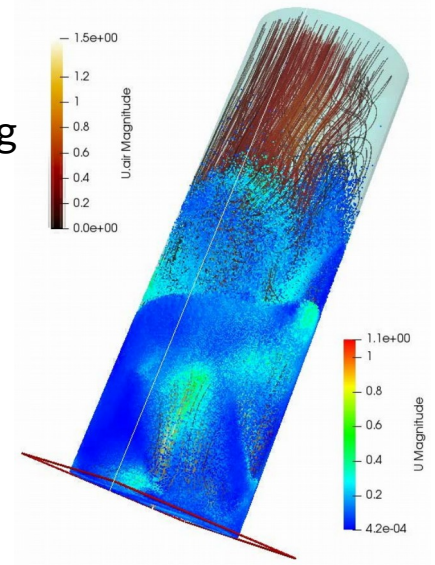


More information:
www.reaxpro.eu

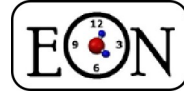
Multiscale Modeling for Heterogeneous Catalysis (2/2)



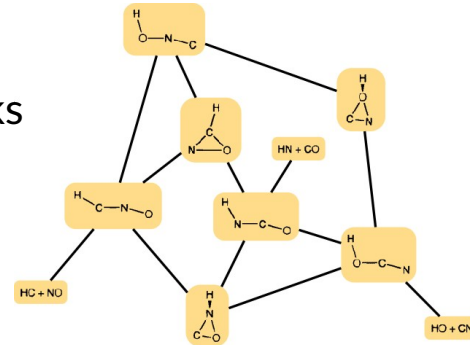
Reactor modeling



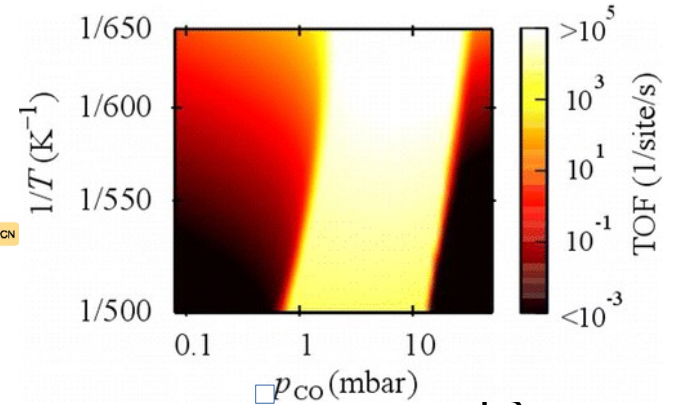
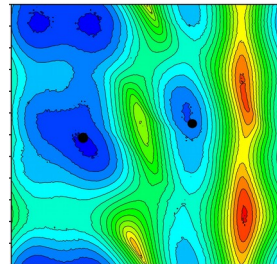
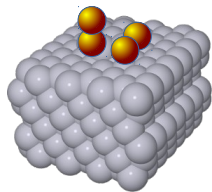
Kinetic Monte Carlo



