First-principles investigation of the impact of high pressure on the structural, electronic and elastic properties of the type-VIII barium-doped silicon clathrate Ba$_8$Si$_{46}$.

Nassim Ahmed MAHAMMEDI $^{1,2}$, Marhou FERHAT $^{2,3}$

$^1$Laboratoire de physique des matériaux LPM, Amar Télidji University of Laghouat, BP37G, Laghouat 03000, ALGERIA.
$^2$École Normale Supérieure de Laghouat (ENSL), RN01, Laghouat 03000, ALGERIA.
$^3$Department of physics, The University of the West Indies, Mona, Kingston 07, JAMAICA.

Corresponding author
Nassim Ahmed MAHAMMEDI
n.mahammedi@lagh-univ.dz

Abstract

By means of density functional theory (DFT) calculations, we have investigated the effect of hydrostatic pressure on the structural, electronic and elastic properties of the barium-doped silicon clathrate Ba$_8$Si$_{46}$ in the type-VIII structure ($\alpha$ phase). Physical properties are calculated under different conditions of pressure (0 GPa to 45 GPa) using the GGA-PBE functional, those calculations have been performed using the Cambridge serial total energy package CASTEP code within the Materials Studio package. Electronic properties have shown that the type-VIII Ba$_8$Si$_{46}$ has metal-like properties with a fundamental bandgap of 1 eV. Under pressure the fundamental bandgap increases slightly and the positions of the valence band maximum VBM and the conduction band minimum CBM remain unchanged. We found that the compound is mechanically stable under the pressure range, but this needs to be confirmed experimentally through synthesis, a comparison with the type-I and the guest-free counterparts has exhibited promising features for the type-VIII Ba$_8$Si$_{46}$.

I. Introduction

The quest for a perfect semiconductor that combines several features is a challenging task and a revived competitive scientific topic for materials scientists and engineers. Silicon is one of the most investigated, handled, and utilized semiconductor in major high-tech devices including not only optics, electronics, microelectronics and photovoltaics, but also thermoelectrics, superconducting and future quantum computing devices. Additionally to the fact that silicon is the second abundant substance on the Earth’s crust (about 25.8% of its global mass), it has several advantages that made its industry mature and very advanced. However, exploring crystalline phases beyond the cubic diamond structure will certainly overcome its several shortages and provide more technological alternatives for expensive and rare semiconductors. Silicon clathrates are regarded as the most promising structures of silicon, they were first synthesized in 1965 by Kasper et al and Cros et al [1, 2], and their discovery released prominent
research for exploring and exploiting their many intriguing features including thermoelectrics [3, 4], electronics, photovoltaics [5], superconductors [6, 7], and even as anodes for Li-based batteries [8]. Si clathrates are cage-like framework of silicon that entraps guest atoms inside. Relatively to their constituents and their chemical formula, a very large family of clathrates can be formed [9]. Etymologically, the term ‘Clathrate’ is derived from the Greek word ‘Klethra’ which originally means ‘alder’ [10] and more precisely from the Latin word ‘Clathrus’, meaning ‘surrounded on all sides’[11], the term was used in the 1940s to describe clathrates of \( \beta \)-hydroquinone compounds by Powell, H.M. [12]. Theoretically, eight types of clathrates are described, and more clathrate-like forms can also be added [4], however, only three types are successfully synthesized so far, i.e. type-I, type-II and type-VIII. Type-I and type-VIII clathrates possess the same chemical formulas \( X_8Si_{46} \). Guest free silicon clathrates are promising materials for semiconducting and optoelectronic applications, as well as for thermoelectrics. Synthesis of guest-free silicon clathrates is still a challenging task, where attempting to grow single crystals, powder, or thin films only succeeded for the type-II configuration [13, 14]. Thin film \( Si_{136} \) was recently grown over a (111) Si substrate [15]. Norouzzadeh et al extensively investigated the type-VIII guest-free \( Si_{46} \) by first-principles calculations, their results showed very promising thermoelectric features of this material [16-18].

In this work, first-principles calculations in the frame of the DFT within the GGA approach are employed to investigate the main structural features and the impact of hydrostatic pressure on the structural, electronic and elastic properties of type-VIII \( Ba_8Si_{46} \) silicon clathrates, thus, this paper is organized as follows: the first section outlines the theoretical approach including our computational details and structural aspects of the clathrate, the second section discusses the electronic properties under pressure, and then the elastic properties under pressure are discussed in detail.

