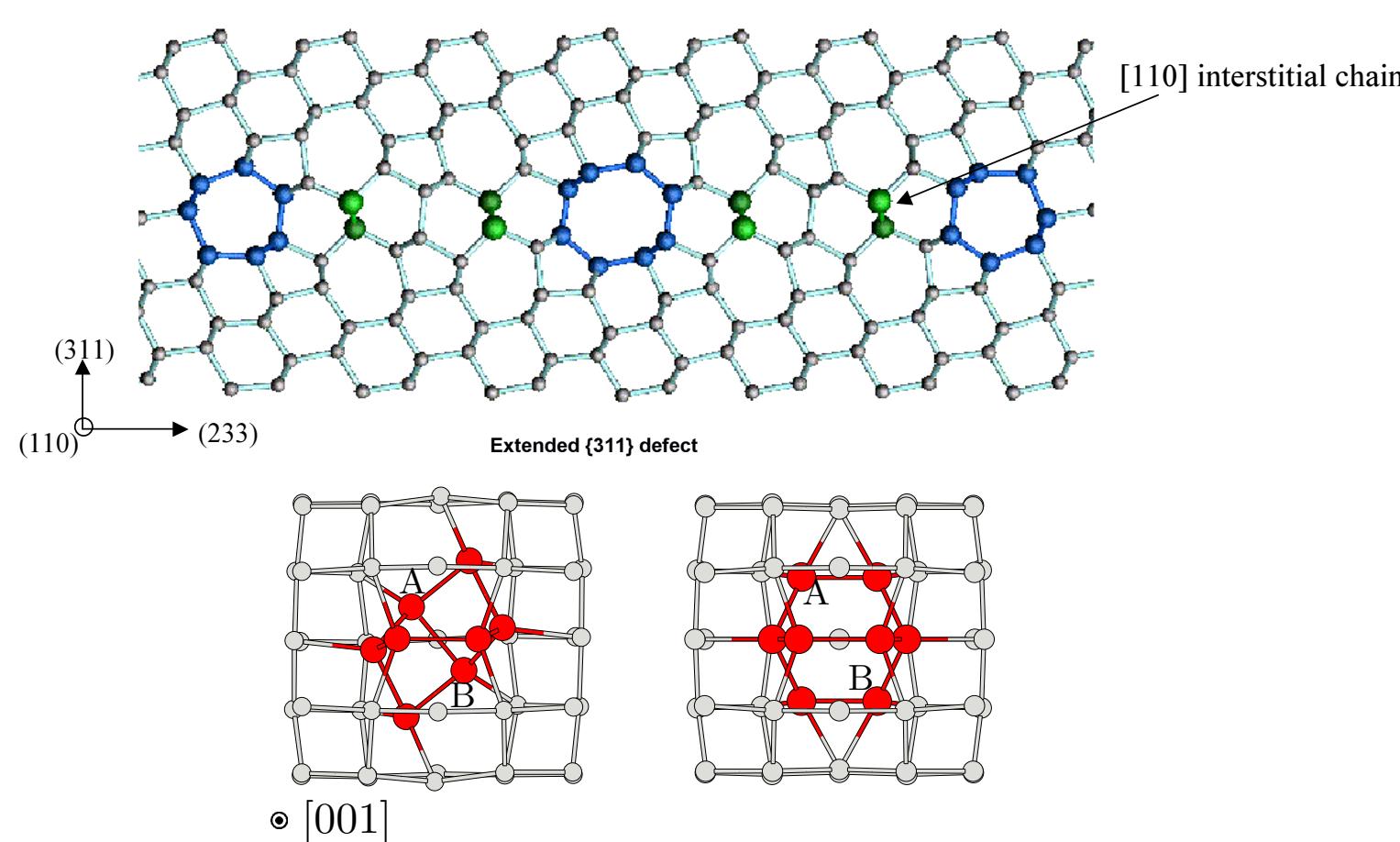


K. P. Driver, W. D. Parker, P. R. Peterson, R. G. Hennig, C. J. Umrigar, J. W. Wilkins

Interstitial Silicon Defects**From point defects to extended {311} defect**

- Ion implanted Si interstitials nucleate {311} defects limiting device fabrication and performance
- Defects identified in TEM/EELS experiment [1]
- Defects from TB-MD and DFT relaxations
- Several dozen new stable interstitial clusters [2]

