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Theoretical Study of Icosahedral *closo*-Borane, -Alane, and -Gallane Dianions $(A_{12}H_{12}^{2-}; A = B, Al, Ga)$ with Endohedral Noble Gas Atoms (Ng = He, Ne, Ar, and Kr) and Their Lithium Salts $(Li[Ng@A_{12}H_{12}]^-$ and $Li_2[Ng@A_{12}H_{12}])$

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Geometries, energies, vibrational frequencies, and magnetic properties have been computed at the B3LYP level with the 6-31G* and 6-311+G* basis sets for a family of endohedral closo-boranes, -alanes, and -gallanes $Ng@A_{12}H_{12}^{2-}$ with noble gas atoms (Ng) located in the centers of icosahedral [B₁₂], [Al₁₂], and [Ga₁₂] clusters. The endohedral structures of most of the systems are minima lying above separated Ng + $A_{12}H_{12}^{2-}$ by 166 $(He@B_{12}H_{12}^{2-}) \ and \ 403 \ (Ne@B_{12}H_{12}^{2-}) \ kcal/mol \ for \ boranes; \ 29 \ (He@Al_{12}H_{12}^{2-}), \ 63 \ (Ne@Al_{12}H_{12}^{2-}), \ 154 \ Al_{12}^{2-}), \ Al_{12}^{2-}$ $(Ar@Al_{12}H_{12}^{2-})$, and 189 $(Kr@Al_{12}H_{12}^{2-})$ kcal/mol for alanes; and 39 $(He@Ga_{12}H_{12}^{2-})$, 71 $(Ne@Ga_{12}H_{12}^{2-})$, and 213 (Ar@Ga₁₂H₁₂²⁻) kcal/mol for gallanes. Three types of transition states are found for the exit of Ng from a cage: via an edge (TS-1), through a face (TS-2), and via a more extensive deformation through a pentagonal cage "neck" (TS-3). The most favorable exit path depends on the rigidity of the cage, the exothermicity of the dissociation, and the relationship between the size of the internal cavity of the cage and the Ng atomic radius. Ng exit via **TS-3** is preferred for $\text{He@Al}_{12}\text{H}_{12}^{2-}$, $\text{Ne@Al}_{12}\text{H}_{12}^{2-}$, $\text{He@Ga}_{12}\text{H}_{12}^{2-}$, $\text{Ne@Ga}_{12}\text{H}_{12}^{2-}$, $\text{Ar@Al}_{12}\text{H}_{12}^{2-}$, and $Kr@Al_{12}H_{12}^{2-}$. Helium exits via a cage edge (TS-1) for $He@B_{12}H_{12}^{2-}$, while for $Ne@B_{12}H_{12}^{2-}$ the neon exits via a triangular face (TS-2). Exit barriers ($H_{\text{exit}}^{\ddagger}$) are high enough (30-60 kcal/mol) for all helium clusters and for Ne@Al₁₂H₁₂²⁻ and Ne@Ga₁₂H₁₂²⁻ to ensure the kinetic stability of these systems. The barriers for $Ar@Al_{12}H_{12}^{2-}$ and $Kr@Al_{12}H_{12}^{2-}$ decrease to 10–15 kcal/mol, while $Ne@B_{12}H_{12}^{2-}$ has a very low exit barrier and is not expected to be stable kinetically. There is a linear dependence of Ng@A₁₂H₁₂²⁻ cage size on the Ng atomic radii; that is, the heavier Ng atoms "bulge" the cages. Nucleus independent chemical shifts (NICS) indicate that all three $A_{12}H_{12}^{2-}$ anions are aromatic but the alanes are the least so. A face- or edge-coordinated external Li+ cation has a moderate effect on the structure and vibrational and magnetic properties of the helium-containing clusters, i.e., Li[He@A₁₂H₁₂]⁻. In contrast, for systems with very large exothermicities of Ng exit, Li⁺ complexation promotes their dissociation. Thus, the internal atom Ne exits from the cage of $Li[Ne@B_{12}H_{12}]^-$ and the salt dissociates into Ne + LiB₁₂H₁₂ $^-$ without barrier. Systems with two Li⁺ ions located initially above opposite cage faces (Li₂[Ng@A₁₂H₁₂]) undergo complex intramolecular rearrangements leading to destruction of the icosahedral closo structures.