II. Theoretical aspects

II.1. Computational procedures

In this work, first-principles calculations within the density functional theory DFT [19, 20] are performed using the Cambridge Serial Total Energy Package (CASTEP) code [21]. Plane-wave based norm-conserving pseudopotentials [22] are used. The electronic configurations in these pseudopotentials are Ba: [Xe] 6s\(^2\) and Si: [Ne] 3s\(^2\)3p\(^2\). The general gradient approximation with the Perdew-Burke-Erzenhorf GGA-PBE [23] approach is selected to estimate the exchange and correlation energy XC. In order to ensure that the self-consistent convergence of the total energy reaches \( 10^{-6} eV/atom \), the kinetic plane-wave cut-off energy is set to 480 eV and the first
irreducible Brillouin zone is sampled using a 6x6x6 Monkhorst-Pack scheme [24]. Structural optimization is primordial in order to get an equilibrium structure at a minimum total energy, for that purpose, the optimizations are being performed using the Broyden-Fletcher-Goldfarb-Shanno (BFGS) algorithm [25-28] and stopped until the Hellmann-Feynman interatomic forces reach $10^{-5}$ eV/Å and the pressure reach a minimum value of 0.001 GPa for all calculations. All physical properties are calculated for equilibrium crystals optimized at minimum of the total energy.

II.2. Crystalline structure

Type-VIII silicon clathrates typically crystallize in the body centered cubic space group I4-3m No 217 [29, 30], their frameworks are initially composed from associations of polyhedral cages of silicon as a host material encapsulating alkali-metals or earth-like guest species. The strongly distorted dodecahedron cages $\text{Si}_{23}$ are the only type of polyhedron composing the type-VIII structure, the silicon atoms occupy the 2a, 8c, 12d and 24g Wyckoff positions, whereas, guests (barium for our case) occupy the 8c positions inside those cages. The host silicon atoms are bonded to each other through the strong sp$^3$ hybridization, while the barium atoms have strong van der Walls interactions with their surrounding Si clusters (due to their relatively larger size). The unit lattice of $\text{Ba}_8\text{Si}_{46}$-VIII (illustrated in figure 1), contains 54 atoms, is used as a basic model throughout this work. We have firstly performed a full structural optimization at zero pressure and minimum of total energy and interatomic forces, our results showed that the lattice constants are $a=b=c=10.395$Å (expected to be slightly overestimated by GGA). The results are compared with the literature and summarized in the table 1, unfortunately, no experimental results for type-VIII $\text{Ba}_8\text{Si}_{46}$ are reported so far. Our calculated structural parameters are in a fair agreement with the available data; however, the slight differences are due to the different DFT codes solicited for each study. In what follows, the Si atoms at 2a, 8c, 12d, and 24g are symbolized as S1, S2, S3 and S4 respectively.
III. Results and discussions

III.1. Structural properties under high pressure

In order to investigate the effect of hydrostatic pressure on the physical properties of VIII-Ba$_8$Si$_{46}$, the equilibrium lattice constants at zero pressure and at minimum total energy are calculated and used as starting structural parameters for what follows. Our results for structural parameters including optimized lattice constants and atomic positions by GGA-PBE are summarized in the table 1 and are compared with available literature. It can be noticed that intercalation of barium atoms will increase the volume of the silicon clathrate network by 2%; this is due to the relatively large size of Ba atoms. Experimentally, only ternary compounds of type-VIII clathrates have been reported, hence, experimental values for the type-VIII Sr$_8$Al$_{10}$Ga$_{6}$Si$_{30-x-y}$ (synthesized by means of arc-melting and spark plasma sintering SPS, $a=10.4444\text{Å}$ for Sr$_8$Al$_{10}$Ga$_6$Sr$_{30}$) by Shimizu et al [31], and for the type-I Ba$_8$Si$_{46}$ (synthesized by HPHT, $a=10.328\text{Å}$) by Yamanaka et al [32], are used for sake of comparison as well. Following structural optimization, we obtained lattice parameters at 0 GPa as $a=10.395\text{Å}$ for type-VIII Ba$_8$Si$_{46}$ and $a=10.121\text{Å}$ for the guest free VIII-Si$_{46}$. The lattice parameter of type-VIII Ba$_8$Si$_{46}$ obtained by GGA-PBE is closer to the experimental value of VIII-Sr$_8$Al$_{10}$Ga$_6$Si$_{30}$ [31] and to that of the type-I Ba$_8$Si$_{46}$. Lattice parameters for the guest-free VIII-Si$_{46}$ are lower than the mentioned experimental values, however, if compared to the experimental value of the guest-free type-II Si$_{136}$ synthesized by Jan Gryko et al [13] (by means of successive vacuum treatment and density separation of Na$_x$Si$_{136}$) (space group Fd-3m, $a=14.62601\text{Å}$), the values are far different.

