



Cite this: *RSC Adv.*, 2019, 9, 27208

Structure, stability, absorption spectra and aromaticity of the singly and doubly silicon doped aluminum clusters $Al_nSi_m^{0/+}$ with $n = 3-16$ and $m = 1, 2^\dagger$

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Structures of the binary Al_nSi_m clusters in both neutral and cationic states were investigated using DFT and TD-DFT (B3LYP/6-311+G(d)) and (U)CCSD(T)/cc-pvTZ calculations. Silicon-doped aluminum clusters are characterized by low spin ground states. For small sizes, the Si dopant prefers to be located at vertices having many edges. For larger sizes, the Si atom prefers to be endohedrally doped inside an Al_n cage. Relative stability, adiabatic ionization energy and dissociation energies of each cluster size were evaluated. A characteristic of most Si doped Al clusters is the energetic degeneracy of two lowest-lying isomers. Calculated results confirm the high stability of the sizes Al_4Si_2 , $Al_{12}Si$ and $Al_{11}Si_2^+$ as "magic" clusters, that exhibit 20 or 40 shell electrons and are thermodynamically more stable as compared to their neighbors. Electronic absorption spectra of isoelectronic magic clusters Al_{13}^- , $Al_{12}Si$, and $Al_{11}Si_2^+$ that have two pronounced bands corresponding to blue and violet lights, have been rationalized by using the electron shell model. The magnetically included ring current density (MICD) analyses suggest that they are also aromatic structures as a result of the "magic" 40 shell electrons.

Received 27th May 2019
Accepted 21st August 2019

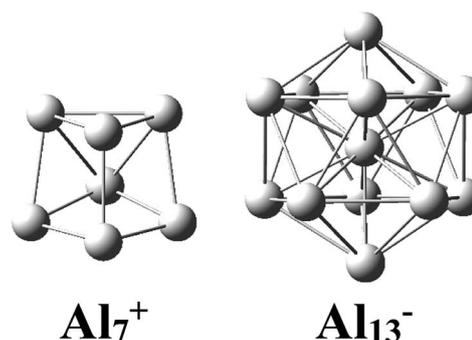
DOI: 10.1039/c9ra04004h

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1. Introduction

There has been considerable interest in the aluminum clusters as witnessed by a large number of experimental and theoretical studies reported on them in the past decades, in part due not only to their appealing physical and chemical properties, but also to their promising abilities for new technological applications.¹⁻¹⁶ Both the pure cationic Al_7^+ and anionic Al_{13}^- clusters (Scheme 1) were revealed as "magic clusters" with an enhanced thermochemical stability as each possesses a closed electron shell structure including 20 and 40 valence electrons, respectively. The neutral Al_{13} , as formed from removal of one electron from the anion Al_{13}^- , was shown to be a superhalogen having a very high electron affinity exceeding those of halogen atoms.^{8,17}

Along with pure Al clusters, several doped Al clusters have also been investigated. For alkali and alkali-earth dopants, while Rao *et al.*¹⁸ presented a theoretical study on their geometries and energetics including adsorption energies, ionization potentials, and electron affinities of Al_nLi and Al_nK , Sun and coworkers¹⁹ examined Al_nBe using the high accuracy method CCSD(T). Wang and coworkers^{20,21} investigated the Al Zintl anion moieties within the mixed sodium as well as magnesium aluminum clusters $Na_mAl_n^{-/0}$ and $Mg_mAl_n^-$ using a combination of anion photoelectron spectroscopy and density functional theory (DFT) calculations. Studies on transition metal (TM) doped Al clusters provide us with a considerable amount



Scheme 1 Structural motifs of the pure magic clusters Al_7^+ and Al_{13}^- .

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[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/c9ra04004h



of results. For copper derivatives Al_nCu with $n = 11\text{--}14$, unlike the alkali atom-doped counterparts in which the alkali dopant prefers an exohedral position, the copper atom tends to reside inside an Al cage.²² For platinum doped aluminum clusters, Zhang *et al.* reported a joint photoelectron spectroscopic and theoretical study of the PtAl^- and PtAl_2^- anions.²³ An experimental study on the argon physisorption ability of TM indicated that Al_nTM^+ clusters attach one argon atom up to a critical cluster size n_{crit} , with $n_{\text{crit}} = 16$ and $n_{\text{crit}} = 19\text{--}21$ for $\text{TM} = \text{V}, \text{Cr}$, and $\text{TM} = \text{Ti}$, respectively, and undergo a geometrical transition in going from exohedrally to endohedrally doped clusters in which the TM atom is located inside an Al cage.²⁴ Experimental results on Al_nTi were confirmed by subsequent theoretical DFT computations.²⁵

More recently, the influence of spin on the properties of TM doped aluminum clusters in a series of low spin clusters was found in which a prominent odd–even oscillation in all calculated properties supports the presence of jellium shell structure. The electron jellium model considers a cluster of atoms as a superatom where the cores of constituent atoms form a constant positive background, the “jellium density”, and only valence electrons are treated explicitly. Under the model, the shell molecular orbitals of the clusters are denoted as 1S, 1P, 1D, 2S, 2P, 1F, ... (usually the notation 1s, 1p, 1d, 2s, 2p, 1f, ... have also been used), rather than as the atomic orbitals 1s, 2s, 2p, 3s, 3p, 3d, ... of hydrogen-like atoms. High spin ground state isomers show more smooth trends similar to the bulk materials.²⁶ Geometrical and electronic structures as well as physical and chemical properties of Al clusters doped by various common nonmetal dopants have also abundantly been investigated.^{27–45}

Interaction of hydrogen molecules with Al_nRh_2^+ with $n = 10\text{--}13$ clusters was recently studied by mass spectrometry and infrared multiple photon dissociation (IRMPD) spectroscopy.⁴⁶ A comparison of the IRMPD spectra with predictions obtained using DFT calculations showed that for $n = 10$ and 11, a single H_2 molecule dissociates upon binding to the cluster, whereas for $n = 12$ and 13, it adsorbs molecularly. Upon adsorption of a second H_2 molecule to the cation $\text{Al}_{12}\text{Rh}_2^+$, both hydrogen molecules dissociate. These results point out a peculiar catalytic effect of doped Al clusters.

Silicon, being contiguous to Al in the periodic table and having one valence electron more than Al, has been and still is massively used in the semiconductors and optoelectronic industries. A mixture of aluminum–silicon atoms gives rise to some quite interesting compounds as previous studies indicated some remarkable properties of Al–Si nanomaterials. For instance, interaction between a Si nanowire and one Al-atom was found to increase the electrical conductivity as compared to pristine Si-nanowire.⁴⁷ Al atoms form an ordered array of magic clusters on the surfaces of $\text{Si}(111)$ ⁴⁸ and a persistent formation of Al–Si nanowires was reported.⁴⁹ However, understanding of geometrical and electronic structures and physico-chemical properties of binary Al–Si clusters remains limited. Earlier studies mainly examined the Al doped Si clusters.^{50–52} whereas only a few studies on the Si doped Al clusters were performed and concentrated on high symmetry structures^{53,54}

that are not necessarily the most thermodynamically stable structural motifs.

In view of the scarcity of reliable results on mixed Al–Si clusters, we set out to carry out a systematic investigation, using both density functional theory (DFT) and wavefunction methods, on a series of small singly and doubly silicon doped aluminum cluster Al_nSi_m , with $n = 3\text{--}16$ and $m = 1\text{--}2$, in both neutral and cationic states. Geometries of the most stable equilibrium structures and their energetic parameters are determined by using both DFT and coupled-cluster theory CCSD(T) methods. It is of importance to emphasize the similarities and differences between the behavior of Al–Si clusters that have a similar size and number of electrons. As for a typical example, as the isoelectronic species Al_{13}^- , Al_{12}Si and $\text{Al}_{11}\text{Si}_2^+$ have closed electronic shells in the jellium model (40 valence electrons), we thus model their optical absorption spectra by using the time-dependent DFT (TD-DFT).

2. Computational methods

All standard electronic structure calculations are carried out using the GAUSSIAN 09 package.⁵⁵ The search for possible structures of clusters is conducted using different approaches. First, we generate many initial structures of Al_xSi_y , denoted (x, y) for simplicity, by modifying the known global minima (GM) geometries taken from the literature as follows: (a) addition of a Si at different positions of the $(x, y - 1)$ GM; (b) addition of Al at different positions of the $(x - 1, y)$ GM; (c) Al-to-Si substitution in $(x + 1, y - 1)$ GM, and Si-to-Al substitution in $(x - 1, y + 1)$ GM. These initial isomers are rapidly optimized using DFT with the hybrid B3LYP functional in conjunction with the 6-31G(d) basis set. The lowest-lying isomers then become the inputs for the following search using a stochastic algorithm, with the aim to survey all possible lower-lying isomers of each Al_nSi_m size.

Geometries and harmonic vibrational frequencies of the lower-lying isomers having relative energies of <10 eV of Si_nAl_m at the small sizes ($n \leq 4$) and their corresponding cations are then reoptimized with the higher spin states (triplet and quintet for closed shell clusters, and quartet and sextet for open shell clusters). However, no high spin states having a remarkable stability are found, due to the fact that both Al and Si are basically non-magnetic atoms. It can thus be predicted that the larger sizes of Al_nSi_m also exist as non-magnetic clusters, and therefore we can neglect the calculations of their high spin states due to the limitations of our computational resource.

The local minima with relative energies of <5 eV with respect to the corresponding lowest-lying isomer of each size are considered again and reoptimized using the same B3LYP functional but with the larger 6-311+G(d) basis set. Harmonic vibrational frequencies are again calculated at this level in order to identify the obtained structures as local minima on the potential energy surface and to evaluate zero-point energy corrections. To obtain more reliable energetic parameters, the lower-lying isomers of each size having relative energies within 1.0 eV are selected for single point electronic energies using the coupled-cluster theory CCSD(T) with the correlation consistent cc-pVTZ basis set, based on B3LYP optimized geometries. The



unrestricted formalism (UHF, UB3LYP, UCCSD) is employed for open-shell structures.