Introduction

The preparation and characterization of polyhedral compounds containing endohedral atoms or ions (X@A_n) not only involve the fullerenes, but also dodecahedrane $C_{20}H_{20}^2$ and metal clusters, such as $Ga_{13}^{-,3}$ Tl_{13}^{11-} , $NiIn_{10}^{10-}$, $ZnIn_{10}^{10-}$, and $Ga_{19}[C(SiMe_3)_3]_6^{4a}$ and $Si@Al_{14}^{4b}$ In addition, a number of more complex (condensed) clusters, which link small isolated molecular clusters and bulk metals, 2.5 also have been examined.6

Bombardment of cages such as fullerenes or dodecahedrane with ionized noble gas beams (Ng⁺) followed by Ng NMR chemical shift measurement is a useful tool for the investigation of cluster structure and properties. In combination with ab initio calculations, this technique has been a fruitful approach to the

study of isomerism, aromaticity, and other cluster properties. Similarly, metals and alloys have been bombarded with Xe beams to form Xe_n clusters in the metal surface layers which

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were then probed with Xe NMR spectroscopy to gain insight into the structure and properties of the metal surface layers and micrograin boundaries.⁹

Computational studies on endohedral polyhedral borane complexes include our 10a examination of He@B12H122- and He@B₁₇H₁₇²⁻ and Jemmis and Balakrishnarajan's recent calculations on B₁₂H₁₂²⁻ and B₁₄H₁₄²⁻ dianions "stuffed" with cations C⁴⁺, B³⁺, Be²⁺, Li⁺, Al³⁺, Mg²⁺, and Zn²⁺, ¹¹ as well as earlier studies on $B_6H_7^-$ and $LiB_6H_7^{10b}$ with proton and Li^+ in the center of the octahedral $B_6H_6^{2-}$ anion. Recently, 12 we reported preliminary results on a family of X@Al₁₂H₁₂²⁻ alane and X@Ga₁₂H₁₂²⁻ gallane endo clusters with inserted noble gas atoms He, Ne, Ar, and Kr, cations Li⁺, Na⁺, Be²⁺, Mg²⁺, Al⁺, and Al3+, and anions H-, F-, Cl-, Br-, O2-, etc. However, these studies only characterized local minima for endo stuctures; the potential barriers for exit of heteroatoms from the cage were not calculated and the question of the kinetic stability of the endo isomers remained open. Among the issues of interest to chemists are the character of the interaction between the heteroatom and the cage, the peculiarities of the internal atom valence states, the influence of heteroatom on the cage, the limits of endohedral-cluster viability (i.e., which atoms inserted into a cage A_n result in kinetically stable species), the mechanism of intramolecular rearrangements between endo and exo isomers in salts, and the three-dimensional aromaticity of clusters with delocalized cage bonding such as fullerenes1 and closoboranes. 10a, 13

We now report the results of density functional (DFT) calculations on icosahedral *closo*-borane and model *closo*-alane and *closo*-gallane dianions $(A_{12}H_{12}^{2-}; A = B, Al, or Ga)$ with noble gas atoms (Ng = He, Ne, Ar, and Kr) inserted in the