We calculated the Gibb’s free energy at 0 GPa using the formula $G_0=E_0+PV-TS$, where $E_0$ is the total energy per unit cell and $T_S$ is the temperature. Since this DFT study is held at a ground state where $T_S=0\text{K}$, the Gibb’s free energy will be equal to the enthalpy $H_0=E_0+pV_0$. The Enthalpy variation with pressure is illustrated in figure2. Not surprisingly, it is observed that the most stable structure is under zero pressure, and then the enthalpy increases linearly with pressure above 5 GPa. Compared with the type-I phase, it was reported that the type-VIII Ba$_8$Ga$_{16}$Sn$_{30}$ clathrate has lower enthalpy and therefore it tend to be more stable under high-pressure [33]. We also estimated the formation energy of the VIII-Ba$_8$Si$_{46}$ from the following equation [16]:

$$\Delta H(M_8Si_{46})=E_{Ba8Si_{46}}-8E_M-46E_{Si} \tag{1}$$

Where: $\Delta H$ is the total formation energy, $E_M$ is the total energy of the intercalated atom calculated in its stable bulk phase, and $E_{Si}$ is the total energy of Si atom in the stable diamond
phase. A negative value is supposed to indicate that the compound is energetically favorable to form, whereas, if it is positive, that means that the material could be synthesized but require an additional amount of energy. We report a value of $\Delta H = -97.23$ eV.

The evolution of the lattice volumes and the interatomic distances Si-Si and Ba-Si under high-pressure are illustrated in figures 3 and 4 respectively. As can be noticed from figure 3, the cell volume tend to normally decrease when pressure is increased, however, the behavior is not linear, from 0 GPa to 5 GPa, the volume is drastically reduced by about 31%. Above 10 GPa, lattice volume will decrease in a quasi-linear manner with compressive pressure. At 30 GPa the overall volume is reduced by about 49%. According to Kim et al [34], a volume collapse above 20 GPa is expected for the type-VIII quaternary substituted silicon clathrate Sr$_8$Al$_8$Ga$_8$Si$_{30}$, therefore, in our case, no such change is observed. Leoni et al [35] has suggested an analytical model to analyze the phase transition from type-I ($\beta$ phase) to type-VIII ($\alpha$ phase) clathrates. High pressure behavior for type-I Ba$_8$Si$_{46}$ by Zhang et al [36] and K$_8$Si$_{46}$ by Tse et al [37] were studied by means of DFT codes. In both studies similar behaviors were observed with higher mechanical stability for the barium doped clathrate as a result of the larger Ba atoms and hence stronger interaction with the Si cage. A study by Li et al [33] on the high-pressure behavior of the type-VIII Ba$_8$Ga$_{16}$Sn$_{30}$ has also predicted that the type-VIII clathrate is more stable than its type-I allotrope, and they found that the phase transition from type-I to type-VIII ($\beta \rightarrow \alpha$) cannot occur under hydrostatic pressure regardless its magnitude. This will be understood in the discussion of mechanical properties section. Figure 3(a) exhibits the variation of the Ba-Si and Si-Si bonds under high pressure. Ba-Si and Si-Si bond lengths have different behaviors towards pressure, thus, the Ba-Si has similar tendencies towards lower values when pressure is increased. This monotonous behavior demonstrates the isotropic compression of the cages under pressure; this is certainly related to the enhanced integrity of the Si cages by the presence of Ba atoms. Nevertheless, generally, the expansion of cages volume caused by the presence of large Ba atoms will loosen the Si-Si bonding, even if the Ba-Si interaction contributes to the integrity of the structure but the Si cage will be weakened.

Si-Si bonds in contrary show different behaviors towards pressure, for example, the Si$_3$(12d)-Si$_4$(24g) and Si$_4$(24g)-Si$_4$(24g) distances decrease slowly comparing to the Si$_1$(2a)-Si$_2$(8c) and Si$_2$(8c)-Si$_4$(24g) bonds. It can be observed that the three Si-Si lengths converge towards similar values at 30 GPa unlike the Si$_1$(2a)-Si$_2$(8c) bond which seems to decrease more rapidly.

