Molecular Dynamics Simulation Report

Optimized Lattice Parameter: 5.2756e-10 m

Melting Point: 125 K Boiling Point: 165 K

Total Runtime: 635.04 seconds

Best Verlet Algorithm: Standard Verlet (Adaptive)

Best Time Step: 5.00e-15 s

Reason: Lowest average potential energy (-1.27e-20 J) and minimal energy drift

(1.63e-22 J) during preliminary simulation.