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center of the relatively small, dense cages (Ng@ $A_{12}H_{12}^{2-}$). We also computed the lithium salts of several of these compounds, with one or two external Li cations (Li[Ng@A₁₂H₁₂] and Li₂-[Ng@A₁₂H₁₂]). Our investigation of the energetic and kinetic stabilities of the Ng@ $A_{12}H_{12}^{2-}$ dianions reveals changes and trends in their structural and energetic parameters, vibrational frequencies, and NMR chemical shifts depending on the size of Ng and the cage, as well as the electronegativity of A. We analyze the influence of external cations on these properties and examine the differences between boranes, alanes, and gallanes. Our predictions on the structure, energetics, and spectroscopic properties of these compounds may be useful guides for the preparation and identification of new clusters. In contrast with the well-characterized dodecaborane dianion $(B_{12}H_{12}^{2-})$, ¹⁴ the corresponding parent alane $(Al_{12}H_{12}^{2-})$ and gallane (Ga₁₂H₁₂²⁻) have not been reported to date. Nevertheless, we expect that our predictions should be applicable to recently synthesized icosahedral alane $(Al_{12}R_{12}^{2-})^{15}$ and gallane $(Ga_{12}R_{10}^{2-})^{16}$ anions which have bulky substituents (R = Bu, N(SiH₃)₂, C₁₃H₉, etc). Our earlier DFT calculations on fluorosubstituted boranes, B₁₂H_iF_{12-i}, ^{13a} and the substituted gallane series, 17 Ga₁₉H₆ ${}^{-}$ -Ga₁₉(CH₃)₆ ${}^{-}$ -Ga₁₉[C(SiH₃)₃]₆ ${}^{-}$, revealed that the skeleton geometry is relatively insensitive to the external substituent.

Computational Methods

Similar to our previous computational studies, 12 all geometries (see Figure 1) were optimized using the hybrid density functional B3LYP method with the standard 6-31G* basis set; vibrational frequencies were computed at the same B3LYP/6-31G* level of theory. The borane anions were then reoptimized with the more flexible 6-311+G** basis set while the alanes and all salts were refined by computing B3LYP/6-311+G** single points, i.e., B3LYP/6-311+G**//B3LYP/6-31G*. Gallanes were only computed at B3LYP/6-31G*. Optimized B $_{12}$ H $_{12}$ ²⁻ and Al $_{12}$ H $_{12}$ ²⁻ $_{12}$ R(A-A) anion bond lengths (Table 1) reproduce X-ray $_{13}$ R(B-B) and $_{13}$ R(Al-Al) values $_{14}$ R(S-1) satisfactorily. B3LYP/6-31G*

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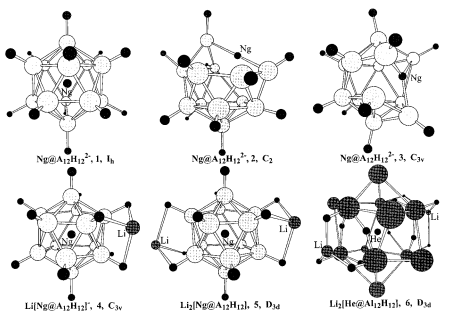


Figure 1. Various Ng@ $A_{12}H_{12}^{2-}$, Li[Ng@ $A_{12}H_{12}$]⁻, and Li₂[Ng@ $A_{12}H_{12}$] geometries.

Table 1. Calculated Bond Lengths (Å) of closo-Boranes, -Alanes, and -Gallanes Ng@A₁₂H₁₂^{2-b} (1,I_h) with Ng Atoms in the Center of the [A₁₂] Icosahedron

В				Al					Ga			
	empty ^d	Не	Ne	empty ^d	Не	Ne	Ar	Kr	empty ^d	Не	Ne	Ar
R(Ng-A)	(1.700)	1.770	1.904	(2.569)	2.597	2.635	2.724	2.758	(2.453)	2.487	2.546	2.635
R(A-A)	1.787	1.861	2.002	2.703	2.730	2.770	2.864	2.900	2.579	2.614	2.677	2.823
R(A-H)	1.202	1.202	1.202	1.605	1.609	1.610	1.613	1.616	1.568	1.569	1.571	1.584

^a At B3LYP/6-311+G** for boranes and B3LYP/6-31G* for alanes and gallanes. The basis set extension from 6-31G* to 6-311+G* does not effect the borane results significantly. ${}^{b}A = B$, Al, Ga. ${}^{c}Ng =$ "empty", He, Ne, Ar, Kr. ${}^{d}Values$ for isolated $A_{12}H_{12}^{2-}$ diamions calculated at the same levels of theory. In parentheses, the distance from the icosahedron center to a vertex atom, A.

magnetic shielding constants and NMR chemical shifts were computed with the GIAO method.¹⁹ All calculations employed the Gaussian 98 program package.²⁰ Results are presented in Tables 1-5, and optimized structures are shown in Figures 1 and 3.

