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$B_xGe_{12}^{0/+}$ Clusters with x = 1-4: Germanium Tubes Stabilized by Three and Four Boron Dopants

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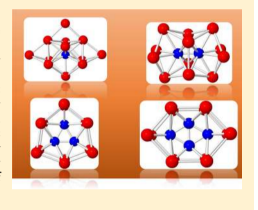
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Supporting Information

ABSTRACT: Some boron-doped germanium clusters $B_xGe_{12}^q$ with x = 1, 2, 3, 3and 4 and q = 0, 1 were designed as stabilized double ring tubes. While the B_2Ge_{12} constitutes the smallest deltahedral germanium cluster, both B_3Ge_{12} and B₄Ge₁₂ clusters present us, for the first time, with an endohedral tubular motif in which either the B_3 or the B_4 cycle is encapsulated inside a Ge_{12} hexagonal prism tube. Both B₃ and B₄ units thus satisfy a geometry requirement to create an endohedral structure within the Ge₁₂ double ring. Keeping their high symmetry, both B₃ and B₄ units generate delocalized bonds upon interaction with the Ge12 tubular framework and thereby induce an aromatic character for the resulting $B_3Ge_{12}^+$ and B_4Ge_{12} , respectively. Their aromaticity was probed by the magnetic responses of electron densities. Such a tubular aromaticity appears to greatly contribute to the high thermodynamic stability of the binary hexagonal germanium tubes.



1. INTRODUCTION

Compounds based on the germanium element are used for applications in semiconductors and optoelectronic industries.^{1,2} Development of new and tiny electronic devices has attracted much interest in the basic studies of the geometric, electronic, thermodynamic, and spectroscopic properties of small Ge clusters and their doped varieties.³⁻¹³ The main objective of such investigations is a search for appropriate cluster units that can be used as building blocks for different assembly materials whose properties can effectively be controlled by a change of the cluster size and its charge as well as the nature of the dopants.

According to numerous previous investigations, the interplay between the dopants and Ge hosts controls the thermodynamic stability and provides a structural flourish for doped Ge clusters. The structural richness perspective of Ge-based clusters is clearly illustrated by, a. o., the geometry of $[M_3Ge_{18}]^q$ clusters, with M = Au, Ni, and Pd, where the Ge_{18} counterpart is actually a combination of two Ge_9 units.^{14–16} Three Ni atoms generate a trimeric filament in $[Ni_3@Ge_{18}]^{4-}$, whereas Au₃ and Pd₃ each form a trimeric cycle in $[Au_3Ge_{18}]^{5-}$ and $[Pd_3Ge_{18}]^{2-}$ clusters, respectively. However, introduction of two Pd atoms to the Ge_{18} host establishes [Pd2@Ge18]4- as the largest single-cage deltahedron in which the Pd_2 dimer is covered by a deltahedral Ge_{18} .

Recently, a similar structural motif is found for $[Co_2@Ge_{16}]^{4-}$ clusters. X-ray experiments that were supported by density functional theory (DFT) computations, pointed out that $[Co_2@Ge_{16}]^{4-}$ exists in two different isomers in which both Co atoms are encapsulated by Ge₁₆ cages.^{18,19}

Beside such deltahedral structures, singly transition metaldoped germanium clusters provide us with a wide range of geometrical features. It is peculiar that the singly doped CoGe10³⁻ and FeGe10³⁻ clusters are stable in a double-ring tube in which either the Co or the Fe atom is centered in a D_{5h} (5/5) Ge₁₀ pentagonal prism.^{20,21} Another remarkable feature is the high stability of $M@Ge_{16}$ (M = Ti, Zr, and Hf) Frank-Kasper polyhedrons.²² In the latter, a transition metal dopant is encapsulated by a Ge₁₆ cage and subsequently induces a high symmetry T_d M@Ge₁₆. A large number of systematic investigations were carried out to elucidate the structural evolution of singly doped germanium clusters at various charged states.²³⁻³³ Accordingly, the interplay between the metal dopant and the Ge-host gives rise to a geometrical richness which varies from incomplete cage through the encapsulated form, and then to the Frank-Kasper polyhedron.

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Within the Ge-based garden, the doped Ge-clusters containing 12 Ge atoms emerge as the most beautiful flower owing to their typical diversity in structure and chemical bonding phenomena. A hexagonal prism shape was found for V@Ge₁₂⁻, Mo@Ge₁₂, and W@Ge₁₂ indicating a certain similarity of germanium and silicon clusters.^{30,31,34} Such an incomplete cage was identified for M@Ge₁₂^q and particularly their shape is quite sensitive with respect to the charge state. Interestingly, the AuGe₁₂⁻ anion cluster presents a high symmetry structure in which the Au dopant is encapsulated by an I_h Ge₁₂ host.³⁵

Recently, a hexagonal cylinder was found for the Si₁₂ cluster when it is doped by three boron atoms giving the three-layer tube $B_3(@Si_{12}^{+,36,37})$ This result brings in an interesting possibility that small boron clusters such as B_2 , B_3 , and B_4 can serve as doping units in an efficient approach to generate stable tubular clusters. In this context, we set out to expand the scope of our approach in successfully stabilizing germanium clusters containing 12 Ge atoms into a hexagonal tubular prism using the effects of both boron clusters B_3 and B_4 as dopants. According to our literature survey, both the $B_3Ge_{12}^+$ and B_4Ge_{12} cases represent, for the first time, endohedral motifs in which all dopants are encapsulated within a Ge_{12} hexagonal tube. In other words, the unstable Ge double hexagonal ring tube is stabilized following doping of B_3 and B_4 units.