We have also examined the evolution of angles in the silicon frameworks under high-pressure and their evolution is plotted in the figure 3(b). It can be seen that the Si(2-1-2) angle in the
pentagonal rings are constant under pressure. Other angles such as the Si(4-2-4) in the six-fold rings and the Si(4-3-4) in the seven-fold rings are not affected by pressure. The Si(4-3-4) in the six-fold rings increases with pressure, whereas, the Si(2-4-3) and Si(4-4-3) in the six- and seven-fold rings respectively decrease rapidly with pressure. At 20 GPa, it can be noticed that the Si(1-2-4) and the Si(2-4-4) in the pentagonal rings became equal and cross at this pressure. Anisotropic behavior of angles is a normal characteristic of cage-like structures and is a consequence of the isotropic behavior of the overall lattice volume.

![Graph (a)](image)

![Graph (b)](image)

**Figure 2.** Evolution of (a) the Total enthalpy and (b) the equilibrium volume of VIII-Ba$_8$Si$_{46}$ under pressure
Figure 3. High-pressure behavior of: (a) Ba-Si and Si-Si distances, and (b) the Si-Si-Si angles for the type-VIII Ba$_8$Si$_{46}$
### Table 1

| Atom   | Site:          | X        | Y        | Z        |
|--------|----------------|----------|----------|----------|
| Si1    | (x, y, z)      | 0        | 0        | 0        |
| Si2    | (x, y, z)      | 0.36713  | 0.36713  | 0.36713  |
| Si3    | (x, y, z)      | 0.25     | 0.5      | 0        |
| Si4    | (x, y, z)      | 0.41891  | 0.41891  | 0.41891  |
| Ba     | (x, y, z)      | 0.19183  | 0.19183  | 0.19183  |

**Lattice constant** $a_0$ (Å)

| Atom   | $a_0$ (Å) | $b_0$ (Å) | $c_0$ (Å) |
|--------|-----------|-----------|-----------|
| Si1    | 10.395    | 10.385    | 10.385    |
| Si2    |           |           |           |
| Si3    |           |           |           |
| Si4    |           |           |           |
| Ba     |           |           |           |
| Formation energy (kJ.mol$^{-1}$) | -97.23$^a$ | -97.17$^b$ | -94.05$^c$ |
| Density (Kg/m$^3$) | 3.82 x 10$^3$ | 3.7 x 10$^3$ |
| Fundamental bandgap (eV) | 1.07$^a$ | 1$^b$ | 1$^c$ |

$a$: our work GGA-PBE  
b: Ref [16] GGA-PBE  
c: Ref [30] GGA-PBE

### III.2. Electronic properties

#### III.2.a. At zero pressure

We have computed by means of GGA-PBE the band structures and the total and partial densities of states DOS of Ba$_8$Si$_{46}$-VIII under zero and high-pressure conditions. According to the space group (I4-3m, N°217), the band structures are calculated along the following path at the first Brillouin zone: $\Gamma$(0,0,0), H(1/2,2/1,-1/2), N(1/2,0,0), P(1/4,1/4,1/4). Figure 4(a) illustrates the band structure and the figure 4(b) the partial DOS at 0GPa. The Ba$_8$Si$_{46}$-VIII clathrate has a similar allure of band structure with the guest-free counterpart, they differ by the position of the Fermi level, generally, inorganic group-IV clathrates obey the rigid band model [38], according to which they will qualitatively have similar allures of band structures regardless the nature and the size of the intercalated species as long as the host framework atoms remain the same. This is due to the fact that guest atoms will contribute with their valence electrons to the framework’s antibonding states, and hence, the Fermi level will be shifted towards the conduction band, creating mostly *n*-type heavily or lightly doped semiconducting materials (upon the concentration of guest atoms). Valance and conduction bands are not overlapped at the Fermi level (set to 0 eV as energy reference). This result is coherent with previous studies by Norouzzadeh et al [16, 30], who has shown that Ba$_8$Si$_{46}$-VIII is a metallic-like compound with a fundamental bandgap of about 1eV (GGA-PBE) appearing between $\Gamma$ and N high-symmetry points. We found that the fundamental bandgap between the two same points is about 1.07 eV.
The total and partial densities of states DOS are illustrated in the figure 4(b). Structural stability of a compound is directly associated with its bonding strength. It can be noticed that near the valence band, the Si atoms contribute with their p-states, while the contribution from Si-s states increases when going toward low energies. Ba atoms contribute through their s and p states mainly to the middle and lower parts of the valence band, and they are less present in the conduction band.