Results and Discussion

Geometric and Energetic Parameters. Local minima exist for most but not all endohedral icosahedral anions Ng@A₁₂H₁₂²⁻. I_h minima include He@B₁₂H₁₂²⁻ and Ne@B₁₂H₁₂²⁻ as well as the entire alane series (He-Kr). The Ng@Ga₁₂H₁₂ $^{2-}$ I_h configuration stability only includes He, Ne, and Ar.21 The remaining Ng@A₁₂H₁₂²⁻ structures have one or several degenerate imaginary frequencies; these endohedral icosahedral

(21) All Ar@Ga₁₂H₁₂²⁻ vibrational frequencies are real at the Hartree-Fock level; however, B3LYP gives one small imaginary frequency of 22i cm⁻¹. The character of this icosahedral stationary point needs to be determined by higher level calculations.

structures are not stable with respect to exit of the inserted atom

As expected, 12 all endohedral icosahedral clusters are higher in energy than the empty cage plus the Ng atom. $E_{\rm diss}$, the zeropoint vibrational energy (ZPE) corrected energies released during dissociation

$$Ng@A_{12}H_{12}^{2-} \rightarrow Ng + A_{12}H_{12}^{2-}$$

are (see Table 2) 166 (He@B₁₂H₁₂ $^{2-}$) and 403 (Ne@B₁₂H₁₂ $^{2-}$) kcal/mol for boranes, 29 (He@Al₁₂H₁₂²⁻), 63 (Ne@Al₁₂H₁₂²⁻), 154 (Ar@Al₁₂H₁₂²⁻), and 189 (Kr@Al₁₂H₁₂²⁻) kcal/mol for alanes, 39 (He@Ga₁₂H₁₂²⁻), 71 (Ne@Ga₁₂H₁₂²⁻), and 213 $(Ar@Ga_{12}H_{12}^{2-})$ kcal/mol for gallanes. While E_{diss} increases with larger noble gas atomic radii R(Ng), the trend with cage size is irregular (boranes > alanes < gallanes). This trend is in line with the cage sizes as gallanes are more compact than the alanes; the lengths of the R(Ga-Ga) edges as well as the peripheral R(Ga-H) bonds are shorter than the corresponding R(Al-Al) and R(Al-H) distances by 0.12 and 0.04 Å, respectively, both in the empty A₁₂H₁₂²⁻ dianions and in the endohedral Ng@A₁₂H₁₂²⁻ systems (Table 1). In turn, the larger size of alanes as compared to gallanes may be related to the order of electronegativities, B > Al < Ga. Al is less electronegative than Ga and forms more ionic bonds with H's, thus making the Al₁₂ cage more electron deficient and weakly bound than Ga_{12} .

The sizes of the endohedral clusters increase upon Ng encapsulation, parallel with the rise in their dissociation exothermicity. Compared with the B-B length in the empty dianion,

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Table 2. Total Energies (au), ZPE, Dissociation Energies of Ng@ $A_{12}H_{12}^{2-}$, Deformation Energies of the Cage $A_{12}H_{12}^{2-}$ due to Insertion of Ng, Relative Energies of Structures **2** and **3**, and Barriers for Exit of Ng from the Cage $A_{12}H_{12}^{2-}$ in *closo*-Boranes, -Alanes, and -Gallanes (kcal/mol)^a