2. COMPUTATIONAL DETAILS

In order to construct the potential energy surfaces of $B_x Ge_{12}^{0/+}$ clusters, we use a stochastic genetic algorithm developed earlier by us.³⁸ We now modify this algorithm by adding a permutation subroutine in which each atom exchanges its position with all the others. Moreover, to ensure that the global energy minimum isomer of each size is correctly found, several series of geometries having nonconventional shapes are also considered with the aim to search for unexpectedly stable structures. The initial structures are generated without any constrain on the largest distance of B-B, B-Ge, and Ge-Ge connectivities, whereas their shortest distances are the sum of covalence radii. We use DFT with both hybrid TPSSh and B3P86 functionals^{39,40} for geometry optimizations in view of previous results that these functionals provide the reliable results.^{41,42} Initial structures are first optimized using the small split-valence 3-21G basis set.^{43,44} The optimized isomers, whose relative energies are lying within a range of 50 kcal/mol with respect to the lowest-lying one, are subsequently reoptimized using the same functionals, but in conjunction with the larger $6-\overline{3}11+G(d)$ basis set.^{45,46} Harmonic vibrational frequencies are analyzed afterward at the same level of theory to ensure the character of the optimized structures as energy local minima and to estimate their zero-point vibrational energies. In order to explore the chemical bonding of the clusters considered, their electron density will be explored by means of the electron localization function (ELF) method,⁴ Mayer bond order (MBO),⁴⁸ as well as natural bond orbital (NBO) atomic charge.⁴⁹ All standard electronic structure theory computations are performed using the Gaussian 09 suite of program.⁵⁰

The magnetic response of the electron density is calculated using the CTOCD-DZ method^{51,52} implemented in the SYSMO program,^{53,54} which is connected to the GAMESS-UK package.⁵⁵ The magnetic ring current maps are then constructed using the B3P86 functional with the 6-311G(d) basis set. In each current density map, the contour and shading show the modulus of the induced current density, and the arrows display its projection on the plotting plane. As for a convention, anticlockwise or clockwise circulations of electron flows correspond to diatropic or paratropic currents, respectively. A diatropic current flow is correlated with an aromatic character, whereas a paratropic current flow suggests an anti-aromatic character.

3. RESULTS AND DISCUSSION

In the search for Ge clusters having a tubular form, we investigate the B_xGe_{12} systems in which x = 1, 2, 3, and 4. As the open-shell doublet state in both neutral BGe_{12} and B_3Ge_{12} radicals usually leads to a distorted geometry, we only report on the corresponding cations BGe_{12}^+ and $B_3Ge_{12}^+$ that exhibit a closed-shell singlet ground state. Let us first briefly report on their geometrical aspects before analyzing their electronic structure. As for a convention, structures mentioned in the following sections are denoted by **x**. **Y** where **x** stands for the number of B atoms, being 1, 2, 3, and 4, and **Y** = A, B, and C, ... for the isomers with increasing relative energy. Thus, structure **x**. **A** invariably refers to the lowest-energy isomer for the dopant B_{xx} , and relative energy of other isomers mentioned are given, unless otherwise stated, with respect to its corresponding **A** isomer.

3.1. BGe_{12}^+ . The most stable isomer 1.A of the BGe_{12}^+ cation is given in Figure 1, whereas some of its lower-lying

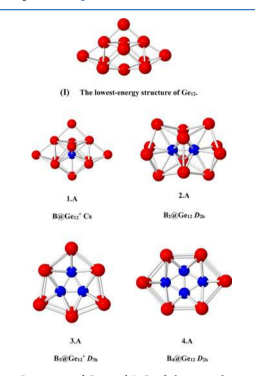


Figure 1. Geometries of Ge_{12} and $\text{B}_x \text{Ge}_{12}^{-q}$ clusters with x = 1, 2, 3, 4, and q = 0, 1. Geometry optimizations were performed using the TPSSh/6-311+G(d) and B3P86/6-311+G(d) methods.

isomers are shown in Figure S1 of the Supporting Information file. The ground state **1.A** (C_s ¹A') is revealed by both TPSSh and B3P86 calculations. **1.A** has an endohedral form in which the B⁺ ion is covered by the most stable structure of the Ge₁₂ cluster (I) displayed in Figure 1. Actually, a similar shape was identified for the isovalent BSi₁₂⁺ in which the most stable isomer Si₁₂ fully covers the B dopant.³⁶ Hence, the latter actually does not induce a significant modification on the geometry of both pure Ge₁₂ and Si₁₂ clusters.

3.2. B₂Ge₁₂. Our extensive DFT optimizations point out that a mixture of two B atoms with 12 Ge atoms induces an endohedral structure 2.A for the neutral B₂Ge₁₂. Some lowerlying isomers are displayed in Figure S1 of the Supporting Information file. Geometric characteristics of 2.A illustrate that both B atoms form a B-B dimer, which is encapsulated by a Ge_{12} host, then giving rise to a high symmetry $D_{2h} B_2 Ge_{12}$. The B-B connection of 2.A, with a bond distance of 1.73 Å, is significantly longer than that of free B_2 molecule (1.61 Å). On the one hand, formation of 2.A reduces the strength of the B_2 molecule. For the pure Ge_{12} system, a D_{2h} isomer having the same skeleton as that in 2.A is highly unstable on its potential energy surface.⁵⁶ An interesting finding is that the two B dopants contribute to stabilize an unstable D_{2h} Ge₁₂ isomer, emphasizing a special effect of a double doping of two B atoms. According to previous reports, the geometry of the B2Ge12 cluster is totally different from that of the B₂Si₁₂. As shown in Figure 2, the B_2Si_{12} is of a double ring structure in which the B_2

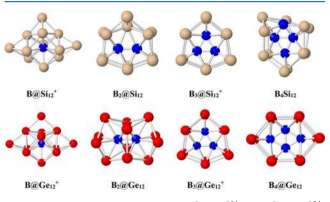


Figure 2. Summary of geometries of $[B_x Si_{12}]^{0/+}$ and $[B_x Ge_{12}]^{0/+}$ clusters, with x = 1, 2, 3, 4.

moiety is located inside a (6×2) Si₁₂ prism, whereas the B₂Ge₁₂ is stabilized in a deltahedral shape. This result emphasizes the different behavior between Si and Ge hosts regarding the B₂ dopant.

