

SACADA Database Code: 78

Topology: [hal](#) 

of independent nodes (IN): 1
Transitivity: [1343]
Space Group: Fm-3m
Pearson: cF96
Coordination Number (CN): 4

Year: 2011

Data

Name	Pressure, GPa	Density, g/cm ³	Gap, eV	Relative energy, eV/atom	Bulk, GPa	Shear, GPa	Vickers, GPa	Refs
hal (SACADA #78)		2.285		0.654	261.7	200.6	26.4	SACADA ¹
CB								doi: 10.1134/s1063776111060173 
fcc-C12					259.39	164.24		doi: 10.1021/jp3064323 
CB								link 

Elasticity tensor (kBar)¹

5458.2520	1195.5033	1195.5033	-0.0000	0.0000	0.0000
1195.5033	5458.2520	1195.5033	-0.0000	0.0000	0.0000
1195.5033	1195.5033	5458.2520	0.0000	-0.0000	0.0000
-0.0000	-0.0000	0.0000	1925.5119	0.0000	-0.0000
0.0000	-0.0000	-0.0000	0.0000	1925.5119	0.0000
0.0000	-0.0000	-0.0000	-0.0000	0.0000	1925.5119

¹ We apply the density functional theory (DFT) approach by using the Vienna Ab Initio Simulation Package (VASP) to calculate the total energy and properties of carbon allotropes.

DFT calculations

We apply the density functional theory (DFT) approach by using the Vienna Ab Initio Simulation Package (VASP) package [6] to calculate the total energy of carbon allotropes. The Generalized Gradient Approximation [7] (GGA) for exchange-correlational functional is used everywhere. The energy cutoff set to 600 eV. Fully automatic Γ -centered k-points mesh with a reciprocal-space resolution of $2\pi \times 0.025 \text{ \AA}^{-1}$ is applied. We used tetrahedron method with Blöchl corrections to perform the k-point integration. The convergence thresholds are set at 10^{-6} eV for energy and 10^{-5} eV $\text{\AA}^{-1}$ for ionic forces. Polycrystalline elastic moduli — the bulk modulus, the shear modulus, Young’s modulus, and the Poisson’s ratio ν — have been calculated within the Voigt-Reuss-Hill [8] approximation. The Vicker’s

hardness H_v has been estimated according to Oganov's model [\[9\]](#).