		Ng@Al ₁₂ H ₁₂ ²	_	
Ng	He	Ne	Ar	Kr
$E_{ m tot}$	-2919.8036	-3045.7949	-3444.2410	-5670.3744
ZPE	60.2	59.2	57.8	56.9
$E_{ m diss}{}^b$	-29.1	-62.9	-154.3	-188.8
$E_{ m def}$	0.9	5.3	26.0	37.3
$E_{\rm rel}(2,C_2)^c$	78.8	39.3	-36.0	-59.2
$E_{\rm rel}(3,C_{3v})^c$	77.2	48.7	-0.4	-21.7
$H_{\mathrm{exit}}^{\ddagger}$	58.5	41.5	15.2	10.5

	Ng@B	12H ₁₂ ²⁻	Ng@Ga	$\frac{1}{12}H_{12}^{2-u}$
Ng	Не	Ne	Не	Ne
$E_{ m tot}$	-308.4141	-434.0679	-23058.8081	-23122.7381
ZPE	106.8	97.2	57.6	55.9
$E_{\mathrm{diss}}{}^b$	-166.0	-402.9	-38.7	-70.9
$E_{ m def}$	15.4	104.1	1.5	11.8
$E_{\rm rel}(2,C_2)^c$	31.2	-170.0		
$E_{\rm rel}(3,C_{3v})^c$	97.1	-88.8		
$H_{ m exit}^{\ddag}$	63.5	2.6	29.1^{e}	

^a Calculations of $E_{\rm tot}$ and $E_{\rm diss}$ were performed at the B3LYP/6-311+G** level for boranes, B3LYP/6-311+G**/B3LYP/6-31G* for alanes, and B3LYP/6-31G* for gallanes, while the other energies (ZPE, $E_{\rm def}$, $E_{\rm rel}$, and $H_{\rm exit}$) were calculated at the B3LYP/6-31G* level. $E_{\rm tot}$ and ZPE for empty clusters are the following: −305.7638, −2919.9334, and −23082.9584 au and 105.0, 58.0, and 54.9 kcal/mol for B₁₂H₁₂^{2−}, Al₁₂H₁₂^{2−}, and Ga₁₂H₁₂^{2−}, respectively. ^b The dissociation limit corresponds to dissociation of Ng@A₁₂H₁₂^{2−} (1) into Ng + A₁₂H₁₂^{2−}. ^c With respect to the energy of the icosahedral Ng@A₁₂H₁₂^{2−} structure 1. ^d For Ar@Ga₁₂H₁₂^{2−} $E_{\rm tot}$, ZPE, and $E_{\rm diss}$ are the following: −23610.1339 au, 58.3, and −217.8 kcal/mol, respectively. ^e Calculated at the B3LYP/SDD level.

the B–B edge in the boranes elongates by 0.08 and 0.21 Å after insertion of He and Ne, respectively. In the alanes, where the empty cage size is larger, the Al–Al edge elongations when He and Ne are inserted are only 0.03 and 0.07 Å, respectively, but this increases to 0.16 and 0.20 Å, respectively, upon insertion of Ar and Kr. The cage expansion in gallanes is similar to that in alanes. The peripheral Ng@A₁₂H₁₂^{2–} A–H bonds are also stretched but only by a few thousandths of an angstrom.

As shown in Figure 2a, the changes in Ng@ $A_{12}H_{12}^{2-}$ edge $\Delta R(A-A)$ and radial $\Delta R(Ng-A)$ distances exhibit nearly linear dependences on the noble gas atomic radius R(Ng)

$$\Delta R(\text{Ng-A}) \sim K_{\text{A}}R(\text{Ng})$$

where the coefficient K_A is largest for boranes, smallest for alanes, and is intermediate for gallanes. Dissociation energies calculated for clusters with internal He, Ne and Ar atoms show a parabolic dependence on $\Delta R(Ng-A)$ with a minor quadratic term (Figure 2b).