Concerning the doubly doped Ge clusters having a cage shape, B₂Ge₁₂ appears to be the smallest system known up to now. As mentioned above, the $[Pd_2@Ge_{18}]^{4-}$ and $[Co_2@$ Ge_{16} ⁴⁻ clusters are of the deltahedral germanium clusters.¹⁷⁻¹⁹ According to our literature survey and own extensive computations, no size smaller than the current case of B₂Ge₁₂ can form endohedral structure with two dopants. The B_2Ge_{12} 2.A no doubt represents the smallest single deltahedron of germanium cluster which can endohedrally encapsulate two dopants. The B₂ moiety also exerts a different influence on the isovalence Si_{12} cluster which is of a C_s cage. Indeed, the most stable B₂Si₁₂ is a distorted hexagonal prism in which the B₂ unit is covered by a (2×6) Si₁₂ prism.^{36,37} Hence, the B₂ dopant brings in strong modification on the geometry of both Ge₁₂ and Si₁₂ bare clusters. The main difference in both doped clusters is that because of its larger size, the Ge₁₂ cage has more available space to fully cover the B₂ unit.

3.3. $B_3Ge_{12}^+$. Following BGe_{12}^+ and B_2Ge_{12} , the geometry of the triply boron-doped $B_3Ge_{12}^+$ with a positive charge is considered. Lower-lying isomers of the $B_3Ge_{12}^+$ cation identified by computations using both TPSSh and B3P86 functionals and the 6-311+G(d) basis set are displayed in Figure S2 of the Supporting Information file. Accordingly, isomers containing a B_3 cycle are highly stable, and among

them, the structure having the highest symmetry **3.A** $(D_{3h}^{-1}A'_1)$ is found to be its ground state. **3.A** contains indeed a D_{3h} B₃ cycle which is placed inside a (6×2) Ge₁₂ hexagonal prism, and finally establishes a D_{3h} tube. While the triatomic B_3 cycle of 3.A is actually similar to the free B_3 cycle, the Ge_{12} counterpart is totally different from its pure ground state (C_s structure (I), Figure 1). Therefore, the B_3 cycle again tends to transform the geometry of Ge_{12} from a C_s cage to a D_{3h} hexagonal prism in the positively charged and singlet state. The $B_3Ge_{12}^+$ cation presents for the first time a hexagonal prism, or a double hexagonal ring tube, with three endohedral dopants for a germanium cluster, and it appears to be the smallest one of the class M3@Ge12. Recent reports also pointed out that a B_3 cycle exerts a similar effect on the Si₁₂ host in which a D_{3h} hexagonal prism was also found as the most stable isomer for the singlet $B_3Si_{12}^+$ cation.³⁶ It can be concluded that the cyclic B₃ dopant consistently induces and stabilizes a tubular structure for both Si_{12} and Ge_{12} in their cationic singlet state.

3.4. B_4Ge_{12} . The geometry of the B_4Ge_{12} cluster presents us with the most interesting finding of the present work. Lowerlying isomers of the neutral B_4Ge_{12} are displayed in Figure S3 of the Supporting Information file. Similar to the $B_3Ge_{12}^+$, geometries of the B_4Ge_{12} isomers emphasize a remarkable result that the four B atoms connect together to form a B_4 cycle inside the Ge_{12} host to produce the most stable doped cluster. Both TPSSh and B3P86 calculations point out that both 4.A (D_{2h} $^1A_{1g}$) and 4.B (C_1 1A) isomers are very close in energy. However, coupled-cluster CCSD(T)/6-311G(d) calculations indicate that 4.B actually lies ~7 kcal/mol above 4.A. Therefore, 4.A is confirmed to be the most stable B_4Ge_{12} isomer.

The geometric shape of **4.A** clearly points out that a tetraatomic D_{2h} B₄ cycle is placed inside of a (2×6) Ge₁₂ prism generating again a hexagonal prismatic shape for B₄Ge₁₂. In the pure form, a Ge₁₂ tube does not exist as a local minimum. In the cationic form, it is much higher in energy in Ge₁₂²⁺ (~30 kcal/mol) and Ge₁₂⁴⁺ (~13 kcal/mol) than their corresponding global minimum Ge₁₂ cages (such as I, Figure 1). Although the B₄ unit of **4.A** exhibits the same shape as that of the free B₄, its B–B bond length of ~1.7 Å is actually longer than that of ~1.5 Å in pure B₄.³⁷ As far as we are aware, the appearance of the **4.A** structure presents us with a particular case of doped germanium clusters, in which an encapsulation of four dopants in a double hexagonal ring tube is found for the first time.