![Figure 4. Electronic properties of VIII-Ba₈Si₄₆: (a) the band structure, and (b) the total and partial DOS.](image)

**III.2.b Under pressure**

In order to study the impact of hydrostatic pressure on the electronic properties, we compare the band structures and densities of states (DOS) of type-VIII Ba₈Si₄₆ under 0, 15, and 30 GPa. In Fig. 5 are displayed the GGA-PBE band structures of the Ba₈Si₄₆-VIII along the Γ-H-N-P high-symmetry directions for (a) 0 GPa (b) 15 GPa and (c) 30 GPa. The Fermi level is set to 0 eV as an energy reference. It can be noticed that within the fundamental bandgap, the top of the valence band (VBM) is located at the N k-point and remains at its position under hydrostatic pressure, whereas, the conduction band minimum is located at Γ-H for 0 GPa and 15 GPa and moves to Γ under 30 GPa. At Γ, higher alignment with other valleys is observed at the CB edge,
and a smaller effective mass at this point is estimated which can lead to higher electronic mobility [16], and therefore to better semiconducting and thermoelectric properties. The fundamental band gap slightly increases under pressure; it goes from 1.07 eV at 0 GPa to 1.12 eV at 15 GPa, and to 1.17 eV under 30 GPa. This can be explained in terms of partial density of states. In Figure 6, contributions from Ba and Si states become stronger under high pressure, Due to the strong hybridization of the Si-s and Ba-p states at Fermi level at higher pressures; at 30 GPa we observe an additional peak of the Ba-p states at 15 eV energy level. Also, the contribution from silicon states near and beyond the Fermi level increases with increasing pressure.

![Figure 5](image.png)

**Figure 5.** High-pressure effect on the band structure of the type-VIII Ba$_8$Si$_{46}$ at: (a) 0 GPa, (b) 15 GPa, and (c) 30 GPa.

![Figure 6](image.png)

**Figure 6.** Influence of high-pressure on the total density of states DOS of the type-VIII Ba$_8$Si$_{46}$. 
III.3. Elastic constants and modulus

III.3.a. At zero pressure

The type-VIII Ba₈Si₄₆ clathrate crystallize into a body centered cubic structure; therefore, only three independent elastic constants ($C_{11}$, $C_{12}$ and $C_{44}$) are sufficient to describe the elastic behavior of the material. Defining these elastic constants and investigating their behavior under high-pressure is primordial to predict the mechanical stability of the investigated material under such conditions during synthesis. For sake of comparison, elastic constants and modulus at zero pressure were also computed for the type-VIII guest-free Si₄₆, and for the type-I Si₄₆ and Ba₈Si₄₆ by means of the same GGA-PBE approach and settings. Results are compared with the available literature. The total energy of a given symmetrical crystal at zero pressure and at its equilibrium volume state $V₀$ is given by:

$$ \Delta E(\varepsilon) = \frac{1}{2} V₀ C_{ij} \varepsilon^2 + 0[\varepsilon^4] \quad (2) $$

Where: $\Delta E(\varepsilon) = E(\varepsilon) - E₀$ is the total energy difference. $C_{ij}$ are the elastic constants and $\varepsilon$ are the strain tensor elements. According to the Hooke’s law of general elasticity $\sigma_{ij} = C_{ijkl} \varepsilon_{kl}$, elastic constants can be calculated through the second derivatives of the total energy. Moreover, the three elastic constants are generally derived from a set of three equations. The resolution of the first two equations, which implicate the bulk modulus $B$ and the shear constant $C'$, will give $C_{11}$ and $C_{12}$ as described hereafter.

$$ B = \frac{1}{3} (C_{11} + 2C_{12}) \quad (3) $$

$$ C' = \frac{1}{2} (C_{11} - C_{12}) \quad (4) $$

The bulk modulus $B$ is obtained by applying a hydrostatic strain tensor and then by fitting the Munaghan equation of state. The shear constant $C'$ is obtained through applying a volume-conserving tetragonal stress tensor:

$$ r = \begin{pmatrix} \varepsilon & 0 & 0 \\ 0 & \varepsilon & 0 \\ 0 & 0 & \frac{1}{(1+\varepsilon)^2} -1 \end{pmatrix} \quad (5) $$

Application of such tensor will change the total energy as follows:

$$ \Delta E(\varepsilon) = 6V₀ C' \varepsilon^2 + 0[\varepsilon^4] \quad (6) $$
The third equation gives the $C_{44}$ constant through applying an orthorhombic stress tensor

\[
\varepsilon = \begin{pmatrix}
0 & \varepsilon & 0 \\
\varepsilon & 0 & 0 \\
0 & 0 & \frac{\varepsilon^2}{1-\varepsilon^2}
\end{pmatrix}
\]  

(7)

This total energy is obtained as:

\[
\Delta E(\varepsilon) = 2V_0 C_{44} \varepsilon^2 + 0[\varepsilon^4]
\]  