In an attempt to probe further the effects induced by the Si dopants on the electronic structure, we perform time-dependent density functional theory (TD-DFT) calculations using the same functional/basis set and the optimized geometries of the closed-shell isoelectronic Al_{13}^- , Al_{12}Si and $\text{Al}_{11}\text{Si}_2^+$ species. These calculations also allow us to construct their absorption spectra. For each species, the number of electronically excited states considered is large enough for the spectra spread out up to a wavelength of 250 nm. The shape of molecular orbitals involved in the electron shells and the density of states of the above clusters are also plotted assisting the analysis of their spectra.

In addition, we use the magnetically included ring current density (MICD) approach which is computed by integrating the electronic current passing through interatomic surfaces located by the atom-in-molecule theory (QTAIM) between neighboring atoms by using the AIMALL suite of programs.⁵⁶ The purpose of the latter computations is to assess the aromatic character of the Al_{13}^- , Al_{12}Si , and $\text{Al}_{11}\text{Si}_2^+$ species. The magnetic current density derived from the MICD method is performed with the magnetic field vector perpendicular the molecular plane and out of that plane. As for a convention, clockwise flow of the magnetic current stands for a diamagnetic current, and anti-clockwise flow for a paramagnetic current.

3. Results and discussion

3.1 Lower-lying isomers of Al_nSi_m clusters in both neutral and cationic states

Since there is a large number of isomers located on the potential energy surface of each size considered, we only present here some of the lowest-lying isomers whose relative energies are close to the corresponding ground state structure (<1.5 eV in relative energy). The shapes of the most stable Al_nSi_m equilibrium structures at both neutral and cationic states in comparison to the ground states of the pure aluminum clusters Al_{n+m} having the same number of electrons, their symmetry point groups are shown in Fig. 1–3. Moreover, the lower-lying isomers and their relative energies obtained at the CCSD(T)/cc-pVTZ + ZPE level are shown in Fig. S1 and S2 of the ESI.†

As for a convention, label **X.n.m.Y** is used to denote the isomers in which **X** = **N** and **C** stands for a neutral and cationic state, respectively, **n** is the number of Al atoms, **m** is the number of Si atoms, and **Y** = **A**, **B**, **C**... referring to the different isomers with increasing relative energy. Accordingly, the notation **X.n.m.A** invariably refers to the most stable isomer **A** of the **X.n.m** series.

3.1.1. Singly silicon doped $\text{Al}_n\text{Si}^{0/+}$. The most stable structures of the singly doped clusters are displayed in Fig. 1 ($n = 3$ –10) and 2 ($n = 11$ –16). Their main characteristics can be summarized as follows:

Al_3Si . (U)CCSD(T) results emphasize that both isomers **N.3.1.A** and **N.3.1.B** are basically degenerate within an energy separation of only 0.06 eV. Unlike the neutral Al_3Ti which favors a tetrahedral configuration due to the effect of the 3d orbital of

Ti dopant,²⁵ both are formed by substitution of an Al position of the planar neutral Al_4 by an Si atom.¹³ Although the isomer **N.3.1.C**, formed by adding an Al^+ cation on the Si-top of the triangular Al_2Si , is 0.38 eV less stable than **N.3.1.A**, its corresponding cation, however, becomes the ground state **C.3.1.A**, with 0.32 eV lower in energy than the next isomer **C.3.1.B**.

Al_4Si . Contrary to Si_4Al having a three-dimensional shape due to the dominance of sp^3 hybridization of Si atoms, lower-lying isomers of neutral Al_4Si still keep the planar forms in their low-spin state. In **n.4.1.A** Si takes the place of central Al atom in the neutral Al_5 (ref. 13) and connects with four remaining Al atoms. Similarly, **C.4.1.A** which is the corresponding cation of **N.4.1.A**, remains the most stable cationic isomer Al_4Si^+ .

Al_5Si . The most stable isomer **N.5.1.A** is generated by substitution of one Al atom in the octahedron of Al_6 (ref. 13) by Si. The next isomer **N.5.1.B** in which Si replaces an Al position of a triangular prism in Al_6 ,⁹ is much less stable than **N.5.1.A** by 0.40 eV. However, geometry optimizations indicate that the structure **N.5.1.A** does not exist as a cationic minimum on the potential surface; **C.5.1.A** which is actually the corresponding cation of **N.5.1.B**, becomes the ground state of the cation Al_5Si^+ . There is thus a reversed energy ordering upon ionization.

Al_6Si – Al_9Si . For the singly doped Al_nSi with $n = 6$ –9, their geometries are mostly based on the pseudo-octahedron Al_5Si in which the Si atom is placed at a top. Accordingly, the lowest-lying isomers for Al_6Si , Al_7Si , Al_8Si and Al_9Si in both neutral and cationic states have structural characteristics of Al_5Si with capping of one, two, three, and four Al atoms, respectively, on different positions of the pseudo-octahedron. (U)CCSD(T) calculations indicate an energetic degeneracy between the two most stable isomers of the neutral Al_6Si and Al_9Si with energy gaps of 0.08 and 0.04 eV, respectively.

The corresponding cations of **N.6.1.A**, **N.7.1.A**, and **N.8.1.A** remain also the lowest-lying isomers of the cationic counterparts. Again, calculated results show that the two most stable isomers exist as energetically degenerate for both cations Al_6Si^+ and Al_7Si^+ with energy gap of only 0.03 and 0.01 eV, respectively. For the cation Al_9Si^+ , **C.9.1.B** which is the corresponding cation of **N.9.1.A**, now lies 0.17 eV higher in energy than the C_{2v} symmetric structure **C.9.1.A** although **N.9.1.C** is the corresponding neutral of **C.9.1.A** and is 0.40 eV less stable than **N.9.1.A**.

Al_{10}Si . The first Si encapsulated in the Al framework occurs at $n = 10$; **N.10.1.A** appears as a doped cage in which Si is located inside a bicapped quadrangular prism Al_{10} cage. The next isomer **N.10.1.B**, being 0.32 eV less stable, has the shape of pure Al_{11} cage¹³ in which one Al position is substituted by a Si atom. For the cationic state, the most stable isomer **C.10.1.A** formed by replacement of the Al atom located at the centre of the hexagonal face of pure Al_{11} by a Si atom. According to geometry optimization, a structure having the form of **C.10.1.A** does however not exist as a neutral minimum on the potential energy surface. Thus the **C.10.1.A** is presumably formed through a rearrangement upon ionization of a neutral Al_{10}Si .

Al_{11}Si . The most stable neutral **N.11.1.A** is obtained by substitution of an Al atom located at the centre of the pure Al_{13} icosahedral framework by Si and simultaneous removal of



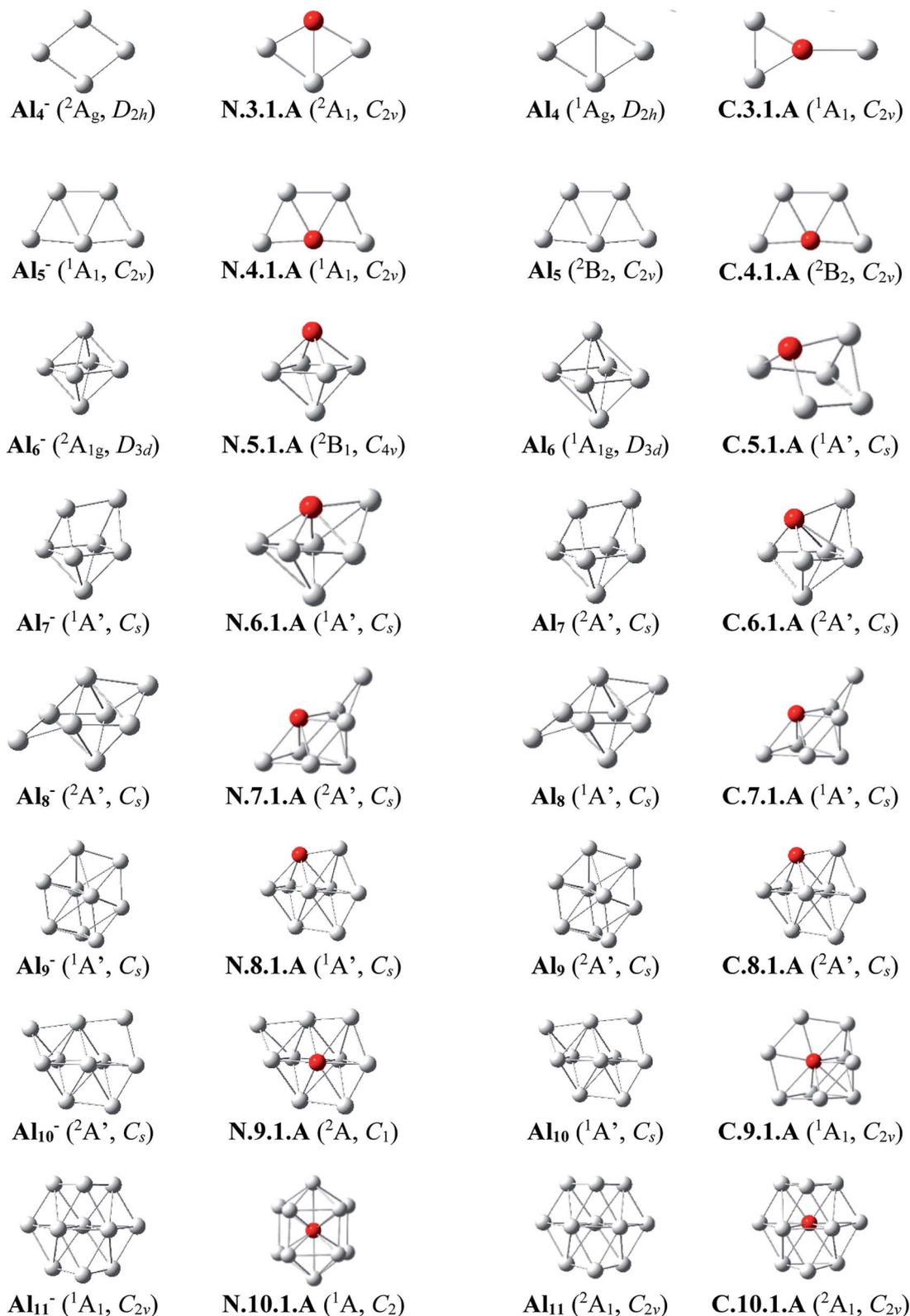


Fig. 1 The shapes and electronic states of the lowest-lying isomers Al_nSi with $n = 3-10$ and the global minimum of Al_{n+1} at the neutral and cationic states for Al_nSi , corresponding to the anionic and neutral states for Al_{n+1} , respectively.