Although a hexagonal prism form is a general structural motif for both $B_3Ge_{12}^+$ and $B_3Si_{12}^+$ clusters, this kind of structure is not found for the B₄Si₁₂ cluster. Actually, the most stable structure of B₄Si₁₂, 4.Si.A isomer, can be considered as a broken version of the B₄Ge₁₂ hexagonal prism. The lower-lying isomers of the B₄Si₁₂ cluster identified by the TPSSh/6-311+g(d) level of theory are given in Figure S4 (Supporting Information file). Accordingly, the **4.Si.A** isomer is found to be the lowest-energy structure, whereas the isomer 4.Si.I, which has the same shape as that of B_4Ge_{12} , is ~12 kcal/mol higher than 4.Si.A. Within 4.Si.A, boron atoms also form a B_4 cycle, but it is not planar, and one B atom replaces a Si site of the (6 \times 2) Si₁₂ hexagonal prism. This result emphasizes that the Si₁₂ double ring is not large enough to contain a B_4 ring, whereas the Ge₁₂ counterpart fits well in this geometric condition. For a complete picture, the geometries of $B_x Si_{12}^{+/0}$ and $B_x Ge_{12}^{+/0}$ are shown in Figure 2.

In summary, geometric features of B_2Ge_{12} , $B_3Ge_{12}^+$, and B_4Ge_{12} clusters indicate the ability of small boron clusters to influence the geometry of the Ge_{12} moiety. While one B atom does not give a significant impact on the Ge_{12} cage, a B_2 dimer converts a C_s Ge_{12} into a higher symmetry D_{2h} shape and produces the smallest deltahedral cage for germanium clusters. The most interesting finding is that the appearance of the **3.A** and especially **4.A** clusters illustrate that both B_3 and B_4 units can be used as dopant units that strongly stabilize a Ge_{12} host into a higher symmetry hexagonal prismatic tube, forcing an adaption of the host to their point group in a suitable charge state.

Concerning the formation of prismatic germanium structures, both B_3 and B_4 units show the importance of geometry requirement. As mentioned in Introduction, three Pd or Au dopants form a Pd₃ or Au₃ triatomic cycle which connect two Ge9 units and then generate the sandwich-type Pd3Ge18 and Au_3Ge_{18} clusters. In contrast, an endohedral structure is now established with a B_3 unit.^{14–16} With bond lengths of 2.9 and 3.0 Å, both Pd₃ and Au₃ cycles are apparently too large to fit inside a Ge_{18} cage, whereas both B_3 and B_4 units, with a B-B bond length of ~ 1.7 Å, are more suitable for endohedral insertion. In addition, a prismatic motif was previously identified for germanium clusters doped by transition metals including a pentagonal prism for M@Ge10^q and a hexagonal prism for $M @ Ge_{12}^{q}$ clusters.^{21,26} Within each of these structures, the metal dopant was found to be located inside a (2×5) Ge₁₀ or a (2×6) Ge₁₂ prism and thereby forming M@ Ge10^{*q*} and M@Ge12^{*q*} endohedral prismatic clusters, respectively. The emergence of 3.A and 4.A structures in which both B_3 and B_4 cycles are located inside a (2×6) Ge₁₂ prism, thus provides us with a novel approach to construct endohedral prismatic structures. Generally, tubular clusters can also be established by doping a planar cluster that has a suitable size.

The high symmetry form of both $B_3Ge_{12}^+$ and B_4Ge_{12} clusters is highly suitable for the assembly materials. Upon extension of both $B_3Ge_{12}^+$ and B_4Ge_{12} hexagonal prisms along the *z*-axis, a novel type of nanotube is established as shown in Figure 3. Accordingly, the final forms of $[B_3Ge_{12}^+]_n$ and $[B_4Ge_{12}]_n$ nanotubes exhibit smaller $(B_3)_n$ and $(B_4)_n$ tubes covered inside the $(Ge_{12})_n$ tube. Because of the effect of $(B_3)_n$ and $(B_4)_n$ tubes, the $[B_3Ge_{12}^+]_n$ and $[B_4Ge_{12}]_n$ tubes can thus

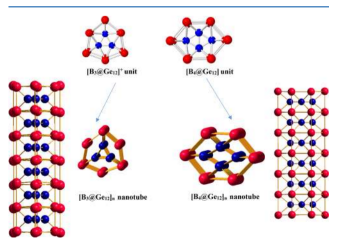


Figure 3. Formation of $[B_3 @ Ge_{12}]_n$ and $[B_4 @ Ge_{12}]_n$ assembly nanotube from $[B_3 @ Ge_{12}]^+$ and $[B_4 @ Ge_{12}]$ units.

be built up, whereas the only-Ge-based tube could not be made yet.

For a quantitative evaluation on the stability of the $[B_xGe_{12}]^{0/4}$ clusters, their binding energies are calculated and given in Table 1. Accordingly, the B_2Ge_{12} deltahedral structure and B_4Ge_{12} hexagonal prism have significantly larger E_b than that of the BGe_{12}^+ and $B_3Ge_{12}^+$ clusters. These neutral mixed clusters are in higher stability than the cations. The embedding energies, which are the energy differences between the doped cluster and the separated fragments $Ge_{12} + B_x^{0/4}$, are also performed and given in Table 1. The $B_3Ge_{12}^+$ hexagonal structure has the largest embedding energy, indicating that of the B_x dopants, introduction a B_3^+ unit into the Ge_{12} host is the most favored approach.

In attempts to understand how a B_x unit can exert such a large influence on the geometry of a Ge_{12} host, their bonding phenomenon is now explored. Investigation of electron densities of the $B_xGe_{12}^{-q}$ clusters using the ELF points out some interesting perspectives in terms of chemical bonding. For the singly doped BGe_{12}^{+} **1.A**, as shown in Figure 2, localization domains associated with B–Ge bonds are observed. However, at the bifurcation value of ELF = 0.65, in the region between B and Ge_{12} host one basin appears covering B atoms indicating the importance of the electrostatic interaction. Hence, the B⁺ dopant establishes a polarized bond with Ge_{12} host. This observation is supported by NBO atomic charge calculated for the B atom which bears a value of -2.1 electron, illustrating that the Ge_{12} host actually undergoes a great charge transfer to the B center.

