

SILVACO

Victory Atomistic

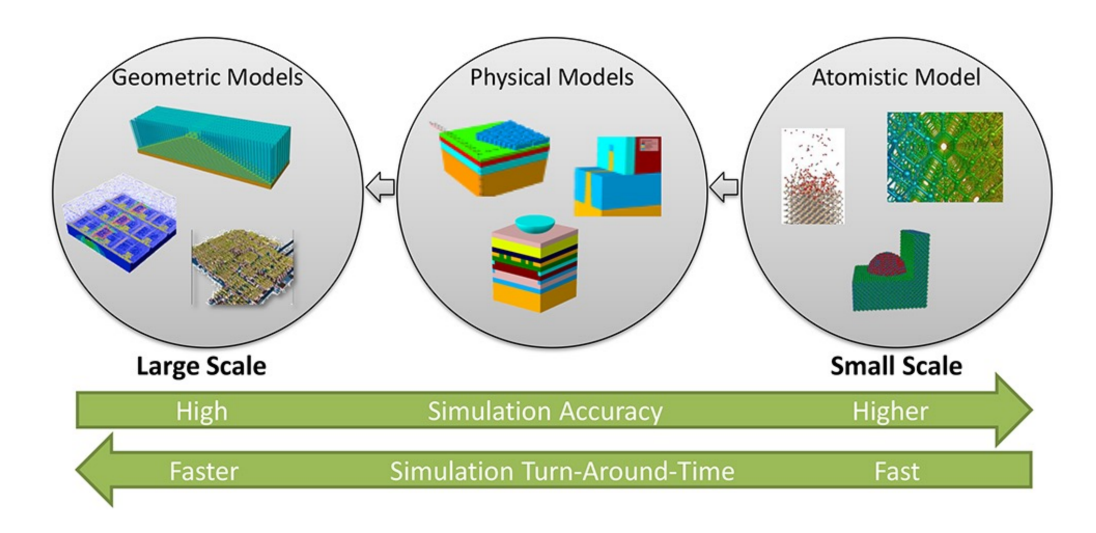
Practical Atomic Scale Simulation

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- TCAD simulation device size continuum
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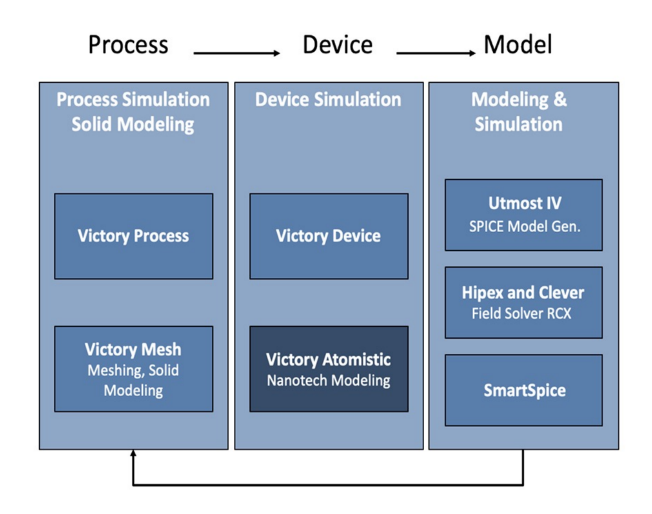
TCAD Simulation Continuum





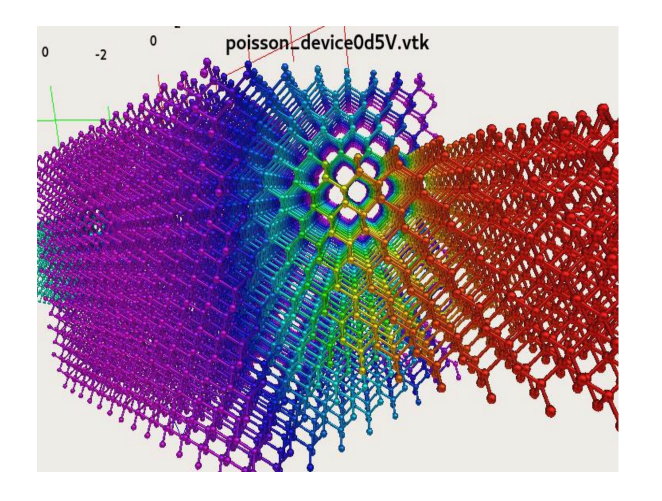
Commercial Tool Flow Integration

- The trusted legacy atomistic simulator, "NEMO" from Purdue University, is now productized and tightly integrated into Silvaco's TCAD tool flow.
- Leverages atomic level simulation accuracy, into final circuit simulations at the SPICE level