Reaction networks



DFT & Force Fields

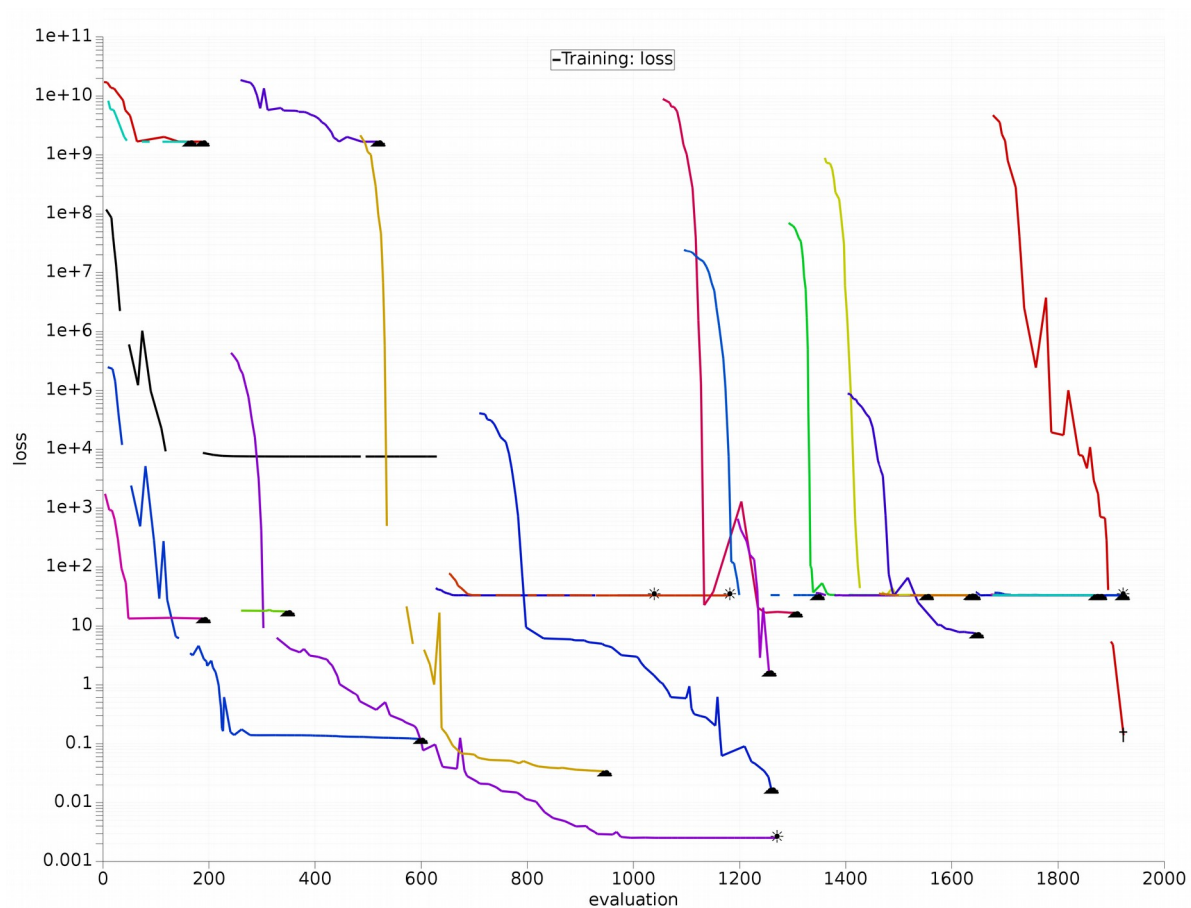


ParAMS for ReaxFF and DFTB parametrizations (1/2)

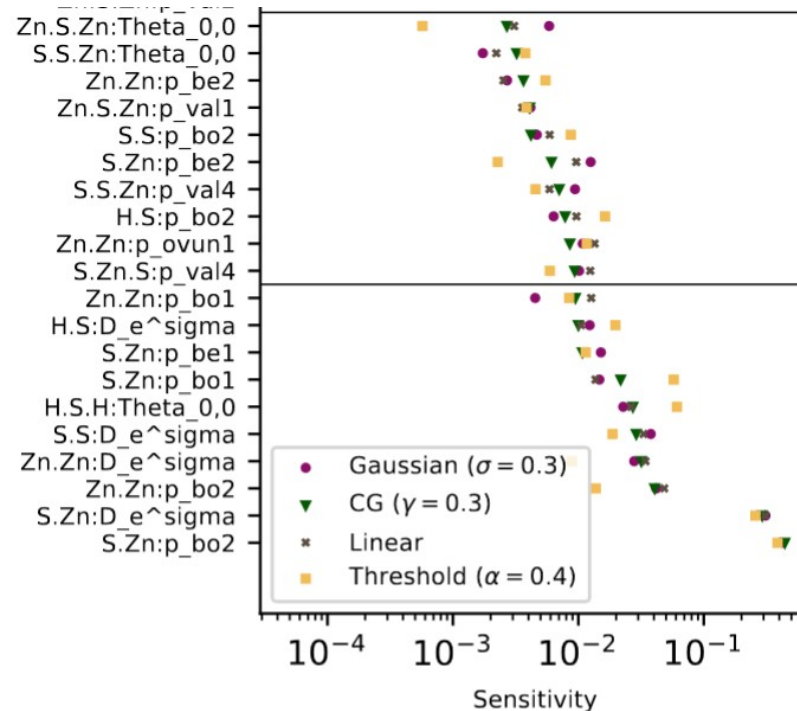
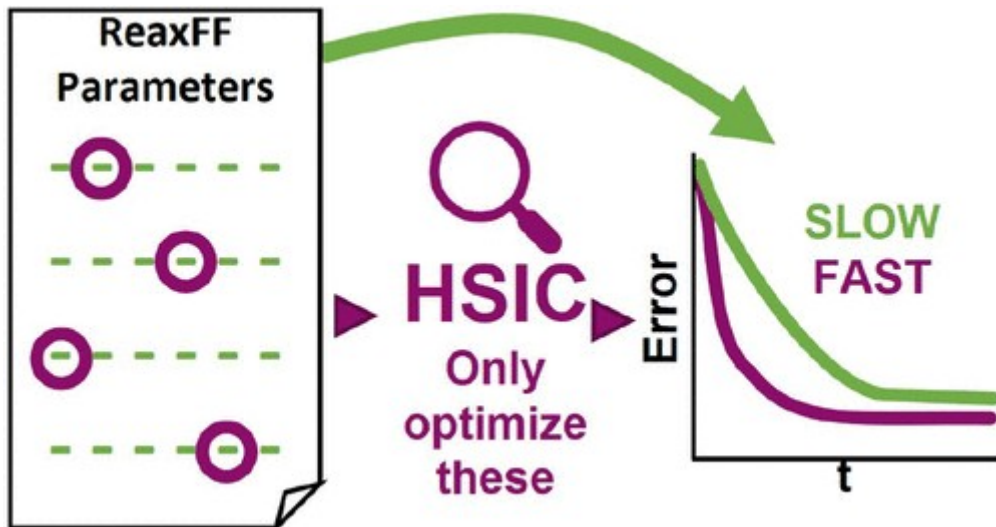