Concerning the smallest germanium deltahedron 2.A of B₂Ge₁₂, two B atoms give rise to a delocalized bond with the Ge₁₂ deltahedral skeleton, as indicated by ELF maps. The ELF map of 2.A (Figure 4) shows the appearance of localization domains in regions between the B atoms and Ge₁₂ host. This observation emphasizes that B dopants form delocalized bonds with the Ge₁₂ counterpart. Indeed, both B dopants in 2.A actually carry a negative NBO charge of -1.8 electron. Similar to BGe_{12}^+ , Ge_{12} transfers its charge to the B_2 unit and thereby enhances its stability through the electrostatic interaction. From this view, 2.A can be considered as a $[B_2^{\delta} - Ge_{12}^{\delta}]$ donor-acceptor complex. Another important feature in B₂Ge₁₂ is that both boron atoms in 2.A connect through a covalence bond rather than existing as two separated atoms. The ELF map of 2.A (Figure 2) also shows a V(B,B) basin located between two B atoms indicating the formation of a B-B valence bond. In addition, the MBO performed for 2.A show a value of \sim 1.0 for this B–B connection. Although this value is much lower than that of the free B_2 molecule (1.9), it suggests that the B–B moiety of B_2Ge_{12} 2.A can be regarded as a B–B single bond.

Regarding the $B_3Ge_{12}^+$ cation 3.A, a novel perspective on the B_3 moiety is now revealed by an ELF examination. With the appearance of a V(B,B,B) basin located over the B_3 cycle, three B atoms actually form a three-center valence bond rather than acting as three separated atoms. The MBO value calculated for the B_3 unit in 3.A has a value of 0.44 indicating the existence of a three-center bond. In comparison to the bare $B_3^{+/-}$ clusters, the MBO value performed for these cycles amount to 1.1/0.97, being a much greater value than those of B_3 moiety of $B_3Ge_{12}^+$. Hence, according to the usual meaning of bond order, interaction with Ge_{12} to establish $B_3Ge_{12}^+$ reduces the strength of the B_3 cycle inside the Ge tube.

	BGe_{12}^+	B_2Ge_{12}	$B_{3}Ge_{12}^{+}$	B_4Ge_{12}	B_2	B ₃ ^{-/+}	B_4
E_{b}^{b}	4.00/3.87	4.52/4.45	4.17/4.10	4.58/4.57			
$E_{\rm D}^{\ c}$	7.71/7.55	7.60/7.50	10.75/10.93	6.38/6.58			
qNBO(B)	-2.10	-1.8	-1.45	-1.30			
d (B-B)		1.73	1.76	1.70/175	1.62	1.55/1.57	1.52
MBO _{BB}		0.91	0.45	0.61/0.52	1.95	2.0/1.64	1.41
MBO_{B_x}			0.44	0.52		1.1/0.97	0.58

Table 1. Binding Energ	r (E _b , eV) and	d Dissociation Energy ($(E_{\rm D}, {\rm eV})$ of ${\rm B}_x {\rm Ge}_{12}^q$ Clusters"
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^{*a*}NBO charge, bonding length of B–B connection (Å), MBO B–B connection (MBO_{BB}), MBO B₃ and B₄ cycle (MBO_{B₂}). ^{*b*}E_b values were calculated by the TPSSh/6-311+G(d) (left), and B3P86/6-311+G(d) (right). ^{*c*}This value was performed for the process: $B_xGe_{12}^q \rightarrow B_x^q + Ge_{12}$. In the reversed direction, these values correspond to the embedding energies.

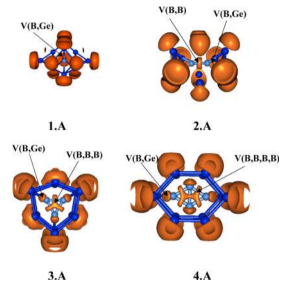


Figure 4. ELF iso-surfaces plotted at the bifurcation value ELF = 0.70 for 1.A, 2.A, 3.A, and 4.A structures. The electron density was obtained by the TPSSh/6-311+G(d) computations.

It is meaningful to consider the connection between both the B₃ cycle and Ge₁₂ hexagonal prism. The B₃ moiety interacts with, and thereby form delocalized bonds with the Ge12 prism according to the ELF result. Localization domains shown in Figure 4, which are located in region between the B_3 and Ge_{12} , illustrate a delocalized bonding character. These localization domains are closer to the B centers illustrating the fact that electrons tend to be concentrated around the B₃ unit. On the other hand, NBO atomic charges of 3.A agree with the ELF picture on the polar character of the bond. Although the $B_3Ge_{12}^+$ has a formal positive charge, each of three B atoms bears a negative net charge of -1.5 electron. This result confirms a polar bonding between the B3 and Ge12 moieties arising from a great charge transfer from Ge12 to B3. The $B_3Ge_{12}^{+}$ cluster can again be regarded as a $[B_3^{\delta-} Ge_{12}^{\delta+}]$ donor-acceptor complex.