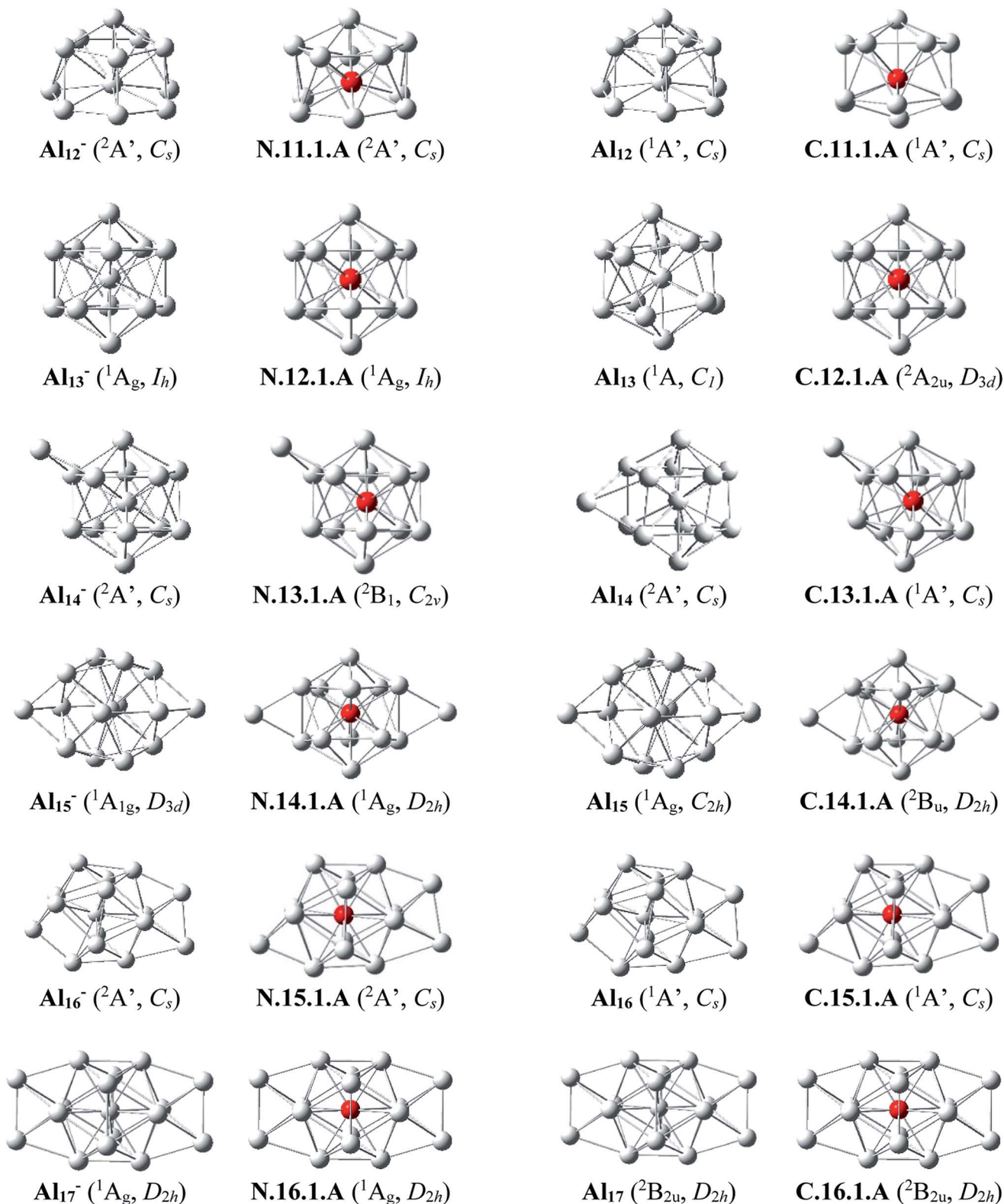


Fig. 2 The shapes, electronic states of the lower-lying isomers Al_nSi with $n = 11-16$ at the neutral and cationic states for Al_nSi , corresponding to the anionic and neutral states for Al_{n+1} , respectively.



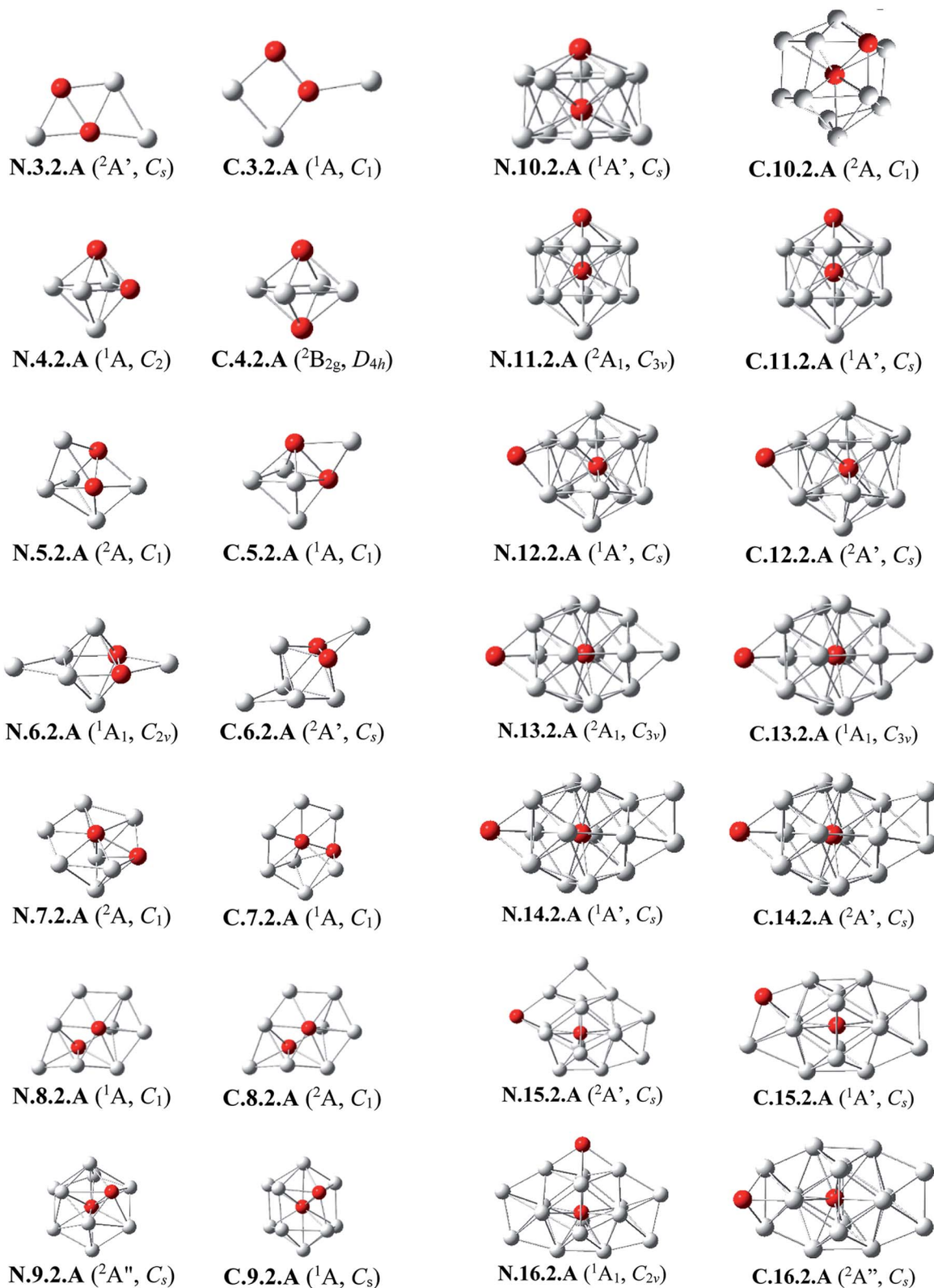


Fig. 3 The shapes and electronic states of the lowest-lying isomers Al_nSi_2 with $n = 3-16$ at the neutral and cationic states. (a) Average binding energy (E_b , eV) of the neutral Al_n , Al_nSi , and Al_nSi_2 computations. (b) Average binding energy (E_b , eV) of the cationic Al_n^+ , Al_nSi^+ , and $Al_nSi_2^+$.



another Al from it. For the cation $\text{Al}_{11}\text{Si}^+$, (U)CCSD(T) calculations indicate that the two endohedral structures, **C.11.1.A** and **C.11.1.B** are practically degenerate with a negligible energy gap of 0.01 eV.

$\text{Al}_{12}\text{Si}-\text{Al}_{16}\text{Si}$. The icosahedral growth pattern appears to dominate the structural motifs of $\text{Al}_n\text{Si}^{0/+}$ clusters for the sizes $n = 12-16$. The global minima of both neutral and cationic $\text{Al}_{12}\text{Si}^{0/+}$ possess the popular icosahedral shape in which the Si atom is centered and connects with 12 Al atoms of the icosahedron cage Al_{12} , and each is at ~ 0.6 eV more stable than the next isomer. Based on that finding, the most stable structures of neutral clusters Al_{13}Si , Al_{14}Si , Al_{15}Si and Al_{16}Si are obtained by successive capping of one, two, three, and four Al atoms on different positions of the icosahedral Al_{12}Si , and their corresponding cations remain the global minima of the positively charged state.