- ▶ Run and control
multiple optimizers
in parallel

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

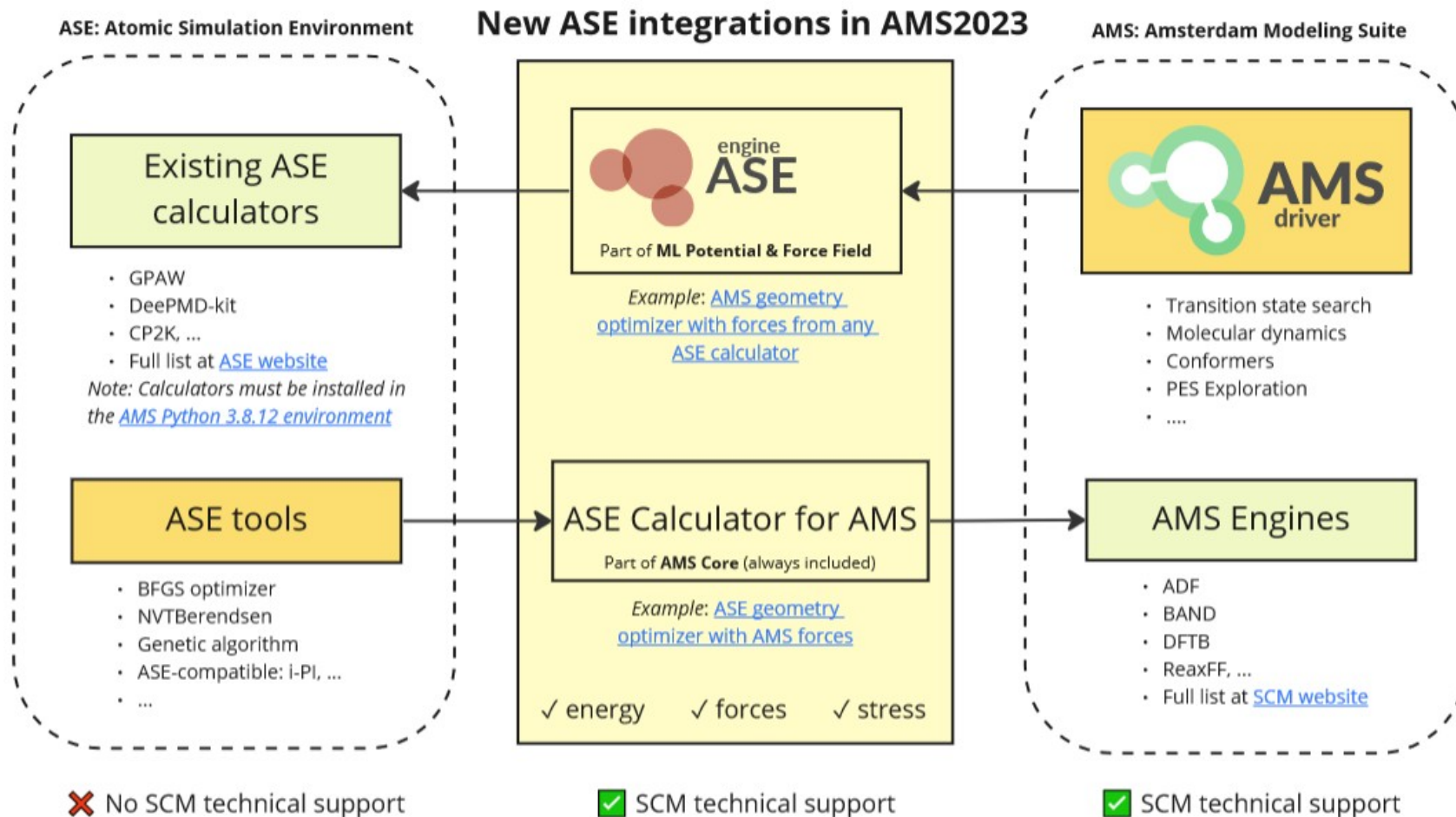


M.F. Gustavo, M. Hellström, T Verstraelen.

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

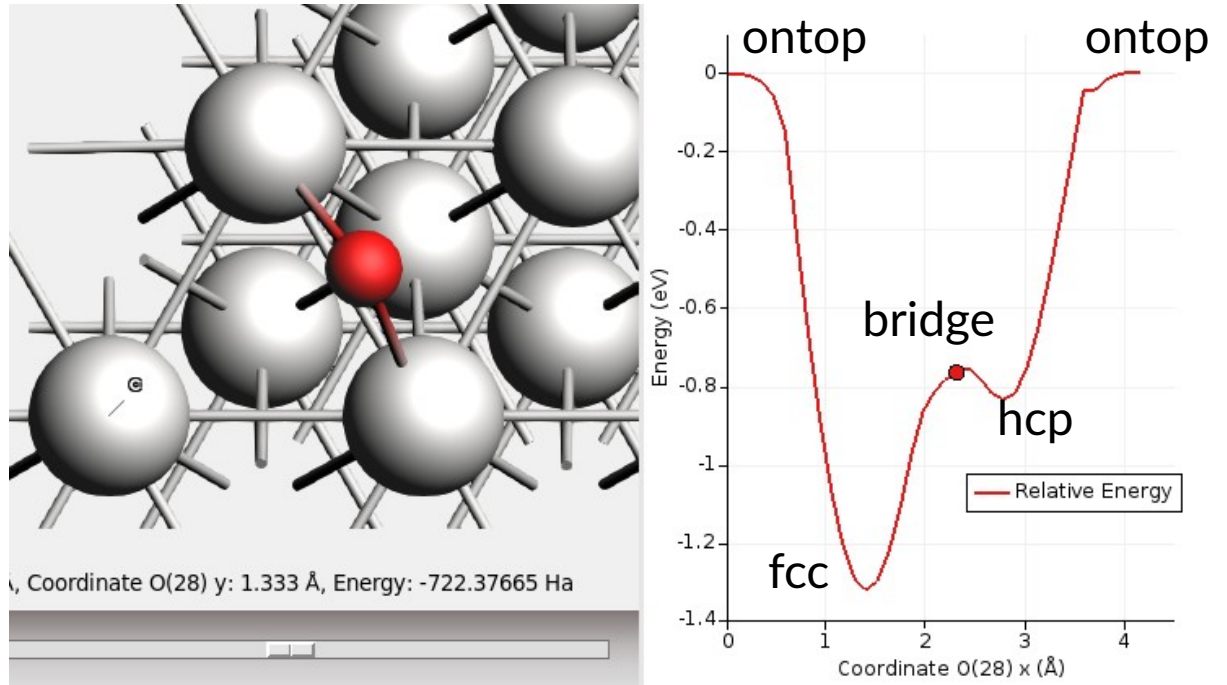
J. Chem. Theory Comput. 2023, 19, 9, 2557–2573

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!

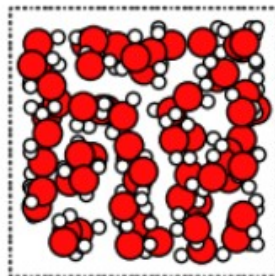
Example PES Scan: O atom diffusion on Pt(111)



- ▶ 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
- ▶ Free trial: www.scm.com/free-trial
- ▶ YouTube: webinars, tip of the week, ...:
www.youtube.com/c/AmsterdamDensityFunctional
- ▶ Questions?