**Quantum Monte Carlo Method****Trial Wave Function and Jastrow Factor**

$$\Psi_T = \underbrace{\mathcal{D}^\uparrow \mathcal{D}^\downarrow}_{\text{Slater determinant}} \times \underbrace{\mathcal{J}(r_i, r_j, r_{ij})}_{\text{Jastrow factor}}$$

$$\mathcal{J}(r_i, r_j, r_{ij}) = \prod_{\alpha i} \underbrace{\exp(A_{\alpha i})}_{\text{electron-ion correlation}} \times \prod_{ij} \underbrace{\exp(B_{ij})}_{\text{electron-electron correlation}}$$

Variance Minimization

- Optimize Jastrow by variance minimization with Levenberg-Marquardt
- More stable and requires less configurations than energy minimization

Diffusion Monte Carlo

- Stochastic method of solving many-body Schrödinger equation
- Projection of ground state

Density-Functional Calculations

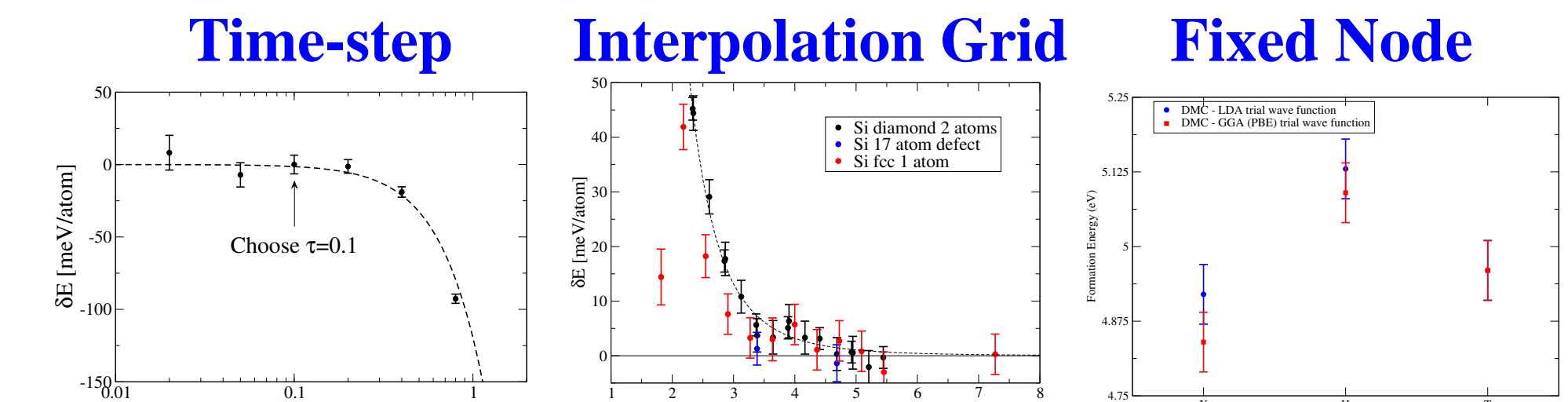
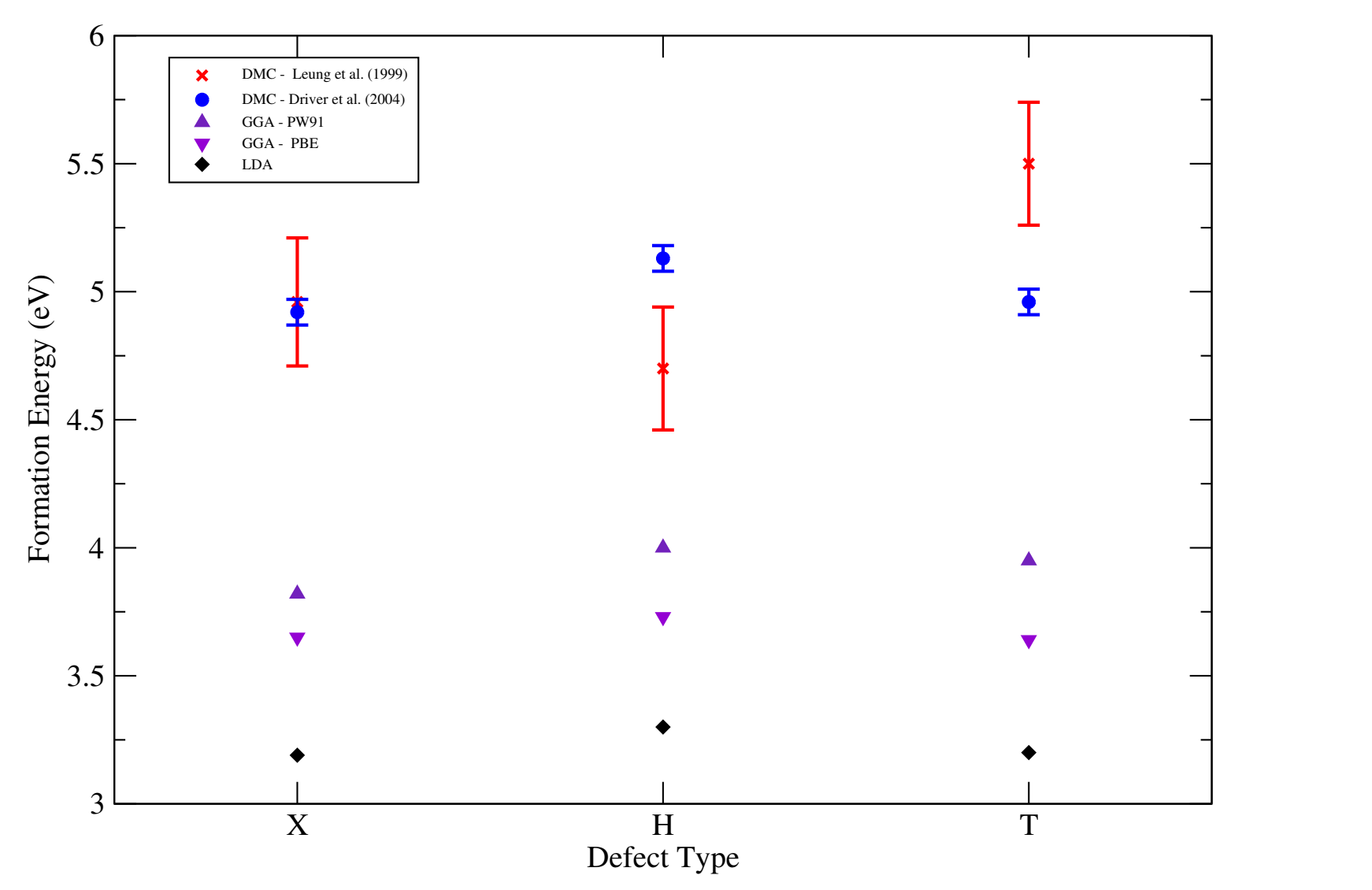
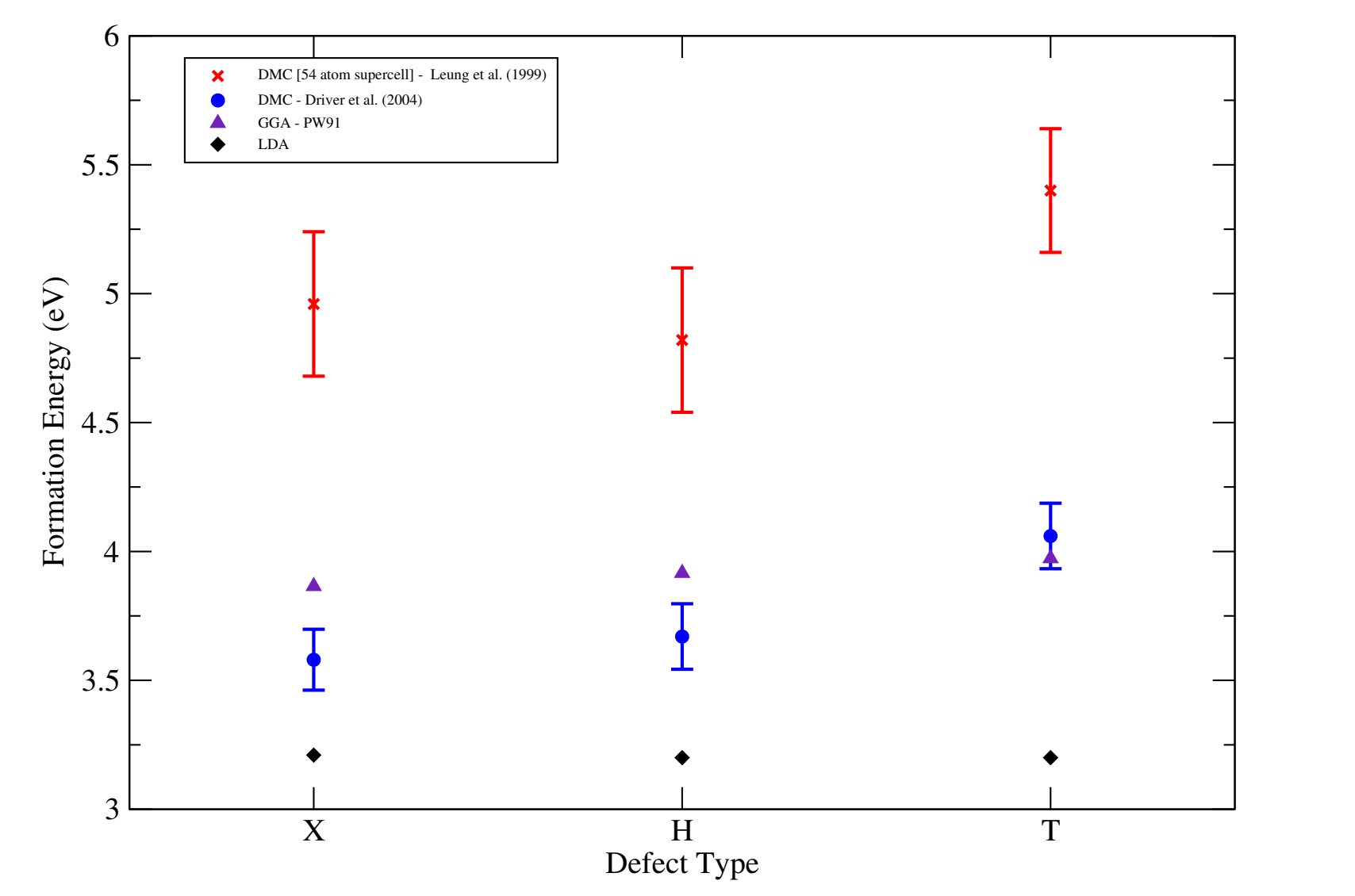
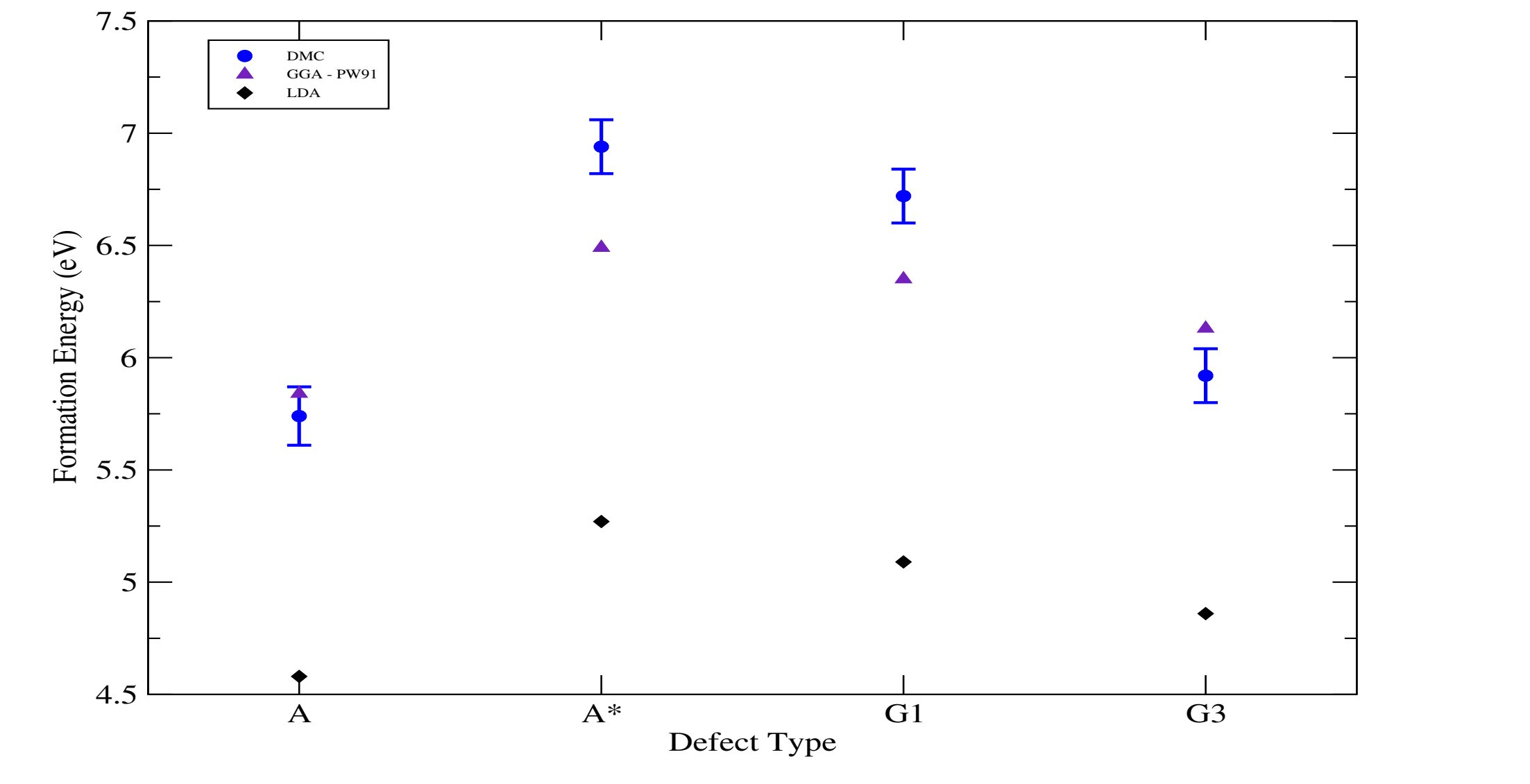
- Use CPW2000 and VASP