(8)

Then the shear modulus $G$ is given for a cubic system as:

\[
G = \frac{1}{5}(C_{11} - C_{12} + 3C_{44})
\]  

(9)

The Young’s modulus $E$ is given as:

\[
E = \frac{9BG}{3B + G}
\]  

(10)

Additionally, the Young’s modulus $E$ along $<100>$ and $<111>$ directions are respectively described as:

\[
E_{(100)} = \frac{(C_{11} + 2C_{12})(C_{11} - C_{12})}{C_{11} + C_{12}}
\]  

(11)

\[
E_{(111)} = \frac{1}{E} \frac{C_{11} - C_{12} - 2C_{44}}{3C_{44}(C_{11} + C_{12})}
\]  

(12)

The average speed of sound $v$ in a material is given as [39]:

\[
v = \left[\frac{1}{3} \left( \frac{2}{v_{\text{tran}}^3} + \frac{1}{v_{\text{long}}^3} \right) \right]^{-1/3}
\]  

(13)

Where $v_{\text{long}}$ and $v_{\text{tran}}$ are the longitudinal and transversal sound velocities, which are expressed as:

\[
v_{\text{long}} = \sqrt{\frac{3B+4G}{3\rho}} \quad \text{and} \quad v_{\text{tran}} = \sqrt{\frac{G}{\rho}}
\]  

(14)

Where: $\rho$ represents the mass density (Kg/m$^3$).

First, we examine the elastic properties of the Ba$_8$Si$_{46}$-VIII at zero pressure. The numerical values of the above-described elastic constants are estimated using the CASTEP code by applying a small set of deformations to the crystal and taking the second derivatives of the total energy as expressed in equation (1). Our results of elastic constants $C_{ij}$, the bulk modulus $B$, the shear modulus $G$, the Young’s modulus $E$, and the average velocity of sound $v$, calculated by GGA-PBE for Ba$_8$Si$_{46}$-VIII at 0 GPa are listed in table 2, and compared with reported data for type-I and type-VIII Si clathrates. Our results are in a fair agreement with the literature. We
notice from table 2 that VIII-Ba8Si46 having a bulk modulus of $B=75.804$ GPa, lower than that of the type-I Ba8Si46 ($B=76$ GPa), and that of the guest-free Si46 type-I ($B=76.92$ GPa) and type-VIII ($B=78.87$ GPa). The bulk modulus $B$, which presents a measure of resistance of the material against external deformation, provides much information about both the hardness and the bonding strength of the studied material. Our results confirm that the type-VIII Ba8Si46 is less resistant to external deformation than its counterparts. However, resistance to plastic deformation is also proportional to the elastic shear modulus $G$. Additionally, the shear modulus $G$, the Young’s modulus $E$, and the average velocity of sound $v$ are all larger for the type-VIII Ba8Si46 than those for the type-I Ba8Si46 but reduced when compared with guest-free counterparts Si46.

Ductility and fragility play a critical role during the synthesis of materials. In general, a ductile material is able to deform a lot before it is broken unlike a fragile material which breaks without or with less deformation, we can therefore predict if the material is fragile or ductile upon the well-known Pugh ratio $G/B$ [40], if this ratio is equal or lower than 1.74 the material is least malleable (fragile), otherwise it is more malleable (ductile). In our case we predict that $B/G = 1.681$, slightly lower than the given value, this indicates that the material should be fragile at zero pressure. Elastic constants $C_{11}$, $C_{12}$ and $C_{44}$ are slightly reduced for the type-VIII Ba8Si46. The first elastic constant $C_{11}$, which quantitatively represents the uniaxial deformation along the $<001>$ direction has the highest value, indicating the incompressible character of the compound under the $<001>$ uniaxial stress and is relatively smaller in type-I than in the type-VIII Si46. One can approximately define the melting temperature $T_m$ of a cubic crystal from $C_{11}$ through the equation suggested by Fine et al [41] where $T_m=553+(5.91 \ C_{11}) \pm 300$ K ($T_m$ in K, $C_{11}$ in GPa), we therefore predict for the type-VIII Ba8Si46 that $T_m=1274.634$ K $\pm 300$ K. Considering the Born’s stability criteria for cubic crystals [42] $C_{11}>0$, $C_{11}-C_{12}>0$, $C_{11}+2C_{12}>0$ and $C_{44}>0$, the two types are mechanically stable. The Si46-VIII seems to be harder than its type-I counterpart. Our results are in good agreement with those reported by Norouzzadeh et al [17], who predicted that Si46-VIII is a strongly-isotropic material, and stable against shear and high-pressure, similarly to the type-I Si46. However, empty clathrates, in general, tend to have lower bulk modulus than the guest-containing clathrates, the guest-host hybridization usually plays an important role that can lead to hard ‘diamond-like’ materials [10]. But in contrary, our result shows that intercalation of 8 Ba atoms into the Si46 framework reduces the bulk modulus by about 17 %, this is in fact due to the expansion of the Si cages caused by the presence of big Ba atoms and therefore the weakening effect of this doping over the overall hardness of the material. Furthermore, it was demonstrated through many experimental and theoretical reports that the presence of guest atoms might decrease the stability of clathrates through these host-guest interactions that weakens the
Si-Si bonding on one hand [43], and in the other hand, the electron exchange enforces the integrity of the clathrates frameworks [37]. The main challenge for obtaining a guest-free clathrate Si$_{46}$, whether in type-I or VIII phases, is the fact that removing the guest atoms from the cages will induce a collapse of the system and hence a phase transformation towards more stable phases such as diamond structure Si$_2$ [10, 37]. Nevertheless, guest-free type-II Si$_{136}$ clathrate, was the only guest-free silicon clathrate to be successfully synthesized as crystalline [13], or thin film forms [15].

