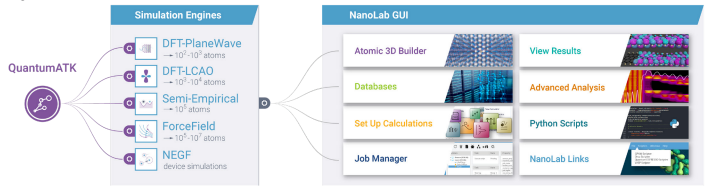


**QuantumATK-M** is a complete and fully integrated software suite for atomic-scale modeling of materials, professionally engineered using state-of-the-art scientific and software-engineering methods. It combines the power of a Python scripting engine with the ease-of-use provided by an intuitive NanoLab graphical user interface (GUI). All simulation engines share a common infrastructure for analysis, ion dynamics, and parallel performance techniques. This datasheet describes QuantumATK-M simulation engines. You can download another datasheet on the NanoLab GUI.



## Core Features of QuantumATK-M

## **NEGF Method of Nanoscale Device and Transport Simulations**

Non-Equilibrium Green's function (NEGF) module works with DFT-LCAO, Semiempirical and ForceField Modules (see next page)

- ▶ Types of Systems: Nanoelectronic Devices & Interfaces (2-Probe), Surfaces (1-Probe)
- ► Include:
  - Surface, defect, interface, grain boundary scattering, and charge transfer effects
  - External electric fields
  - An effective doping scheme to simulate doped semiconductors
  - Metallic gate electrodes and dielectric screening regions
  - Electron-phonon coupling (EPC) effects (with BTE, MD-Landauer and STD-Landauer methods)
  - Relaxation of device geometries
  - Multilevel parallelism up to 1000s of cores
  - Options to balance performance vs. memory usage

## Calculate:

- Band-alignment, Schottky barriers, and work functions for thin-film/surface heterostructures
- Electronic surface states in external electric fields
- Insight into catalytic activity of transition metal and transition metal oxide surfaces
- Thermal conductance/conductivity of materials, nanostructures, interfaces, etc.
- Electrical characteristics, such as I-V curves, on/off ratio, subthreshold slope, drain-source saturation voltage, drain-induced barrier lowering, transconductance
- Transmission spectrum, eigenvalues, eigenchannels, device density of states (DOS), etc
- Phonon-limited mobility and resistivity of materials
- Tunnel magnetoresistance ratio, spin-polarized tunneling current, and bias-dependent spin-transfer torque for material stacks that comprise the magnetic tunnel junctions
- Photocurrent and open circuit voltage) for solar-cell devices and LEDs

## Core Features of QuantumATK-M

- **DFT-LCAO:** Simulation engine for density functional theory (DFT) using pseudo-potentials and linear combinations of atomic orbitals (LCAO) basis sets.
- DFT-PlaneWave: Simulation engine for DFT using pseudo-potentials and plane-wave basis sets.
- **SemiEmpirical:** Semi-empirical simulation engine for using DFTB, extended Hückel, Slater-Koster, and other tight-binding models.
- ForceField: Simulation engine for atomic-scale simulations (e.g. molecular dynamics) using classical potentials.

**DFT-PlaneWave**→ 10²-10³ Atoms

**DFT-LCA0**  $\rightarrow 10^3-10^4 \text{ Atoms}$ 

→ 10³-10⁴ Atoms **SemiEmpirica**l

10<sup>5</sup> Atom

orceFiel

•	•	•	•

▶ Types of Systems: Molecules, (poly)crystals, polymers, nanostructures, nanoparticles, alloys	•	•	•	•
<ul> <li>Works with NEGF method for simulations of nanoelectronic devices, interfaces, surfaces, solar-cells</li> </ul>		•	•	•
▶ Use: More than 300 LDA/GGA/MetaGGA DFT functionals	•	•		
► Hybrid functionals using the ACE approximation	•			
► Van der Waals models: DFT-D2 and DFT-D3	•	•		
► Hubbard U term for LDA and GGA		•		
<ul> <li>Methods for accurate band gap calculations of semiconductors and insulators (MetaGGA, DFT-1/2, Pseudopotential Projector shift methods)</li> </ul>	•	•		
Norm-conserving Troullier-Martins PPs (FHI, SG15/PseudoDojo)	•	•		
<ul> <li>Predefined numerical atomic orbital basis sets for different accuracy levels</li> </ul>		•		
▶ Plane wave basis sets with automatic default cut-offs	•			
► Spin settings: unpolarized, polarized, noncolinear, noncollinear spin-orbit	•	•	•	
Built-in Slater-Koster models for group IV and III-V semiconductors, extended Hückel model with over 300 basis sets, and Tight-Binding models for strained systems			•	
<ul> <li>More than 300 empirical classical potentials included, Python interface for adding your own or literature potential, support for custom combination of potentials</li> </ul>				•
▶ <b>Perform:</b> Calculation of band structure, fat band structure, DOS, Fermi surface, cohesive energy, defect formation energies and transition levels, etc.		•	•	
<ul> <li>Geometry and unit cell optimization using LBFGS and FIRE methods</li> </ul>	•	•	•	•
► Molecular dynamics (MD) (NPT, NVT, NVE,) simulations for studying mechanical (e.g. creep simulation), thermal properties, and physical processes (e.g. thin film growth); part of the MD-Landauer simulations to include electron-phonon coupling (EPC) effects	•	•	•	•
<ul> <li>Nudged elastic band (NEB) for reaction path optimization (e.g. to get insight into catalytic activity of transition metal and transition metal oxide surfaces)</li> </ul>	•	•	•	•
Adaptive kinetic Monte Carlo (AKMC) for long timescale kinetics (e.g. diffusion kinetics)	•	•	•	•
Phonon modes (part of the STD-Landauer method to include EPC effects), phonon bandstructure, DOS, and thermal transport simulations	•	•	•	•
▶ Benefit from: Multilevel parallelism (threading + MPI) up to 1000s of cores. Parallelize over k-points/basis functions/NEB images/AKMC searches/bias points, etc.	•	•	•	•



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