For the neutral Al_{14}Si , calculations continue to emphasize an energetic degeneracy with an energy gap of 0.03 eV between the high symmetry structure **N.14.1.A** (D_{2h}) and **N.14.1.B** in which a Si and an Al atom are exohedrally attached on the icosahedron Al_{13} . In other words, it can be considered that **N.14.1.B** is formed by substituting a Si atom at an Al-top of the high symmetrical (D_{3d}) anion Al_{15}^- and this may lead to the stability of **N.14.1.B**.¹² The Al_nSi can be regarded to be generated upon substitution of the central Al position of the icosahedron Al_{13} of the Al_{n+1} species, with $n = 12-16$, by a Si atom.¹²

Generally, since the Si element has one valence electron more than the Al, the neutral Si is isoelectronic with the anion Al^- . Therefore, it is observed in Fig. 1 and 2 that in small neutral Al_nSi clusters ($n \leq 9$), the Si atom prefers a substitution at an Al position, which exhibits a higher coordination number, of the pure aluminum framework of the isoelectronic anion Al_{n+1}^- . The neutral Al_{10}Si represents however an exception because its most stable structure significantly differs from the Al_{11}^- framework. The ground state structure of a cationic Al_nSi^+ cluster turns out to be also quite similar to those of the pure neutral Al_{n+1} . Moreover, for the cluster Al_nSi with $n \geq 10$, the Si dopant prefers to be located at the inner cage formed by Al atoms. According to Weigend *et al.*,⁵⁷ this can be rationalized by the fact that the Si atom has larger nuclear charge as compared to Al atom, and its valence orbitals (3s, 3p) are lower-lying in energy, more compact to its nucleus as compared to Al atom, causing a higher electron density at the center as compared to that in the surface of the Al_nSi cluster.

3.1.2. Doubly silicon doped $\text{Al}_n\text{Si}_2^{0/+}$. The main structural characteristics for these systems are displayed in Fig. 3 and 4.

Al_3Si_2 . The lowest-lying isomers of both neutral and cationic states of Al_3Si_2 are characterized by a planar structure. **N.3.2.A** in which both Si atoms replace two Al positions in the neutral Al_5 is the most stable isomer. According to geometry optimizations, the shape of **N.3.2.A** does not exist as a minimum on the potential energy surface of Al_3Si_2^+ . For the cation Al_3Si_2^+ , both isomers **C.3.2.A** and **C.3.2.B** are degenerate within a tiny energy gap of 0.07 eV. Both cationic structures are formed by adding an Al atom on a Si-top of the rhombic Al_2Si_2 .

Al_4Si_2 . The most stable structures **N.4.2.A** and **N.4.2.B** are obtained by substitution of two Al atoms in the octahedron of

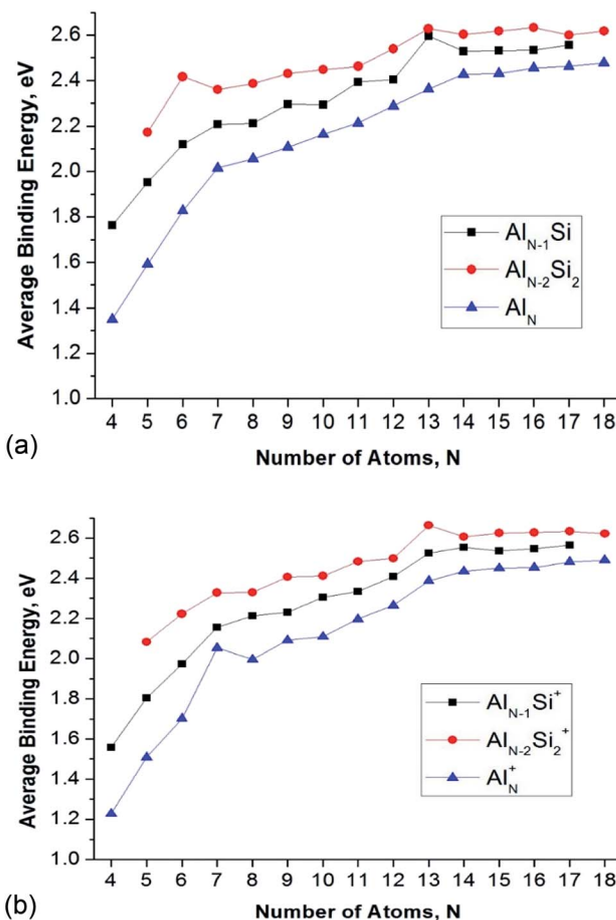


Fig. 4 Average binding energy (E_b , eV) of the $\text{Al}_n\text{Si}_m^{0/+}$ ($n = 3-16$, $m = 1-2$) clusters using (U)CCSD(T)/cc-pVTZ + ZPE. (a) MO diagram of Al_{13}^- anionic cluster. (b) MO diagram of Al_{12}Si neutral cluster. (c) MO diagram of $\text{Al}_{11}\text{Si}_2^+$ anionic cluster.

Al_6 by two Si. Again, the cation having the shape of **N.4.2.A** does not exist as a cationic energy minimum. The high symmetry (D_{4h}) isomer **C.4.2.A** generated by ionization of **N.4.2.B** emerges as the most stable cation.

$\text{Al}_5\text{Si}_2-\text{Al}_8\text{Si}_2$. Similar to the singly doped Al_nSi with $n = 6-9$, for the Al_nSi_2 clusters consisting of a total number of seven to ten atoms, their geometrical structures are based on the octahedron Al_4Si_2 in which two Si atoms replace two Al positions of the pure octahedron Al_6 . The most stable isomers for Al_5Si_2 , Al_6Si_2 , Al_7Si_2 and Al_8Si_2 in both neutral and cationic states are thus obtained by capping of one, two, three and four Al atoms, respectively, on different positions of Al_4Si_2 . Due to the presence of two Si dopant atoms, the octahedron Al_4Si_2 is however slightly distorted within a few Al_nSi_2 structures. (U)CCSD(T) calculations point out again an energetic degeneracy in several sizes including Al_5Si_2 , Al_6Si_2 , Al_6Si_2^+ , Al_7Si_2^+ and Al_8Si_2 , with the largest energy gap of only 0.08 eV between the two corresponding lowest-lying isomers.

Al_9Si_2 . From size $n = 9$, the doubly Si doped Al cluster tends to favor an icosahedral structure. **N.9.2.A** is accordingly formed following substitution of an external Al atom by a Si atom and



simultaneous removal of two other Al atoms of the icosahedron Al_{12}Si in which the Si atom is situated at its central position. For the cation, due to the fact that they have the same number of atoms and valence electrons, the most stable **C.9.2.A** has a similar shape to that of the ground state of neutral Al_{10}Si **N.10.1.A** in which the additional Si atom now replaces an Al position.

$\text{Al}_{10}\text{Si}_2$. (U)CCSD(T) results consistently point out a degeneracy in energy between both isomers **N.10.2.A** and **N.10.2.B** for the neutral Al_{10}Si , and among **C.10.2.A**, **C.10.2.B**, and **C.10.2.C** for the cation, with tiny energy gaps of 0.01 and 0.06 eV, respectively. Both neutrals **N.10.2.A** and **N.10.2.B** are generated from the same motif of Al_9Si in which an Al atom is replaced by a Si atom and another Al atom is extracted from the icosahedral Al_{12}Si . The cationic isomer **C.10.2.A** is formed by addition of an Al atom into the cation Al_9Si_2^+ **C.9.2.A**, isomer **C.10.2.B** is obtained by capping a Si atom into the neutral Al_{10}Si , and isomer **C.10.2.C** is the corresponding distorted cation of **N.10.2.A**.

$\text{Al}_{11}\text{Si}_2$. Again, the icosahedral growth pattern prevails in generating the structures of Al_nSi_2 clusters for $n = 11$ –16. Both the neutral and cationic most stable isomers $\text{Al}_{11}\text{Si}_2^{0/+}$, **N.11.2.A** and **C.11.2.A**, prefer an icosahedral structure in which one Si atom is located at the central position while the remaining Si substitutes an external Al position of the icosahedron Al_{13} and they have a remarkably high thermodynamic stability in comparison to the remaining isomers.

$\text{Al}_{12}\text{Si}_2$. Both isomers **N.12.2.A** and **N.12.2.B** are found to exist as two degenerate equilibrium ground states for the neutral $\text{Al}_{12}\text{Si}_2$ with a tiny energy difference of 0.05 eV. **N.12.2.A** is obtained by capping a Si atom onto a surface of the icosahedron Al_{12}Si framework, whereas **N.12.2.B** is generated upon capping of an Al atom onto an exohedral Al–Si edge of the icosahedron $\text{Al}_{11}\text{Si}_2$. Similarly, their corresponding cations, namely **C.12.2.A** and **C.12.2.B**, are also energetically degenerate with an even smaller energy separation of 0.01 eV.

$\text{Al}_{13}\text{Si}_2$ – $\text{Al}_{16}\text{Si}_2$. For $n = 13$ –16, the lowest-lying structures in both neutral and cationic states are constructed by preserving the icosahedron Al_{12}Si skeleton and by adding the second Si atom and remaining Al atoms onto its different external positions. The global minima of $\text{Al}_{13}\text{Si}_2$, $\text{Al}_{14}\text{Si}_2$ and their corresponding cation have a similar geometric shape. For both neutral and cationic states of $\text{Al}_{15}\text{Si}_2$ and $\text{Al}_{16}\text{Si}_2$, (U)CCSD(T) calculations result again in energetic degeneracies of the two lowest-lying isomers, with energy gap <0.1 eV.

Similar to the growth pattern of singly silicon doped $\text{Al}_n\text{Si}^{0/+}$, it can be considered that the structures of doubly silicon doped $\text{Al}_n\text{Si}_2^{0/+}$ are obtained following substitution at two Al positions of the pure corresponding aluminum framework Al_{n+2} by two Si atoms. For $n = 12$ –16, $\text{Al}_n\text{Si}_2^{0/+}$ are built up on the basis of an icosahedral growth path in which one Si atom is situated at the centre of the icosahedral Al_{12}Si , whereas the additional Si atom is now externally attached.