To estimate cage deformation energies due to Ng insertion, $E_{\rm def}(A_{12}H_{12}^{2-})$, we carried out single-point B3LYP/6-31G* calculations on the expanded empty $A_{12}H_{12}^{2-}$ dianion geometries based on the equilibrium structures of endohedral Ng@ $A_{12}H_{12}^{2-}$ clusters. These were compared with the energies of the fully relaxed $A_{12}H_{12}^{2-}$ dianions (energies given in Table 2). As expected from the results discussed above, the alane cage appears to be more "elastic" with respect to expansion than the borane cage while the gallane cage is intermediate. In the rigid borane clusters, even replacement of He with Ne leads to elongation of R(B-B) by 0.13 Å and sharp increases in $E_{\rm diss}$ (~237 kcal/mol) and in $E_{\rm def}$ (~90 kcal/mol). In contrast,

replacement of He with a much larger analogue, Kr, in the "elastic" alane cage elongates R(Al-Al) to a similar extent (0.16) Å), but the $E_{\rm diss}$ and $E_{\rm def}$ energies increase by only ~ 150 and ~36 kcal/mol, respectively. Apparently, the higher rigidity of the Ga₁₂H₁₂²⁻ dianion is responsible for the instability of the icosahedral Kr@Ga₁₂H₁₂²⁻ cluster. The dependence of the change of empty cluster total energy $[\Delta E_{tot}(A_{12}H_{12}^{2-})]$ for the boranes and alanes families on the change in their size $\Delta R(oA)$ (o is the center of the icosahedron) is described by two parabolic curves, shown in Figure 2c, the flat one for the "elastic" alanes and the steeper one for the more rigid boranes. The good fit, together with the linear dependence of Ng@A₁₂H₁₂²⁻ cluster size on R(Ng), indicates that steric repulsion is the dominant factor determining the compactness and strain energy of a cage. Since the steepest rise in R(Ng) occurs between Ne and Ar, the largest change in endohedral alane structural and energetic parameters also takes place when Ne is replaced with Ar. However, replacements of He by Ne and Ar by Kr result in smaller changes.

Clearly, cage deformation energy constitutes only a small fraction of the energy released when an internal Ng atom is expelled from a cage. $E_{\rm def}$ is small for He@Al₁₂H₁₂²⁻ and He@Ga₁₂H₁₂²⁻, moderate for their Ne analogues (5–10 kcal/mol), and becomes significant only with Ng = Ar. Rigid boranes again behave differently; for He@B₁₂H₁₂²⁻ $E_{\rm def}$ is 15 kcal/mol and increases to 104 kcal/mol for Ne@B₁₂H₁₂²⁻. Obviously, strong repulsive interactions between the noble gas atom and the cage contribute strongly to $E_{\rm diss}$.

Vibrational Frequencies. Vibrational frequencies involving Ng@A₁₂H₁₂²⁻ cage atoms and peripheral bonds may be shifted (either blue or red) compared with the corresponding frequencies of the empty A₁₂H₁₂²⁻ anions (Table 3). The frequencies usually shift monotonically with increase in Ng atomic number. For example, the lowest 1hg, 1gg, and 2hg frequencies in the alane series decrease by 50-60 cm⁻¹ in going from He to Kr; the larger Al-H stretching frequencies (4hg, 2t_{2u}, 2a_{1g}, and 4t_{1u}) decrease uniformly by ~40 cm⁻¹, and the 3h_g frequencies increase by $40\ cm^{-1}$. Only four t_{1u} frequencies are IR active due to the high I_h symmetry. Among these, the changes of most intense vibrations ($3t_{1u}$ and $4t_{1u}$) are small and the vibration with the largest frequency shift (1t_{1u}) has a low intensity. Therefore, the $2t_{1u}$ vibration, in the 925-1030 cm⁻¹ range for boranes and 340-560 cm⁻¹ for alanes and gallanes, which for the most part corresponds to motions of the inserted Ng atom and of moderate intensity, should be the most informative in IR spectroscopic studies of the endohedral clusters.