A similar characteristic is also identified for B_4Ge_{12} **4.A** in which four B atoms prefer to connect together and act as a B_4 block. As indicated in Table 1, the MBO values performed for B_4 unit of **4.A** and free B_4 are nearly identical, being equal to ~0.5. The ELF map of **4.A** (Figure 4) gives a similar observation. In fact the ELF map of **4.A** reveals the V(B,B,B,B) basin populated over four B atoms illustrating the formation of a four-center bond. Similar to the cases of $B_3Ge_{12}^+$ and B_2Ge_{12} , the B_4 cycle in **4.A** also brings in a strongly delocalized bonding with the Ge_{12} moiety, in constructing the hexagonal

tubular prism. Actually, localization domains appeared in the region between both B₄ unit and Ge₁₂ prism is associated with the emergence of delocalized bond between the B₄ dopant with the Ge₁₂ host. In terms of charge transfer, the Ge₁₂ host again supplies its electrons to the B₄ moiety, in such a way that each B atom bears a negative net charge of -1.3 electron, emphasizing again the importance of induced electrostatic interaction within an apparent $[B_4^{\delta-} Ge_{12}^{\delta+}]$ donor-acceptor complex.

In summary, the above analysis gives an emphasis that the small B_2 , B_3 , and B_4 clusters can be used as efficient dopants to turn Ge hosts into high symmetry clusters. Boron atoms introduced into a Ge₁₂ host maintain both geometric and bonding characteristics as in free B_2 , B_3 , and B_4 clusters according to calculated ELF and MBO results. These small B clusters connect with Ge₁₂ host through delocalized bonds. The Ge₁₂ host carries out a great charge transfer to B_x and thereby produces a kind of $[B_x^{\delta-} \text{Ge}_{12}^{\delta+}]$ donor–acceptor complex. The B_x units can be considered as negatively charged islands surrounded in positively charged Ge₁₂ tubes.

3.5. Molecular Orbital Interactions between Ge12 and B_x Units. The special geometries of B_2Ge_{12} , $B_3Ge_{12}^+$, and B_4Ge_{12} clusters deserve a further analysis in order to more deeply understand the stabilizing interactions as well as the inherent electronic requirements. Orbital interaction diagrams between B_x dopants and their respective Ge_{12} skeletons are constructed and displayed in Figures 5-7. The orbital shapes of 2.A, 3.A, and 4.A structures associating to orbital interaction between B_r and Ge_{12} are given in Figures S6–S8 of Supporting Information file. For the smallest germanium deltahedron, orbital interactions of the B_2 dimer with the deltahedral Ge_{12} emphasize an interesting perspective on its stability. As displayed in Figure 5, the doubly occupied molecular orbitals (MOs) of the Ge_{12} host involving the $5a_g$ and $4a_g$ MOs enjoy stabilizing interactions with the $2\pi_u$ of B_2 and subsequently establish the fully occupied $5a_{g}$, $6a_{g}$, and $7a_{g}$ levels for B_2Ge_{12} 2.A.

Similarly, the $2b_{2g}$ MO of B_2Ge_{12} **2.A** is constructed by combination of the $2b_{2g}$ level of Ge_{12} and $2\pi_g$ of B_2 , whereas the $1\pi_g$ of B_2 enjoys an interaction with the $3b_{1g}$ eigenstate of Ge_{12} host and generate the $2b_{1g}$ MO of B_4Ge_{12} , which is occupied by two electrons. The $2\sigma_g$ vacant level of B_2 is an orbital overlap with $4b_{2u}$ eigenstate of Ge_{12} , which is occupied by two electrons, and produces subsequently the doubly occupied $3b_{2u}$ MO of B_2Ge_{12} . These interactions are associated with an electron donation from Ge_{12} to B_2 and give rise to an enhanced stability for the resulting B_2Ge_{12} . This finding is in agreement with NBO calculations discussed above.

Orbital interaction diagrams also reveal another special bonding feature of the B_2Ge_{12} deltahedron. The B_2 highest

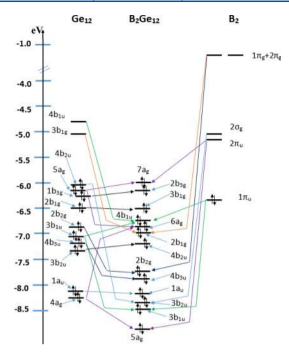


Figure 5. Orbital interaction diagram of B_2 and Ge_{12} in D_{2h} symmetry forming the **2.A** structure. The eigenvalues were obtained by TPSSh/ 6-311+G(d) calculations.

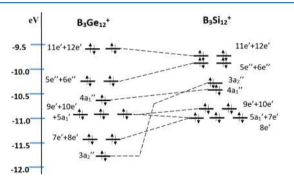


Figure 6. Quantitative correlation diagram between $B_3Ge_{12}^+$ and $B_3Si_{12}^+$ clusters. The eigenvalues were obtained by the TPSSh/6-311+G(d) calculations.

occupied MO (HOMO) is involved in an interaction with the fully occupied $3b_{1u}$ and empty $4b_{1u}$ MOs of Ge_{12} , and subsequently induce two fully occupied MOs, namely, $3b_{1u}$ and $4b_{1u}$, for D_{2h} B₂Ge₁₂ **2.A.** As a result, the stability of B₂Ge₁₂ is also contributed by electron backdonation from the B₂ unit.