3.2 Relative stability of clusters

In order to evaluate the inherent thermodynamic stability of the clusters considered, the average binding energies (E_b) are

determined and compared to those of the relevant pure aluminum clusters. The average binding energies (E_b) can conventionally be defined as follows (eqn (1)–(6)):

$$E_b(\text{Al}_n\text{Si}) = [nE(\text{Al}) + E(\text{Si}) - E(\text{Al}_n\text{Si})]/(n + 1) \quad (1)$$

$$E_b(\text{Al}_n\text{Si}^+) = [(n - 1)E(\text{Al}) + E(\text{Al}^+) + E(\text{Si}) - E(\text{Al}_n\text{Si}^+)]/(n + 1) \quad (2)$$

$$E_b(\text{Al}_n\text{Si}_2) = [nE(\text{Al}) + 2xE(\text{Si}) - E(\text{Al}_n\text{Si}_2)]/(n + 2) \quad (3)$$

$$E_b(\text{Al}_n\text{Si}_2^+) = [(n - 1)E(\text{Al}) + E(\text{Al}^+) + 2xE(\text{Si}) - E(\text{Al}_n\text{Si}_2^+)]/(n + 2) \quad (4)$$

$$E_b(\text{Al}_n) = [nE(\text{Al}) - E(\text{Al}_n)]/n \quad (5)$$

$$E_b(\text{Al}_n^+) = [(n - 1)E(\text{Al}) + E(\text{Al}^+) - E(\text{Al}_n^+)]/n \quad (6)$$

where $E(\text{Si})$, $E(\text{Al})$, and $E(\text{Al}^+)$ are the total energies of the Si-atom, Al-atom and the cation Al^+ , respectively. Since the ionization energy of Al atom (1.56 eV) is lower than Si atom (2.1 eV), the total energies of the cation Al^+ is thus using to calculate the average binding energy instead of the total energies of the cation Si^+ . For their part, $E(\text{Al}_n\text{Si})$, $E(\text{Al}_n\text{Si}^+)$, $E(\text{Al}_n\text{Si}_2)$, $E(\text{Al}_n\text{Si}_2^+)$, $E(\text{Al}_n)$, and $E(\text{Al}_n^+)$ are total energies of the neutral Al_nSi , cationic Al_nSi^+ , neutral Al_nSi_2 , cationic Al_nSi_2^+ , neutral Al_n , and cationic Al_n^+ structures, respectively. All energetic values are obtained at CCSD(T)/cc-pVTZ with ZPE corrections. Geometrical parameters of the pure aluminum clusters at both neutral and cationic states are obtained from the previous studies.^{11,13} Where needed, geometries are reoptimized using (U)B3LYP/6-311+G(d) calculations. The plots illustrating the E_b evolution are displayed in Fig. 4.

The E_b value tends to rise with increasing cluster sizes. For singly doped Al_nSi clusters, the neutral Al_{12}Si reveals the largest E_b value as compared to the remaining singly doped neutral species. Of the neutral Al_nSi_2 clusters, Al_4Si_2 presents an enhanced E_b value as compared to those of the remaining doubly doped species. In the series of cations considered, $\text{Al}_{11}\text{Si}_2^+$ gets a maximum peak in the E_b plot which demonstrates its higher thermodynamic stability.

The enhanced relative stability of these clusters can simply be rationalized in terms of “magic clusters” in which the number of valence electrons amounts to 2, 8, 18, 20, 34, 40 and 58... etc. Accordingly, the closed electron shells [$1\text{S}^2 1\text{P}^6 1\text{D}^{10} 2\text{S}^2 1\text{F}^{14} 2\text{P}^6 \dots$] of the neutral Al_4Si_2 contains the same number of 20 valence electrons but an atom fewer in comparison to the pure cation Al_7^+ , while both Al_{12}Si and $\text{Al}_{11}\text{Si}_2^+$ have each 40 valence electrons and the same icosahedral structure with a total number of 13 atoms such as the anion Al_{13}^- . As a result, they behave as “magic clusters” with an enhanced thermodynamic stability.

Fig. 4 also displays a comparison of the binding energies of the pure Al_n , Al_nSi , and Al_nSi_2 using (U)CCSD(T)/cc-pVTZ + ZPE calculated results in which the E_b value of Al_nSi is larger than the E_b value of Al_n and smaller than Al_nSi_2 at the same number of atoms. Accordingly, it appears that the Si dopant tends to



Table 1 Adiabatic Ionization Energies (AIE, eV) of singly and doubly silicon doped aluminum clusters in comparison with pure aluminum clusters Al_n (CCSD(T) calculations)

Number of atoms, k	$Al_{k-1}Si$	$Al_{k-2}Si_2$	Al_k
	$N.k - 1.1.A$	$N.k - 2.2.A$	
4	6.75	—	6.43
5	6.67	6.39	6.36
6	6.80	7.10	6.69
7	6.30	6.17	5.68
8	5.94	6.40	6.42
9	6.55	6.16	6.09
10	5.83	6.31	6.49
11	6.59	5.71	6.13
12	5.88	6.43	6.23
13	6.85	5.46	5.63
14	5.60	5.88	5.84
15	5.89	5.84	5.66
16	5.71	6.03	5.97
17	5.79	5.36	5.63
18	—	5.87	6.05

increase the cluster stability with respect to various fragmentation processes.

3.3 Adiabatic ionization energies

Calculated total energies carried out at the (U)CCSD(T) level are now used to evaluate the adiabatic ionization energies (AIE), that are obtained from differences between total energies of the neutrals Al_nSi_m and their corresponding $Al_nSi_m^+$ cations. In general, the AIE values are rather low, but there is no consistent trend in going from one dopant to two dopants. In view of the frequent energetic degeneracy of many neutral sizes, Table 1 displays the AIE values derived from the two most stable isomers Al_nSi_m and these values are compared to those of pure aluminum clusters. Due to the close AIE values, the corresponding photoelectron (PE) spectra of these neutrals are

expected to be rather complicated having broad bands. Only an appropriate treatment considering the vibrational progressions upon ionization can usefully predict their PE spectra. Let us note that both “magic clusters” Al_4Si_2 and $Al_{12}Si$ are remarkably characterized by the largest AIE values, being 7.1 and 6.9 eV, respectively, that are no doubt due to an enhanced stability of these neutrals.

3.4 Dissociation energies

To evaluate further the thermodynamic stability, the dissociation energies (D_e) for the various fragmentation channels of the clusters considered are determined. Results calculated from (U)CCSD(T) energies are listed in Table 2.

The D_e values of the neutrals Al_nSi for the Si removal channel (1) $Al_nSi \rightarrow Al_n + Si$ turn out to be larger than those for the Al-loss channel (2) $Al_nSi \rightarrow Al_{n-1}Si + Al$ proving the stronger Si–Al bond. From the obtained knowledge in solid state, the loss of Al is a lower energy channel than the loss of Si, and this can be understood by the fact that the cohesive energy of Si (4.63 eV per atom) is larger than that of Al (3.39 eV per atom). Similar observations can be made for the positively charged species in that Al_nSi^+ tend to be fragmented generating one Al element or its cation Al^+ plus a smaller $Al_{n-1}Si^{+/0}$ along the fragmentation channels (2) and (4).

Again, for doubly doped neutral Al_nSi_2 clusters, dissociation energies for Si-loss channels (1) $Al_nSi_2 \rightarrow Al_nSi + Si$ are constantly larger than those for Al-elimination pathways (2) $Al_nSi_2 \rightarrow Al_{n-1}Si_2 + Al$. Similarly, the cations $Al_nSi_2^+$ follow a preferential fragmentation to form one $Al^{0/+}$ plus a smaller $Al_{n-1}Si^{+/0}$ as described in the channels (2) and (4) (*cf.* Table 2).

Moreover, from the values given in Table 2, we can infer two exchange reactions:



and

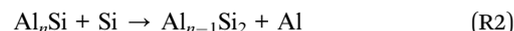


Table 2 Dissociation energies (D_e , eV) for various fragmentation channels of Si_nAl_m ($n = 3-16$, $m = 1-2$) from CCSD(T)/cc-pVTZ calculations

n	$D_e(1)$ remove a Si				$D_e(2)$ remove a Al				$D_e(3)$ remove a Si^+		$D_e(4)$ remove a Al^+	
	Al_nSi	Al_nSi^+	Al_nSi_2	$Al_nSi_2^+$	Al_nSi	Al_nSi^+	Al_nSi_2	$Al_nSi_2^+$	Al_nSi^+	$Al_nSi_2^+$	Al_nSi^+	$Al_nSi_2^+$
3	3.56	3.22	3.82	4.18	n/a	n/a	n/a	n/a	4.91	5.53	n/a	n/a
4	4.36	4.12	4.74	4.31	2.71	2.79	3.63	2.92	5.79	5.74	1.98	2.47
5	4.75	4.31	3.81	4.44	2.95	2.82	2.02	2.96	6.06	5.75	2.09	1.79
6	4.49	4.88	3.64	3.54	2.74	3.24	2.57	2.34	6.29	5.35	2.38	2.11
7	3.59	3.33	4.17	3.95	2.25	2.61	2.77	3.02	5.76	6.12	2.25	2.56
8	4.22	4.09	3.81	4.06	2.97	2.36	2.62	2.46	5.77	5.61	2.36	2.25
9	3.97	4.23	4.16	4.27	2.26	2.99	2.61	3.2	6.25	6.55	2.37	2.83
10	4.68	4.58	4.16	4.32	3.39	2.62	3.39	2.67	6.2	5.83	2.73	2.9
11	4.49	4.75	5.32	5.73	2.53	3.24	3.68	4.65	6.72	7.96	2.59	4.16
12	6.28	5.66	2.7	3.67	4.9	3.93	2.28	1.86	7.53	4.92	3.99	2.34
13	4.69	4.72	3.87	3.63	1.66	2.91	2.83	2.88	7.2	6.14	2.00	2.93
14	4	3.94	4.15	4.01	2.58	2.29	2.86	2.67	6.22	6.23	2.63	2.77
15	4.07	4.01	3.66	4.02	2.55	2.73	2.06	2.74	6.46	6.41	2.78	2.64
16	4.19	4.37	3.66	3.59	2.93	2.85	2.93	2.42	6.5	5.9	3.08	3.01



The energies the exchange reaction (R_1) and (R_2) can easily be calculated *via* the differences between $D_e(1)$ and $D_e(2)$ of Al_nSi and Al_nSi_2 . Generally, all energetic values of (R_1) and (R_2) are approximately in a range of 0.5–2 eV. Remarkably, the maximum value of 3.03 eV, obtained at $n = 12$ for (R_1), corresponds to that of $Al_{12}Si$. It proves the very particular case of neutral $Al_{12}Si$ for which the energy of (R_1) is much higher than for any other case.