Orbital Energies, Population Analysis, and Charge Dis**tribution.** Endohedral alanes and gallane ionization potentials (IP), evaluated using Koopmans' theorem (Table 4), show only a weak dependence on Ng and are close to those of the empty $Al_{12}H_{12}^{2-}$ and $Ga_{12}H_{12}^{2-}$ dianion IPs (1.0–1.1 eV). In contrast, the Ng@B₁₂H₁₂²⁻ IPs decrease significantly from the empty $B_{12}H_{12}^{2-}$ (~1.6 eV) to Ne@B₁₂H₁₂²⁻ (~1.1 eV). The HOMO-LUMO separations decrease with increasing Ng radius and range from 8 to 9.5 eV in alanes and gallanes and 10-11.5 eV in boranes. The skeleton valence molecular orbitals (MO) energy levels and those of the atomic orbitals (AO) of the internal He and Ne atoms are well separated and do not mix. The Ar and Kr AO and the skeleton valence MO levels do approach each other. Hence, the cage bonding t_{1u} MO includes some contribution from a p AO of these noble gas atoms. For example, the Ar 3p AO and Kr 4p AO population in the alane cage t_{1n} MOs are \sim 0.3 and \sim 0.4 e, respectively. The situation is similar for the Ng@ $B_{12}H_{12}^{2-}$ clusters (the contribution of the $1s_{He}$ and $2p_{Ne}$

Table 3. Calculated^a Vibrational Frequencies v_i (cm⁻¹) of closo-Boranes, -Alanes, and -Gallanes Ng@A₁₂H₁₂^{2-b} (1, I_b) with Ng Atoms^c in the Center of the [A₁₂] Icosahedron

			В				Al				Ga	
		$empty^d$	Не	Ne	$empty^d$	Не	Ne	Ar	Kr	$empty^d$	Не	Ne
h _u	1 2	521 959	439 932	291 879	174 486	163 491	149 489	115 481	103 473	92 488	80 497	60 492
h_{g}	1 2 3 4	575 765 962 2517	573 713 912 2526	369 687 888 2526	185 328 466 1804	204 316 472 1810	204 302 486 1805	192 274 504 1790	183 269 513 1775	114 219 450 1883	128 209 464 1880	122 196 482 1861
g_{u}	1 2	747 879	612 883	328 834	307 426	294 471	271 468	219 488	204 487	200 426	188 475	163 477
$g_{\rm g}$	1 2	662 946	569 898	404 867	246 450	233 450	218 458	181 458	170 456	148 433	136 441	117 448
$\mathbf{t}_{1\mathrm{u}}$	1 2	710 (7)	574 (1) 1031 (77)	294 (0) 925 (71)	309 (5)	283 (3) 456 (10)	233 (1) 357 (6)	190 (1) 376 (10)	158 (1) 338 (10)	212 (0)	193 (0) 585 (12)	161 (0) 376 (3)
	3	1089 (64) 2535 (1149)	1343 (15) 2544 (1225)	1003 (53) 2542 (1359)	578 (288) 1810 (1656)	580 (287) 1816 (1700)	579 (303) 1812 (1742)	565 (294) 1798 (1884)	550 (264) 1784 (1964)	567 (218) 1883 (1656)	562 (227) 1880 (1679)	569 (254) 1863 (1719)
t_{2u}	1 2	765 2510	743 2519	747 2520	253 1800	255 1806	266 1802	267 1788	270 1774	155 1876	156 1872	167 1857
t_{1g}	1	961	941	899	420	425	422	414	407	387	389	373
$a_{\rm g}$	1 2	744 2572	704 2583	659 2580	299 1834	297 1841	297 1836	273 1823	274 1811	208 1910	204 1907	198 1890

^a At the B3LYP/6-31G* level. Intensities of IR-active t_{1u} vibrations (km/mol) are given in parentheses. The Ar@B₁₂H₁₂²⁻ and Ng@Ga₁₂H₁₂²⁻ clusters with Ng = Ar and Kr possess one or several degenerate imaginary frequencies. b A = B, Al, Ga. Rg = "empty", He, Ne, Ar, Kr. Data for the isolated $A_{12}H_{12}^{2-}$ species at the same level of theory.