Approximations**Controlled**

- Statistical (increase MC steps)
- Finite-size (larger systems)
- Time-step (smaller time step)
- Population control (more walkers)
- Grid-size (decrease grid spacing)

Uncontrolled

- Fixed node
- Pseudopotential
- Pseudopotential locality

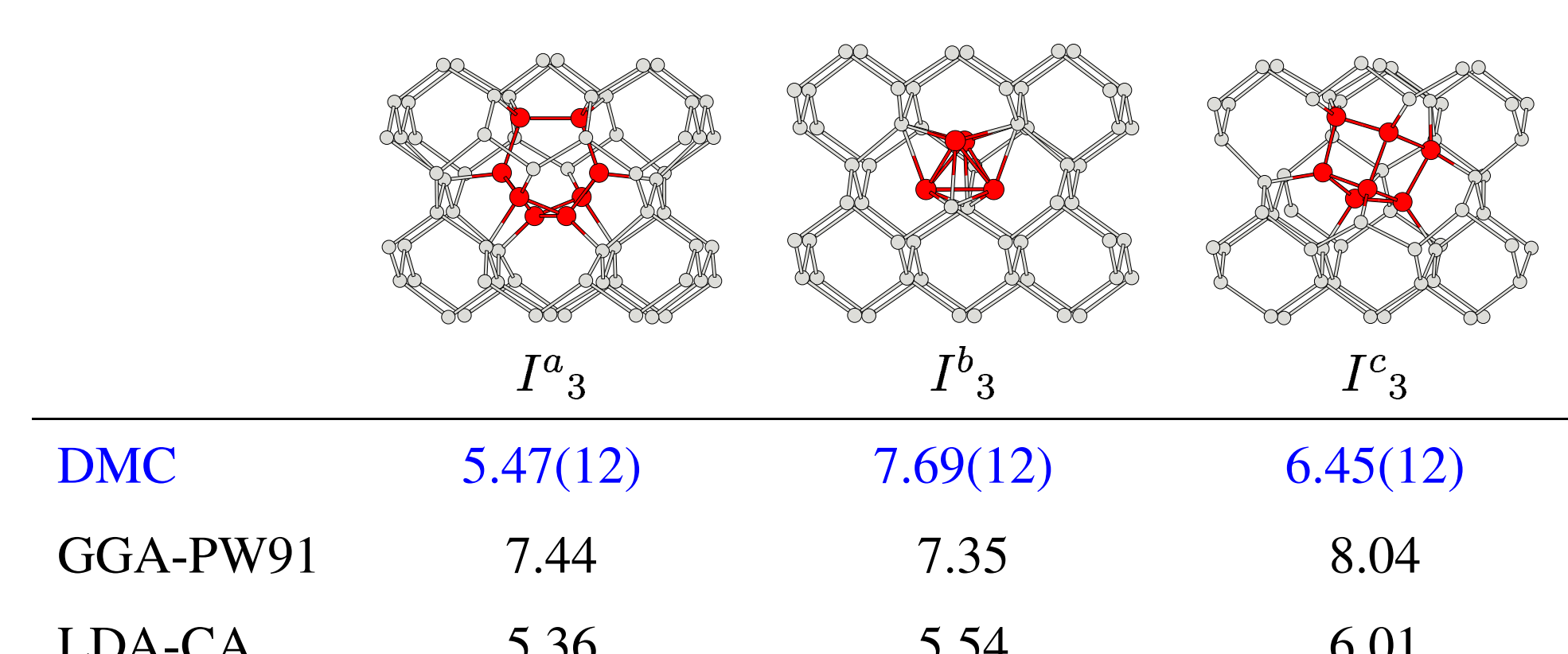
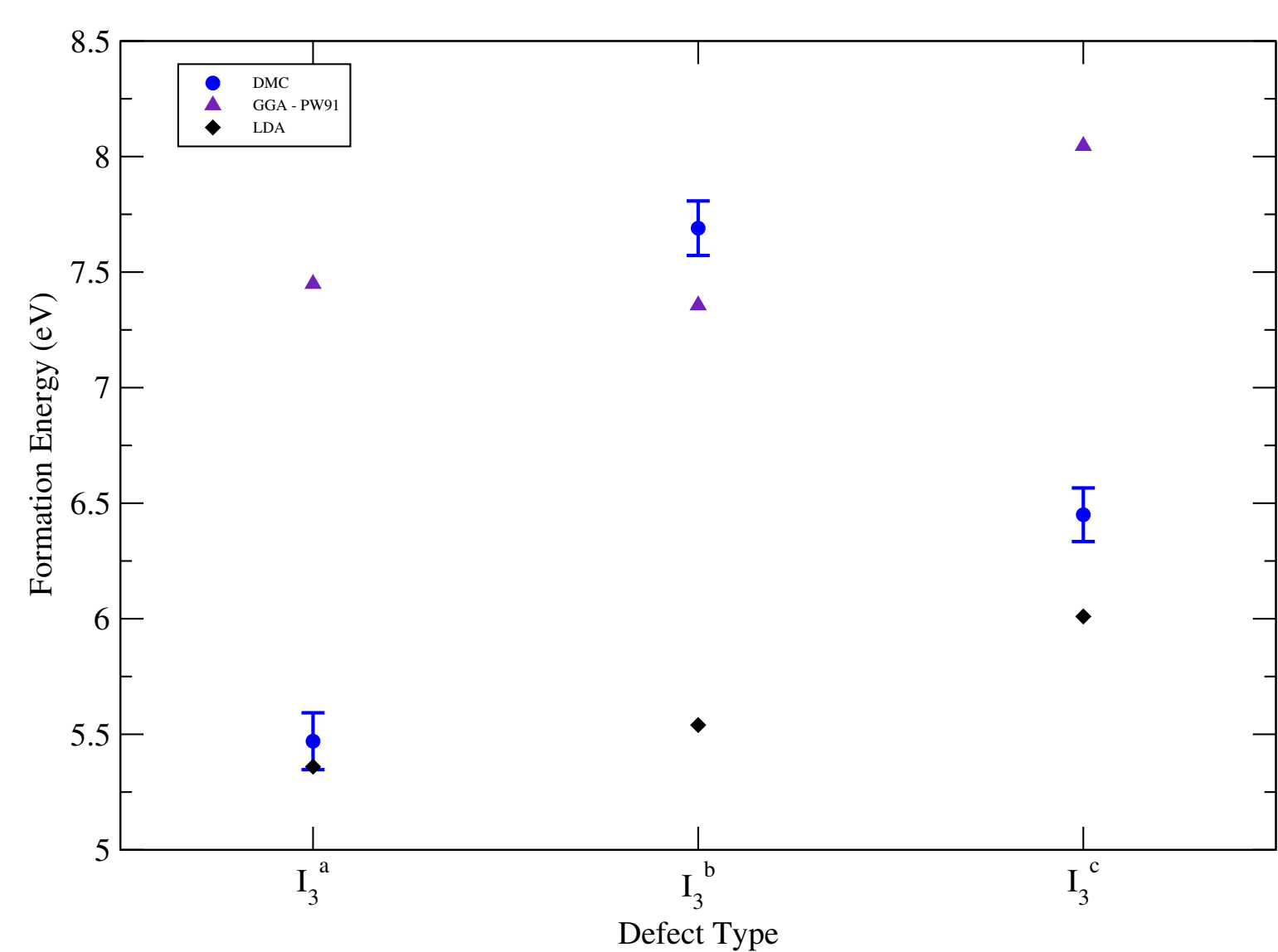
**Formation Energies of Interstitial Silicon Defects****Single interstitial in 16-atom cell****Single interstitial in 64-atom cell****Di-interstitial in 64-atom cell**