3.5 Electronic structures of the isoelectronic Al_{13}^- , $Al_{12}Si$, and $Al_{11}Si_2^+$ having 40 valence electrons

While a single atom possesses discrete electron energy levels and bulk solids have continuous energy levels, the electron energy levels of atomic clusters could be distributed in a non-uniform manner. Clusters having high symmetry frequently result in pronounced electron shells, such as 1S, 1P, 1D, 2S, 2P, 1F... This formation of electron shells is manifested in many physical properties of finite, quantal many-electron systems, such as, for example, their stability, ionization energies, chemical reactivity, or conductance, *etc.* A brief review of the abundant literature on the phenomenon reveals that both cationic Al_7^+ and anionic Al_{13}^- aluminum clusters (Scheme 1) behave as “magic clusters” with enhanced thermodynamic stability, in part due to the fact that they possess the closed electron shells including 20 and 40 valence electrons, respectively. As stated above, the neutral Al_{13} appears as a ‘superhalogen’ characterized by a very large electron affinity exceeding those of halogen atoms.^{8,15,58–60}

Of the whole series of the Al_n^- , $Al_{n-1}Si$, and $Al_{n-2}Si_2^+$ clusters, let us consider a typical series of high symmetry isoelectronic structures Al_{13}^- , $Al_{12}Si$ and $Al_{11}Si_2^+$ to emphasize on how an assemblage of degenerate or nearly degenerate levels is manifested in their optical absorption spectra.

In order to investigate the electronic structure of Al_{13}^- , $Al_{12}Si$, $Al_{11}Si_2^+$, their MO diagrams are first plotted and displayed in Fig. 5. The electron configuration of the outer electron shell of each cluster could then be written as [...1S² 1P⁶ 1D¹⁰ 2S² 2P⁶ 1F¹⁴...], in which the high fold degeneracy levels such as the 1F, 1D, even the 1P and 2P could be split and mixed with other levels. To confirm that the number of 40 electrons in each cluster shell are arising from the valence electrons of Al atoms, we calculate the contributions of atomic orbitals (AO) of Al atoms to the shell MOs by using Natural Bonding Orbital (NBO) analysis.

The contributions of s, p and d AOs of Al atoms into the shell MOs of the anion Al_{13}^- are found as follows: 1S = 94% s + 5% p + 0.1% d; 1P = 81% s + 18% p + 0.22% d; 1D = 89% s + 10% p + 0.49% d; 2S = 97% s + 3% p + 0.25% d; 1F (three lower 1F shell orbitals) = 94% s + 5% p + 0.22% d; 2P = 10% s + 89% p + 0.74% d and 1F (four higher ones) = 0% s + 98% p + 1.6% d. This result shows that the lower shell orbitals 1S, 1P, 1D, 2S, 1F (three lower ones) are mainly constructed from the s(Al) AOs and the higher shell orbitals 2P and 1F (four higher orbitals) are mainly constructed from p(Al) AOs. In other words, the 26 s-electrons of 13 Al atoms form the lower shells [1S² 1P⁶ 1D¹⁰ 2S² 1F⁶...] and 14 p-electrons including the adding one (to form the negative

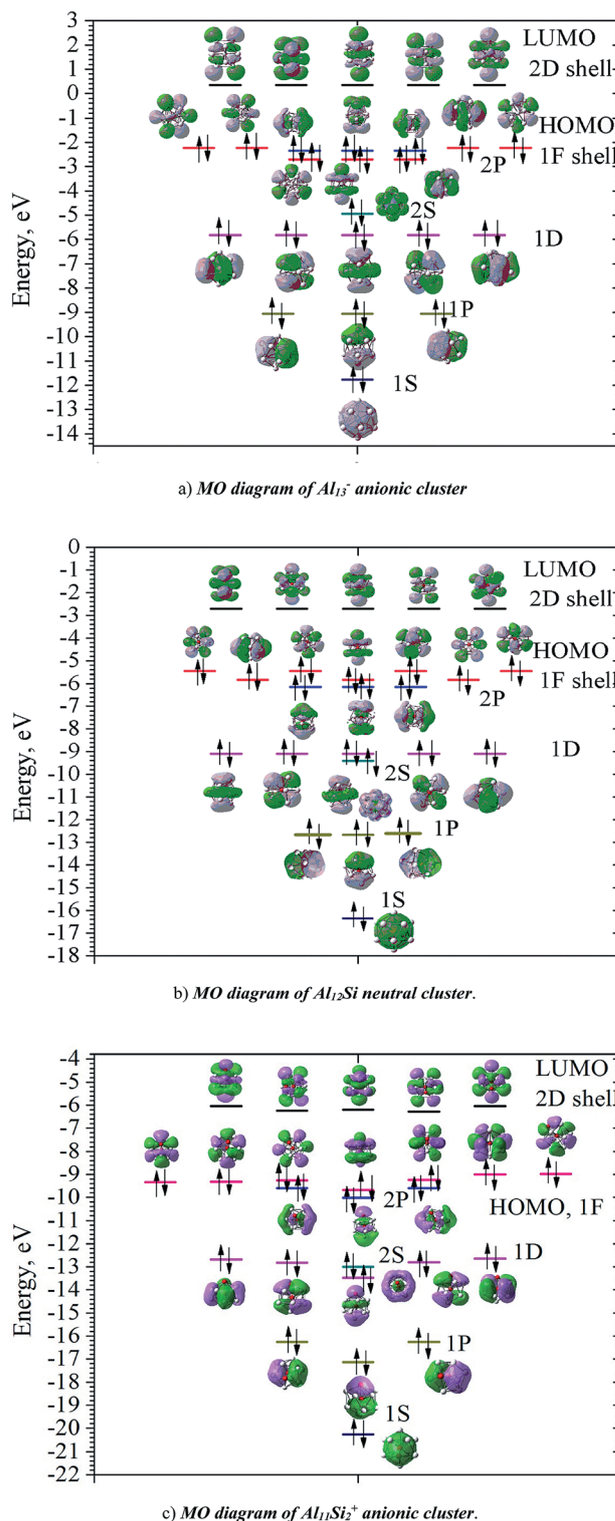


Fig. 5 MO diagram of Al_{13}^- , $Al_{12}Si$, and $Al_{11}Si_2^+$ cluster. Black lines represent LUMO; the filled pink, blue, dark cyan, magenta, dark yellow and navy lines represent orbitals corresponding to the 1F, 2P, 2S, 1D, 1P, and 1S shells, respectively. Black arrows represent electrons (alpha spin: up arrow; beta spin: down arrow).



charge) form the higher shell [$2P^6 1F^8$]. It is worth to note that while for the single Al atom, the $3s$ – $3p$ energy gap is 3.60 eV,⁵⁰ in the Al_{13}^- cluster, according to our calculations, the gap between the $1F$ lower shell, which is formed from $3s(Al)$ AOs and the $2P$ shell which is formed from $3p(Al)$ AOs, is now reduced to 0.36 eV.

Similarly, the contributions of s , p and d AOs of Al and Si atoms into the shell MOs of $Al_{12}Si$ are found as follows: $1S = 90\% s(Si) + 9\% s(Al)$; $1P = 22\% p(Si) + 12\% p(Al) + 66\% s(Al)$; $2S = 96\% s(Si) + 4\% s(Al)$; $1D = 8\% p(Al) + 91\% s(Al)$; $2P = 40\% p(Si) + 8\% p(Al) + 52\% s(Al)$; $1F = 6\% p(Al) + 94\% s(Al)$ (three lower orbitals); and $1F = 2\% d(Al) + 98\% p(Al)$ (four higher orbitals). Introduction of a Si atom into the center of Al_{12} also splits the $1F$ level into two sub-levels in which one sub-level has four-fold degeneracy (higher $1F$) and the other is three-fold (lower $1F$), as it is shown in Fig. 5a and b. The energy gap between them is ~ 0.47 eV for Al_{13}^- and ~ 0.36 eV for $Al_{12}Si$. Unlike the $1F$ shell orbitals, the shells $1P$, $1D$ and $2P$ do not split upon introduction of a Si atom into the center of the Al_{12} icosahedron. The shapes of higher $1F$ shell-orbitals (four-fold degeneracy) are similar to those of the $4f(y(3x^2 - y^2))$, $4f(yz^2)$, $4f(xz^2)$ and $4f(x(3y^2 - x^2))$, and the shapes of lower $1F$ shell orbitals are similar to those of $4f(xyz)$, $4f(z(x^2 - y^2))$ and $4f(z^3)$ AOs. The splitting of $1F$ shell orbitals described above can be understood on the basis that the central ions (Al^{3+} and Si^{4+}) cause the orbitals with small angular momentum (they have larger densities in the inner part) more energetically favorable.

Herein the three lower $1F$ shell orbitals are constructed mainly from $s(Al)$ AOs having small angular momentum, and the four higher $1F$ shell orbitals are composed of $p(Al)$ AOs having larger angular momentum. It is also worth noting that in the formation of $2P$ shell orbitals, the $3p$ AOs of the Si atom contribute up to 40% and the contribution of $3s$ AOs of Al atoms drops to 52%, while in the formation of the three lower $1F$ shell orbitals, the $3s(Al)$ AOs are the dominant contributor (94%). This constitutes a reason that in the $Al_{12}Si$ cluster, the $2P$ orbitals are located below the three lower $1F$ orbitals as it is shown in Fig. 5b. Similarly, the MO diagram of the $Al_{11}Si_2^+$, also having 40 valence electrons, is shown in Fig. 5c and described in detail in ESI.†

As for a comparison, orbital energies of the Al_{13}^- , $Al_{12}Si$, and $Al_{11}Si_2^+$ clusters are also illustrated in Fig. 5. Let us briefly state again that, for the Al_{13}^- anion and the $Al_{12}Si$ neutral, owing to an icosahedral structure with I_h symmetry, orbital degeneracy is observed for each of the $1P$, $1D$ shells, and the $1F$ shell which is split into two subshells. As compared to the case of neutral $Al_{12}Si$ cluster, orbital energy levels of the anion Al_{13}^- are significantly higher due to the repulsion induced by the extra electron.