Regarding the $B_3Ge_{12}^+$ hexagonal prism 3.A, an orbital interaction diagram between the D_{3h} B_3^+ and Ge_{12} skeleton is constructed and shown in Figure S5 of the Supporting Information file. Because of the fact that both isovalent $B_3Ge_{12}^+$ and $B_3Si_{12}^+$ clusters exhibit the same type of geometry,³⁶ it is meaningful to comparatively explore their electronic structure, and identify the stabilizing effect of the B_3 dopant. Empty delocalized MOs of B_3^+ gain electrons from the Ge_{12} moiety through orbital overlap, and subsequently give rise a strong stability for the $B_3Ge_{12}^+$ cluster. Actually, six empty levels of B_3^+ including lowest unoccupied

Actually, six empty levels of B_3^+ including lowest unoccupied MO (LUMO) (a'_1) , LUMO + 1,1' (e') and LUMO + 2,2' (e'') enjoy stabilizing interactions with LUMO and LUMO + 1,1' of Ge_{12} , and consequently induce thermodynamic stability for the resulting $B_3Ge_{12}^+$ hexagonal prism. Quite similar to $B_3Si_{12}^+$, the

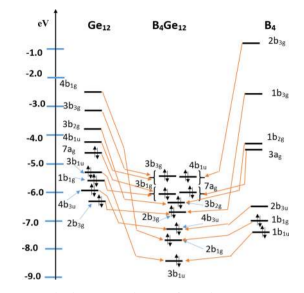


Figure 7. Orbital interaction diagram of B_4 and Ge_{12} in D_{2h} symmetry forming **4.A** structure. The eigenvalues were obtained by TPSSh/6-311+G(d) calculations.

stability of $B_3Ge_{12}^+$ is enhanced by the interaction of bonding MOs of the B_3^+ unit with Ge_{12} . As displayed in Figure S5 (Supporting Information), three bonding $1a_2''$ and 2e' + 3e' levels of B_3^+ are actually involved in an orbital overlap with the HOMO - 1 (a_2'') and HOMO - 4,4' (e') of D_{3h} Ge_{12} . These interactions subsequently generate the $3a_2''$ and 9e' + 10e' eigenstates and thereby give rise to the stable $B_3Ge_{12}^+$ cluster in the high D_{3h} symmetry. Figure 4 displays a qualitative correlation between both $B_3Si_{12}^+$ and $B_3Ge_{12}^+$ clusters in terms of eigenstates. Hence, through orbital interactions with Ge_{12} and Si_{12} tubular skeletons, both empty and occupied MOs of the B_3^+ cycle are strongly stabilized and thereby produce the stabilized D_{3h} $B_3Ge_{12}^+$ and $B_3Si_{12}^+$ tubes.

Orbital interaction diagram of both B_4 and Ge_{12} units in D_{2h} symmetry is now explored to understand how a B_4 cycle actually stabilizes a Ge_{12} block into a D_{2h} shape. It is illustrative that both B_4Ge_{12} and $B_3Ge_{12}^+$ hexagonal prisms share a similar stabilizing mechanism in which both empty and occupied MOs of the B_4 unit are stabilized. The B_4 cycle receives an electron donation from the Ge_{12} host through the orbital overlap. The $1b_{3g}$, $3a_g$, and $2b_{3u}$ empty levels of the B_4 cycle are involved in stabilizing interactions with the doubly occupied $2b_{3g}$, $7a_g$, and $4b_{3u}$ MOs of Ge_{12} , and subsequently produce the $7a_g$, $2b_{3g}$, and $4b_{3u}$ eigenstates, which are fulfilled by 6 electrons, of the D_{2h} B_4Ge_{12} tubular structure **4.A**. These interactions emphasize that a fulfilment of empty MOs of B_4 plays an essential factor for the stabilization of the binary B_4Ge_{12} cluster in a D_{2h} shape.

Orbital interactions also indicate another important contributor to the stabilizing mechanism where empty levels of the D_{2h} Ge₁₂ tube are important. As shown in Figure 7, orbital overlaps of the $4b_{1g}$, $3b_{2g}$, and $3b_{1u}$ MOs of Ge₁₂ with the $1b_{1g}$, $1b_{2g}$ and $1b_{1u}$ MOs of B₄ tend to stabilize the B₄Ge₁₂ tubular cluster. These interactions give rise to the doubly occupied $3b_{2g}$, $2b_{1g}$, and $3b_{1u}$ MOs of r B₄Ge₁₂. Therefore, occupancy of the empty levels of the Ge₁₂ skeleton as well as reinforcement of bonding MOs of the B₄ cycle constitute important contributors to the thermodynamic stability of the B₄Ge₁₂ hexagonal prism **4.A**.

The abovementioned analysis on bonding trends points out a general perspective on an electronic requirement for the

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formation of the tubular geometry. Establishment of delocalized bonds between the B_x^{q} dopant and the Ge_{12} skeletons ends up in an enhanced stability for the doped $B_xGe_{12}^{q}$ clusters. The transition metal-doped MGe_{12}^{q} clusters also share such a stabilization mechanism. Depending on the dopant characteristics, MGe_{12}^{q} structures actually tend to favor a puckered hexagonal prism, bicapped pentagonal prism, or icosahedral geometry. Investigation on the correlationship between geometry and bonding suggests that a maximization of electron density overlaps between the Ge_{12} cage and metal dopant is a driving force for formation of MGe_{12}^{q} in different geometries.⁵⁸ Although each B_x dopant basically generates a different geometry with a Ge_{12} host, they attain in all cases a stabilized electronic structure which is contributed by highly delocalized bonds made through strong orbital interactions.

A general principle to design tubular clusters can thus be proposed upon comparison to other tubular clusters. The M_2Si_{12} hexagonal prism,⁵⁹ Mn_2Si_{15} triple ring,⁶⁰ M_2B_{14} double ¹ and the Rh_2B_{18} teetotum⁶² all have the most stable form ring, in tubular geometry and such an enhanced stability is contributed by two main effects. The first effect is that both antibonding and bonding MOs of M2 are stabilized through orbital interactions with delocalized eigenstates of the corresponding tube. The second factor is that the vacant eigenstate of the Si12, Si15, B14, and B18 tubes becomes occupied in the resulting $M_2 Si_{12}\text{, }Mn_2 Si_{15}\text{, }M_2 B_{14}\text{, and }Rh_2 B_{18}$ structure, respectively, and this also contributes to a stabilization of the latter. It is clear that the M_2 and B_x dopants induce the delocalized bond with either Ge, Si, or B hosts to generate the tubular frames. Hence, an electronic principle for formation of tubular clusters is that the dopants are able to generate delocalized bonds with their host in a tubular shape, and subsequently attain a high-symmetry stable configuration.