	X	H	T
DMC	4.92(5)	5.13(5)	4.96(5)
DMC ³	4.96(24)	4.70(24)	5.50(24)
GGA-PW91	3.82	4.00	3.94
GGA-PBE	3.65	3.73	3.64
LDA-CA	3.19	3.30	3.20

- 16-atom cell results are similar to earlier DMC by Leung et al.
- DFT underestimates defect energies:
 - LDA underestimates by about 1.5 eV
 - GGA (PW91 and PBE) underestimates by about 1 eV

- In 64-atom cells we find the DMC energies to be close to GGA and LDA values. This contrasts with Leung et al. observations in 54- atom cells.
- Energy versus volume curves of bulk Si suggest dependence of energy on pseudopotential.

- Di-interstitial results appear to be consistent with the single interstitial 64-atom cell results.
- GGA values lie within about 0.5 eV of DMC values.
- LDA underestimates defect energies by about 1 eV.

Tri-interstitial in 64-atom cell

	I ^a ₃	I ^b ₃	I ^c ₃
DMC	5.47(12)	7.69(12)	6.45(12)
GGA-PW91	7.44	7.35	8.04
LDA-CA	5.36	5.54	6.01

- Preliminary tri-interstitial DMC results show no systematic trend relative to LDA and GGA.

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Acknowledgements

We thank J. L. Martins for providing his density-functional code and D. Alfe, C. Filippi, R. Needs, P. Kent, and S. Goedecker for helpful discussions. Computational resources were provided by the Ohio Supercomputing Center, the Cornell Theory Center, and NERSC. The research was supported by NSF, DOE, and Sandia National Laboratory.