For $Al_{12}Si$, the level ordering of $2S$ and $1D$ is reversed (relative to those in Al_{13}^-), and the $2P$ levels obviously get lower energy. For $Al_{11}Si_2^+$, the $2P$, $1D$ and $1F$ states markedly split due to the Si substitution and the lower symmetry, and a mixture of both $2P$ and $1F$ orbitals, as well as a mixture of the $2S$ level with the $1D$ levels are observed. However, the widths of the $1D/2S$ and $1F/2P$ sub-bands are still narrower than the spacings between other bands. In fact, this is the main feature for keeping the jellium

model valid. The ellipsoidal shell structures in metal clusters were previously discussed by Brack and Clemenger^{61,62} and crossovers of the subshells in the deformed structures are common.

Calculated results also show that the clusters considered probably become semiconductors as suggested by the HOMO–LUMO energy gap ~ 2.6 eV, though aluminum in its bulk form is a good conductor. The HOMOs of these clusters are the $1F$ and $2P$ shells, while the LUMOs are the $2D$ shells allow us to expect that electronic transitions from HOMO to LUMO is allowed $\Delta = \pm 1$.

Moreover, our TD-DFT calculations indicate that the absorption spectra of the clusters considered, as illustrated in Fig. 6 and listed in Table S1 in ESI,† have two main characteristic absorption bands in the visible region. The absorption spectrum of the Al_{13}^- cluster is characterized by two strong bands which are centered at ~ 345 (peak A) and ~ 435 nm (peak B). The first peak emerges at ~ 345 nm due to electronic transitions from $2P$ to $3S$ shells. The second band corresponds to electronic transitions collectively from the $1F$ shell orbital (HOMOs) as well as $2P$ shell orbitals to the $2D$ shell orbitals (LUMOs).

It is worth noting that the absorption spectra of small ($n \leq 5$) Al_n clusters are dominated by discrete orbital transition, while for the larger size clusters, more collective electron transitions are responsible for their absorption spectra.⁶³

The absorption spectra of both clusters $Al_{12}Si$ and $Al_{11}Si_2^+$ are very similar to each other and characterized by two absorption bands centered at 316–403 and 403–394 nm, respectively. Analogous to the case of Al_{13}^- anion, the longer wavelength absorption band at 403–394 nm is also due to electronic transitions collectively occurred from the $2P$ and $1F$ –the higher shell orbitals to the $2D$ shell orbitals. The $2P \rightarrow 3S$ electronic transition is responsible for the shorter wavelength peak. As compared to the Al_{13}^- anion, the corresponding absorption peaks of these two Si-doped clusters shift to the shorter wavelength (Fig. 6). This is reasonable because the metal Al atom(s) are substituted by the semi-metal Si atom(s)

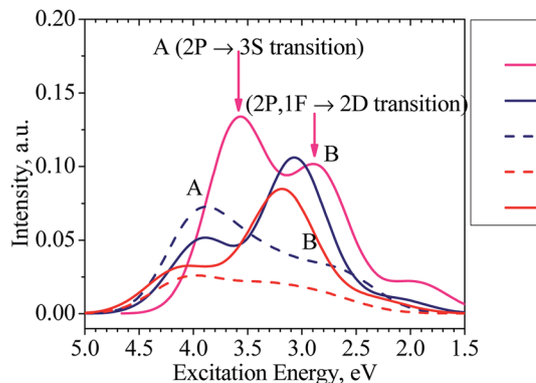


Fig. 6 Absorption spectra of the clusters Al_{13}^- (magenta line), $Al_{12}Si_{10}^0$ (navy solid line for isomer N.12.1.A and navy dashed line for isomer N.12.1.B), and $Al_{11}Si_2^+$ (red solid line for isomer C.11.2.A and red dashed line for isomer C.11.2.B).



and the net charge of the cluster is changed from negative to neutral and positive values.

For the purpose of comparison, the absorption spectra of the second isomers of the Al_{12}Si , isomer **N.12.1.B**, (D_{5h}) and $\text{Al}_{11}\text{Si}_2^+$ **C.11.2.B**, (C_1) clusters are also computed and plotted in Fig. 6. Similar to other isomers, the absorption spectra of **N.12.1.B** and **C.11.2.B** are also characterized by two absorption bands (A and B) in the visible region. For these isomers, the electronic transition from $1F \rightarrow 2D$ (HOMO \rightarrow LUMO) gives rise to a low intensity absorption band centered at around 450 nm. While the absorption band A remains almost unchanged (*cf.* Fig. 6), the absorption band B shifts from higher to lower intensity and from shorter to longer wave length, as compared to the isomers

N.12.1.A and **C.11.2.A**. In view of the fact that **N.12.1.B** is 0.66 eV less stable than **N.12.1.A** and **C.11.2.B** is 0.43 eV higher in energy than **C.11.2.A**, their absorption bands (Fig. 6) are probably not observed, in case that the isomers are thermally populated. However, for other sizes where both lowest-lying isomers are energetically degenerate, each resulting UV-Vis absorption spectrum is likely produced by a superposition of both spectra of both corresponding **A** and **B** isomers.

The magnetic maps of Al_{13}^- , Al_{12}Si and $\text{Al}_{11}\text{Si}_2^+$, obtained by using the MICD approach, are shown in Fig. 7. The clockwise flows of the magnetic current point out that Al_{13}^- , Al_{12}Si , and $\text{Al}_{11}\text{Si}_2^+$ are aromatic structures thanks to the presence of 40 valence electrons. Furthermore, the MICD maps for each shell

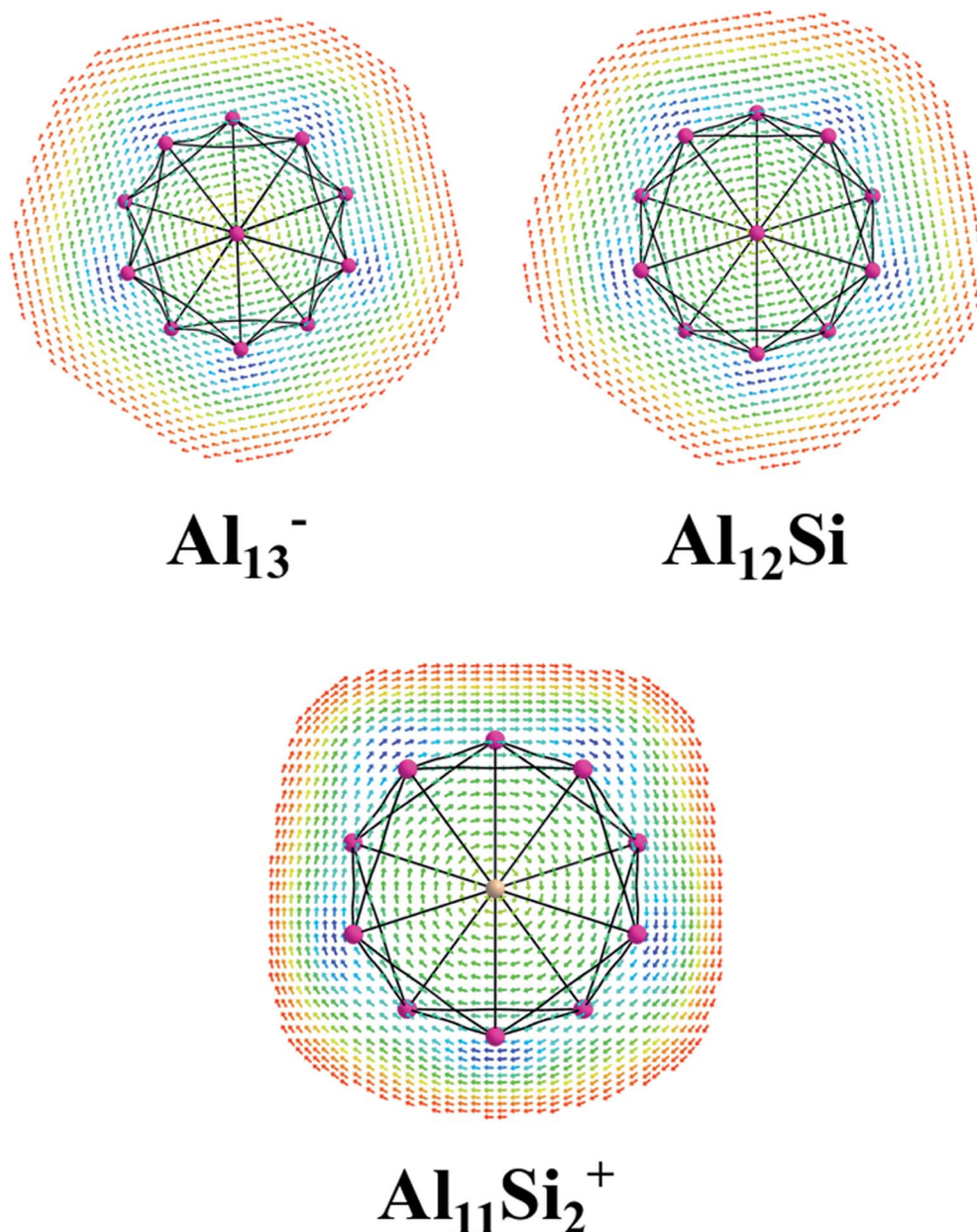


Fig. 7 Profiles of current density for Al_{13}^- , Al_{12}Si , and $\text{Al}_{11}\text{Si}_2^+$ at the planes located above the center atoms by one Bohr. Red to blue arrows represents weak to strong current density with the range: 0 to 0.0006 a.u.