3.6. Aromaticity of B₃Ge₁₂⁺ and B₄Ge₁₂ Clusters. With the appearance of delocalized bonds, it is insightful to further explore the aromatic features of both $B_3Ge_{12}^+$ and B_4Ge_{12} tubular prisms. Previous reports indicated that aromaticity constitutes a main reason for the planar shape of boron-based clusters.⁶³ Because it is rather difficult to classify the MOs of either $B_3Ge_{12}^+$ or B_4Ge_{12} into different σ and π sets, we now explore their magnetic ring current maps with respect to three different planes. The first plane contains either the B₃ or B₄ cycle, denoted as P_0 plane. The second plane is located about 1 a.u. above the P_0 plane, called P_1 plane. Finally, the third plane involves a Ge₆ hexagon and named as P₂ plane. Exploration of the ring current density projected into these planes, in particular the P_1 plane, provides us with an intrinsic aromatic character of these novel type of clusters. The total magnetic ring current maps of $B_3Ge_{12}^+$ 3.A and B_4Ge_{12} 4.A that are projected onto the P_0 , P_1 , and P_2 planes are displayed in Figure 8.

Regarding both P_0 and P_1 planes, both $B_3Ge_{12}^+$ and B_4Ge_{12} sustain diatropic ring current flows induced by an external magnetic field, thus indicating a certain aromatic character for these tubular clusters. For both free B_3 and B_4 cycles, appearance of diatropic induced current densities in both P_0 and P_1 planes shows both σ and π aromatic characters, respectively. However, because of the involvement of MOs of Ge_{12} skeletons in $B_3Ge_{12}^+$ and B_4Ge_{12} , the total ring current maps projected into P_0 and P_1 planes cannot separately be considered as σ and π contributions. Diatropic current density flows are identified in the P_2 -plane indicating an aromatic

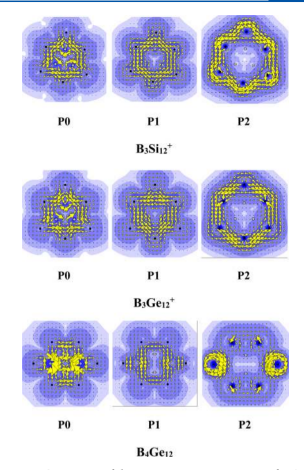


Figure 8. Comparison of the magnetic ring current maps of $B_3Si_{12}^+$, $B_3Ge_{12}^+$ **3.A** and B_4Ge_{12} **4.A** in different planes. P_0 is the plane containing B_3 or B_4 cycle, P_1 is the plane 1 a.u above P_0 , and P_2 is the plane containing a Ge₆ face. Ring current maps were obtained from B3P86/6-311+G(d) electron densities.

feature of the Ge₆ hexagon. It is illustrative that diatropic ring currents for both $B_3 {\rm Ge}_{12}{}^{\scriptscriptstyle +}$ and $B_4 {\rm Ge}_{12}$ are not only discernible on the B_3 and B_4 cycles but also on the hexagonal face of the Ge_{12} skeleton. This result emphasizes that the aromatic character of both tubular clusters $B_3Ge_{12}^+$ and B_4Ge_{12} is not solely contributed by the B_3 and B_4 dopants, but the Ge_{12} tubular skeleton equally has an important role. Indeed, the D_{2h} Ge_{12} skeleton and B_4 dopant of B_4Ge_{12} each separately show a strongly paratropic current density associating with an antiaromatic feature. Combination of both moieties end up with a high-symmetry stable aromatic cluster, clearly indicating that aromaticity plays a more crucial role in the design principle of tubular clusters. It is related to orbital interactions leading to formation of delocalized bonds, that are regarded, as discussed above, to be a driving force in stabilizing tubular clusters. The magnetic ring current calculations suggest that aromaticity, which is resulting from electron delocalization, constitutes another essential factor in the stabilization mechanism.

4. CONCLUDING REMARKS

In the present theoretical study, we successfully designed the boron-doped germanium $B_x Ge_{12}{}^q$ clusters with x = 1, 2, 3, and 4 and q = 0, 1. While the $B_2 Ge_{12}$ structure constitutes the smallest deltahedral germanium cluster, both $B_3 Ge_{12}^+$ and $B_4 Ge_{12}$ clusters present us, for the first time, with an endohedral tubular motif where either the B_3 or the B_4 cycle

Within D_{3h} and D_{2h} geometries, the B_3 and B_4 clusters generate delocalized bonds with the Ge_{12} framework and thereby induce an aromatic character for the resulting $B_3Ge_{12}^+$ and B_4Ge_{12} . Such a tubular aromaticity appears to greatly contribute to the high thermodynamic stability of the binary hexagonal Ge tubes. Together, both characteristics constitute a design electronic principle for making stabilized tubular clusters.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.9b05483.

Geometrical shapes of lower-lying isomers of the $[B_xGe_{12}]^{0/+}$ and B_4Si_{12} and their relative energies, the MO shapes of **2.A**, **3.A**, and **4.A**, orbital interaction diagram of $[B_3Ge_{12}]^+$, and Cartesian coordinates of the optimized ground-state structures of $[B_xGe_{12}]^{0/+}$ (PDF)

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Notes

The authors declare no competing financial interest.

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