**III.3.b. Under high-pressure**

In order to probe the impact of hydrostatic pressure on the mechanical behavior of VIII-Ba$_8$Si$_{46}$, we have calculated (using the same GGA-PBE functional with the same computational procedures as precedent calculations) the elastic constants and modulus for optimized structures under high-pressure (from 0 GPa to 45 GPa). Figure 7(a) illustrates the evolution of calculated elastic constants $C_{ij}$ under high-pressure, unfortunately, comparison with other experimental or theoretical data is not possible since such studies are not yet reported. It can be noticed that $C_{11}$ and $C_{12}$ have similar monotonous pressure dependence behaviors, however their magnitudes are quite different, whereas, $C_{44}$ increases slowly and is less affected by pressure. The $C_{44}$ variation with pressure shows that a maximum value of 55.72 GPa is reached at 10 GPa, and then it tends to stabilize under higher values of pressure. Considering the previously pointed stability criterion for cubic crystals, the $C_{ij}$ constants remain all positive under pressure. However, the previous criterion does not provide alone a precise prediction of the high-pressure stability of the material in question, therefore, the following conditions will be applied and the results are checked afterwards, and if fulfilled, the cubic material might be stable under such isotropic-pressure conditions [44]:

\[
\tilde{C}_{44} > 0, \quad \tilde{C}_{11} > |\tilde{C}_{12}|, \quad \tilde{C}_{11} + 2\tilde{C}_{12} > 0
\]

(15)

Where $\tilde{C}_{11}$, $\tilde{C}_{12}$ and $\tilde{C}_{44}$ present the elastic constants under pressure, and are given as:

\[
\tilde{C}_{11} = C_{11} - P, \quad \tilde{C}_{12} = C_{12} + P, \quad \tilde{C}_{44} = C_{44} - P
\]

(16)

$P$ is the pressure in GPa.

Figure 7(b) illustrates the evolution of $\tilde{C}_{11}$, $\tilde{C}_{12}$ and $\tilde{C}_{44}$ along with the third criterion $\tilde{C}_{11} - \tilde{C}_{12}$ that is equal to: $C_{11} - C_{12} - 2P$ and which presents the precedent stability criterion under high-pressure. When pressure increases, our calculations show that the values of $C_{11} - C_{12} - 2P$ will not remain positive under all the pressure range, and the conditions are not fulfilled though,
therefore, we predict that type-VIII $\text{Ba}_8\text{Si}_{46}$ will not be stable for pressures above 24 GPa. Henceforward, the compound VIII-$\text{Ba}_8\text{Si}_{46}$ is expected to remain mechanically stable during and after synthesis below 24 GPa, however, we have to clarify that this criterion is not sufficient alone to predict any phase transition that can occur below 24 GPa, which is needed to be proven through experiments.