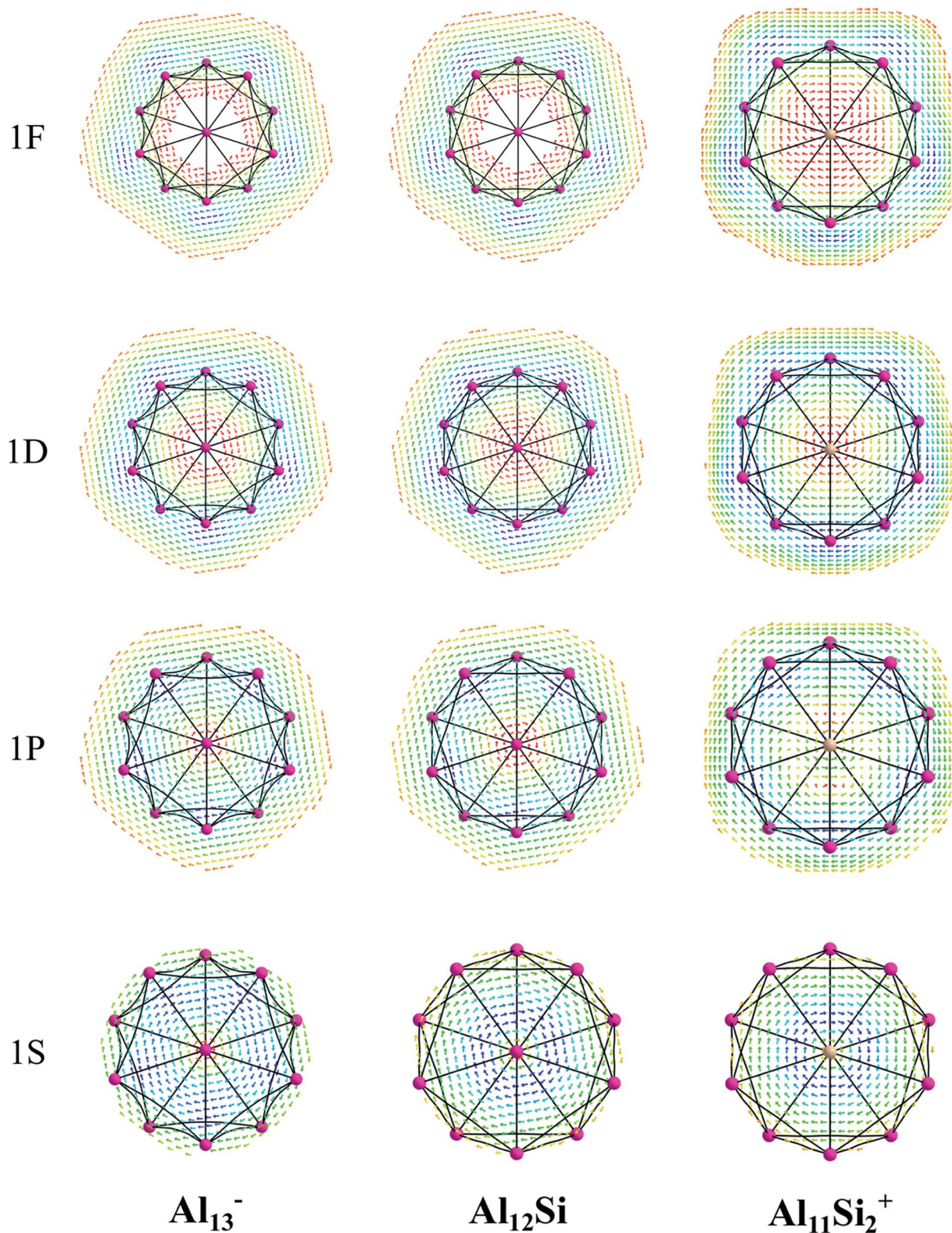


Fig. 8 Profiles of current density for Al_{13}^- , Al_{12}Si , and $\text{Al}_{11}\text{Si}_2^+$ for the 1S, 1P, 1D, and 1F shell at the planes located above the center atoms by one Bohr. Red to blue arrows represents weak to strong current density.

of Al_{13}^- , Al_{12}Si and $\text{Al}_{11}\text{Si}_2^+$, that are shown in Fig. 8 and 9, indicate that they are all aromatic. The blue arrows correspond to the strongest current density in each map. Fig. 8 displays the

MICD maps for the 1S, 1P, 1D, and 1F shells of Al_{13}^- , Al_{12}Si , and $\text{Al}_{11}\text{Si}_2^+$. The positions of blue arrows in these structures move from the center to the outer of each structure when the electron



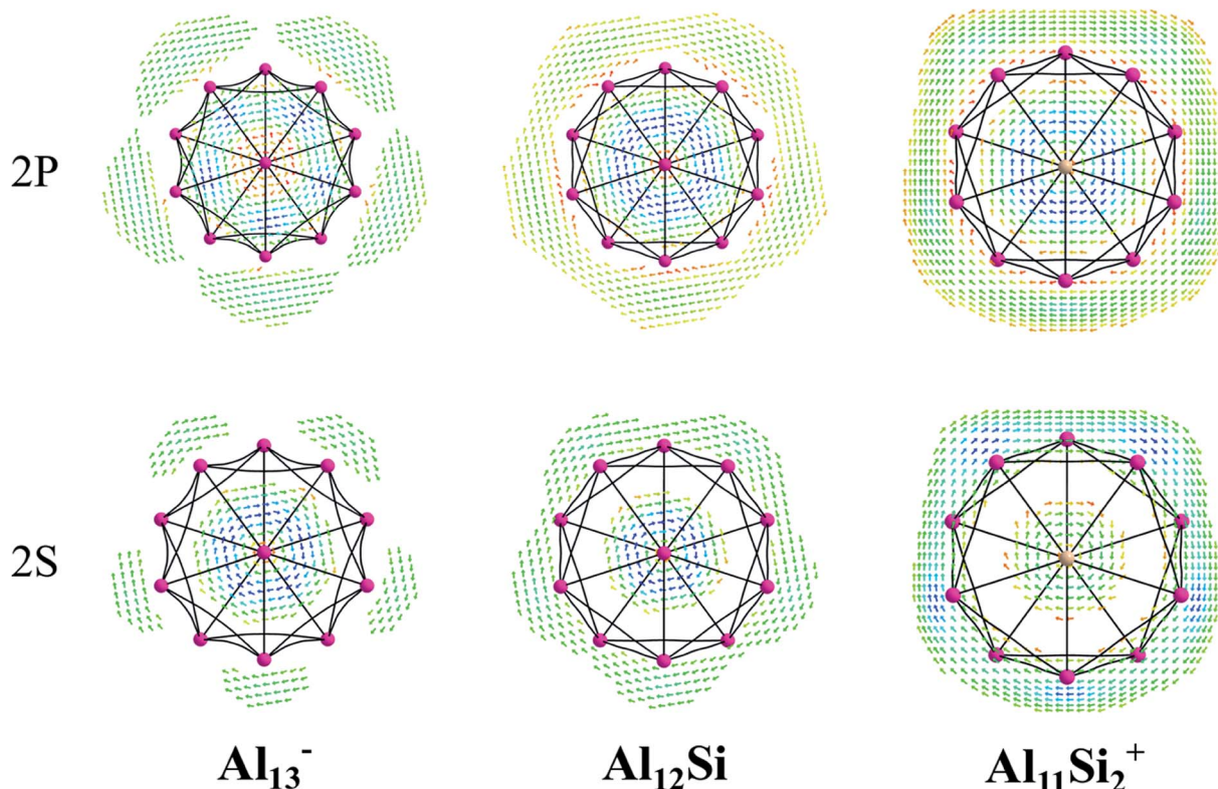


Fig. 9 Profiles of current density for Al_{13}^- , Al_{12}Si , and $\text{Al}_{11}\text{Si}_2^+$ for the 2S, and 2P shell at the planes located above the center atoms by one Bohr. Red to blue arrows represents weak to strong current density.

shell change from the 1S to 1P to 1D and to 1F. The MICD maps confirm the validity of the shell model for these structures. As for a comparison, the MICD maps for both 2S and 2P shells are shown in Fig. 9. Obviously, the profiles of MICD maps for all the shells considered in Fig. 8 and 9 denote the electrons of 2S and 2P shells effect at both inside and outside the cage, whereas the electrons of 1S, 1P, 1D and 1F shells only induce effect within the cage of clusters.

4. Concluding remarks

In the present theoretical study, geometrical structures of lower-lying isomers of Al_nSi_m clusters in both neutral and cationic states were investigated using DFT, TD-DFT with the (U)B3LYP/6-311+G(d) functional/basis set, and coupled-cluster theory (U)CCSD(T)/cc-pVTZ calculations.

All the doped clusters considered are characterized by low spin ground states, singlet for closed-shell species, and doublet for open-shell counterparts. For small size clusters, the Si atom prefers to be located at vertices having many edges. For larger sizes, the Si atom prefers to be endohedrally doped inside the corresponding Al_n cages.

A typical characteristic in most Si doped Al clusters is a consistent degeneracy of energies of the two lowest-lying isomers, either in the neutral or the cationic state. Such a characteristic suggests the coexistence of two distinct isomeric equilibrium structures, which is expected to give rise to

complicated spectra, such as vibrational, electronic or photo-electron spectra.

Relative stabilities of the clusters considered, as compared to their neighbours, were also determined. Adiabatic ionization energy and dissociation energy of each cluster size were evaluated. Calculated results confirm the high stability of the Al_4Si_2 , Al_{12}Si and $\text{Al}_{11}\text{Si}_2^+$ species as “magic clusters”, that are clusters having 20 or 40 valence electrons distributed over their outer electron shells.

The UV-Vis absorption spectra of the isoelectronic magic clusters Al_{13}^- , Al_{12}Si , and $\text{Al}_{11}\text{Si}_2^+$ have also been modelled, showing that they have two pronounced bands centered in the regions of ~ 440 and ~ 350 nm, corresponding to blue light and violet light, respectively. A rationalization for the absorption spectra was assisted by an analysis of the jellium shell model of the clusters involved. Moreover, MICD maps reveal that the isoelectronic series Al_{13}^- , Al_{12}Si and $\text{Al}_{11}\text{Si}_2^+$, having an aromatic shell, are aromatic structures possessing 40 valence electrons.

Conflicts of interest

The authors declare no competing financial interest.

Acknowledgements

This research is funded by Foundation for Science and Technology Development of Ton Duc Thang University (FOSTECT),



website <http://fostect.tdtu.edu.vn>, under Grant FOSTECT.2017.BR.03.

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