Commercial Tool Flow Integration

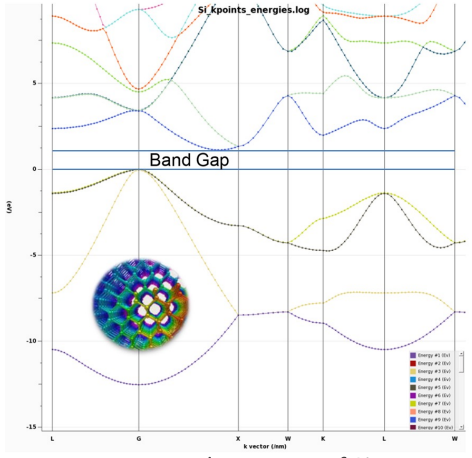
- Uses the familiar Silvaco user interface
- Newly developed in house atomistic plotting capability with other custom features
- Silvaco style manuals
- Customer support
- Examples





Full Band Structure Calculations

- Victory Atomistic uses calibrated tight binding parameters to calculate accurate band structure
- Removes the limitation of relying on the constant effective mass approximation

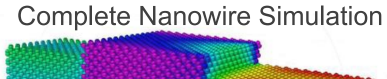


VA -Band structure of Si



Carrier Transport Device Physics

- Quantum transport:
 - QTBM ballistic
 - NEGF scattering
 - Electron phonon self energy
- Strain dependency
- Schrodinger Poisson
 - Self consistent
- Density of states
- Transmission probability
- Optics



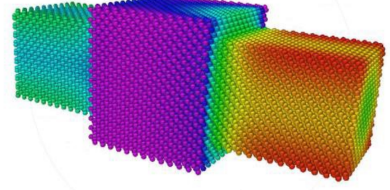
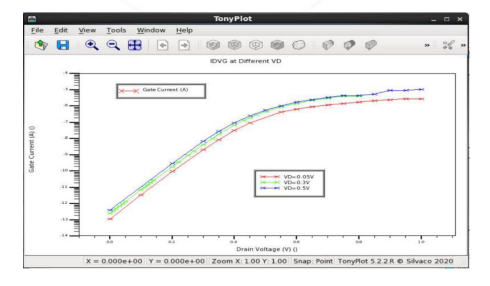


Illustration of Potential





Atomistic Simulation Speed for Everyday Use

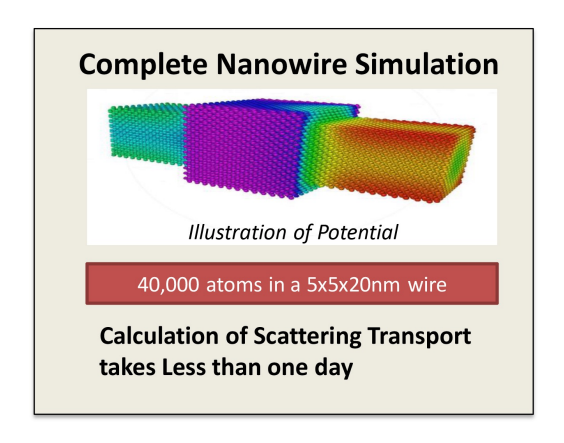
- Advanced numerical acceleration techniques like mode space and Büttiker probe
- Uses MPI and OpenMP for parallel processing

Ballistic I(V)

2x2x20nm	5 min / 30 CPUs
5x5x20nm	15 min / 120 CPUs

Scattering I(V)

2x2x20nm	120 min / 30 CPUs
5x5x20nm	480 min / 240 CPUs





Summary

- Novel approaches for greatly reduced simulation time
- Practical whole device simulations now possible
- Efficient hyper scaling and cloud based computing
- Supported commercial product with ease of use
- Link atomistic simulations to circuit SPICE models
- Simulate 2D materials, quantum dots, optical devices
- Cutting edge FinFETs, nanowires, slabs, tubes
- Knowledgeable team to assist with your project