**Figure 7.** High-pressure evolution of (a) elastic constants $C_{11}$, $C_{12}$ and $C_{44}$, and (b) the stability criterion $C_{11}-C_{12}-2P$ for the type-VIII $\text{Ba}_8\text{Si}_{46}$
Figure 8. High-pressure evolution of major elastic modulus including the bulk modulus $B$, the shear modulus $G$ and the Young’s modulus $E$, for the type-VIII Ba$_8$Si$_{46}$

The major elastic modulus of VIII-Ba$_8$Si$_{46}$ under high pressure are computed and displayed in figure 8. It can be observed that the bulk modulus $B$ increments monotonously with pressure in a linear manner, this was expected and is due to the fact that $B$ is proportional to the applied pressure and inversely proportional to the volume change through the common physical equation of the bulk modulus ($B = \Delta P/\Delta V$). Young’s modulus $E$ which represents the resistance of the material to uniaxial strains and provides the strength degree is found to rapidly increase from 0 to 10 GPa and then starts to slowly increase beyond that pressure, this indicated the incremented stiffness of the material towards pressure. The shear modulus $G$ which describes the resistance towards shearing forces is less affected by pressure, but it tends to slightly increase and stabilizes after 10 GPa at about 56.50 GPa. The values of $E$ and $G$ both have similar behaviors towards pressure, even slowly, this means that, the capacity of the resist stiffness and shear deformation of type-VIII Ba$_8$Si$_{46}$ are increased with the increase of pressure respectively. However, the type-VIII Ba$_8$Si$_{46}$ is globally more resistive to volume compressions than to shear deformation since in all cases $B > G$ for the pressure range.
IV. Conclusion

In this paper, first-principles calculations in the frame of the Density Functional theory DFT were solicited to investigate the high-pressure behavior of the structural, electronic and mechanical properties of the type-VIII Ba$_8$Si$_{46}$ silicon clathrate. The structural evolution shows a stable behavior with pressure with no significant indices of possible phase transition or volume collapse. Electronic properties are highly affected by the presence of pressure, the metallic character of the compound is not changed throughout the pressure range, however, the interaction between barium atoms and their surrounding Si environment becomes stronger with pressure, and as a consequence, the Fermi level decreases. Elastic properties are also affected by pressure, the major mechanical properties were computed for relaxed and stressed materials, and our results showed that the type-VIII Ba$_8$Si$_{46}$ is relatively hard and resistant against deformation.

| Clathrates | Type-I | Type-VIII |
|------------|--------|-----------|
|            | Si$_{46}$ | Ba$_8$Si$_{46}$ | Si$_{46}$ | Ba$_8$Si$_{46}$ |
| $C_{11}$ (GPa) | 134.89$^a$ 140$^b$ | 135$^f$ | 146.17$^a$ 149.28$^b$ 139.3$^c$ | 122.104$^a$ 125.46$^d$ |
| $C_{12}$ (GPa) | 47.93$^a$ 47$^b$ | 46.3$^f$ | 45.22$^a$ 49.22$^b$ 46.3$^c$ | 42.154$^a$ 43.26$^d$ |
| $C_{44}$ (GPa) | 47.8$^a$ 47$^b$ | 24.7$^f$ | 45.88$^a$ 46.83$^b$ 42.3$^c$ | 45.825$^a$ 46.5$^d$ |
| $B$ (GPa) | 76.92$^a$ 78$^b$ 78.5$^c$ 87$^d$ | 76$^f$ 93$^g$ | 78.87$^a$ 82.57$^b$ 80.3$^c$ | 75.804$^a$ 76.66$^d$ |
| $G$ (GPa) | 46.07$^a$ 46$^b$ | 32.6$^f$ | 47.72$^a$ 48.08$^b$ | 45.085$^a$ 45.17$^d$ |
| $E$ (GPa) | 115.21$^a$ 115.33$^b$ | 85.6$^f$ | 119.133$^a$ 120.80$^b$ | 108.436$^a$ 109.57$^d$ |
| $v$ (m/s) | 4402$^a$ 4931$^b$ 5345$^b$ | 3390$^f$ | 5261.7$^a$ 5196$^b$ 5053$^b$ | 4695$^a$ 4509$^d$ |

Table 2: The calculated elastic constants $C_{ij}$, Bulk modulus $B$, Shear modulus $G$, Young’s modulus $E$ and the mean sound velocity $v$ for the type-VIII silicon clathrate Ba$_8$Si$_{46}$ by GGA-PBE at 0 GPa in comparison with Si$_{46}$ and the literature.

a: This work GGA-PBE (CASTEP)
b: data from ref [17] GGA-PBE (ABINIT)
c: data from ref [45] GGA-PBE (QUANTUM-ESPRESSO)
d: data from ref [30]
e: data from ref [46] LDA (SIESTA)
f: data from ref [47] Hartree–Fock HF (CRYSTAL 95)
g: data from ref [36]
h: data from ref [34]
i: data from ref [48]
and shear forces and might collapse if pressure exceeds 24 GPa, however, our predictions are to be confirmed through experiments.

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