Table 4. Orbital Energies (au) of A₁₂H₁₂²⁻ MOs and Effective Atomic Charges (e) of A₁₂H₁₂²⁻ and Ng@A₁₂H₁₂^{2-b} Calculated at the HF/ 6-31G*//B3LYP/6-31G* Level

		В			Al					
	empty ^a	Не	Ne	empty ^a	Не	Ne	Ar	Kr		
MO										
a_{g}		-0.956	-1.819		-0.825	-1.835	-1.204	-1.071		
t_{1u}			-0.791			-0.761	-0.565	-0.526		
a_{g}	-0.706	-0.493	-0.563	-0.466	-0.433	-0.446	-0.419	-0.405		
t_{1u}	-0.523	-0.517	-0.421	-0.382	-0.382	-0.367	-0.332	-0.313		
h_{g}	-0.327	-0.325	-0.327	-0.287	-0.288	-0.289	-0.293	-0.295		
t_{2u}	-0.217	-0.230	-0.253	-0.254	-0.256	-0.258	-0264.	-0.266		
a_{g}	-0.301	-0.184	-0.236	-0.232	-0.213	-0.221	-0.210	-0.201		
t_{1u}	-0.202	-0.212	-0.145	-0.193	-0.197	-0.185	-0.156	-0.131		
h_{g}	-0.105	-0.111	-0.120	-0.123	-0.125	-0.127	-0.138	-0.144		
g_u	-0.056	-0.049	-0.038	-0.037	-0.037	-0.037	-0.038	-0.038		
g* LUMO	0.501	0.482	0.441	0.276	0.274	0.270	0.262	0.259		
Z										
Ng		0.07	0.04		0.02	0.06	0.32	0.31		
A	-0.05	-0.05	-0.02	-0.00	-0.00	-0.00	-0.02	-0.02		
Н	-0.12	-0.12	-0.15	-0.16	-0.17	-0.17	-0.17	-0.18		

^a Data for the isolated $A_{12}H_{12}^{2-}$ species at the same level of theory. ^b A = B, Al. Ng = "empty", He, Ne, Ar, Kr.

AO into the a_g and t_{1u} MO are 0.3 and 0.2 e, respectively) since the B₁₂H₁₂²⁻ dianion valence orbital levels lie much lower than those in Al₁₂H₁₂²⁻. The a_g and t_{1u} MOs show the strongest dependence of orbital energetic levels on the nature of the internal Ng atoms.

Both Mulliken and natural bond orbital (NBO) population analyses on the alanes and gallanes show nearly zero charges on He and Ne. Only a small charge transfer from their heavier analogues (which increases slightly from Ar to Kr but does not exceed 0.1 e) occurs to the A₁₂H₁₂ cage. A similarly small charge transfer takes place in boranes where the He and Ne charges are +0.05 and +0.10 e. The overlap populations Q(Ng-A) are negative and decrease in alanes from He@Al₁₂H₁₂²⁻ (-0.02 e) to Kr@Al₁₂H₁₂²⁻ (-0.15 e). Positive Q(A-A) overlap populations decrease slowly when the cage strain increases, while Q(A-H) populations for the peripheral bonds are rather insensitive to the nature of Ng. These results once more indicate that steric repulsion between Ng and cage is the dominant force in endohedral clusters with noble gases.

Magnetic Shielding Constants and NMR Chemical Shifts. Depending on the size, shape, and charge of the cluster, ³He chemical shifts inside fullerene cages are -8 to -48 ppm compared to free He.1 The negative sign of the 3He NMR chemical shift δ (He) is due to the shielding effect induced by the cyclic electron delocalization in the π -electronic system (i.e., aromaticity) of the C_n cluster. In the icosahedral Ng@A₁₂H₁₂²⁻ clusters, the internal Ng and cage atoms are closer together than in the larger endo-fullerenes. closo-Boranes are known to exhibit

Table 5. Magnetic Shielding Constants σ and Nucleus Independent Chemical Shifts (NICS) of $A_{12}H_{12}^{2-}$ Closo Anions (A = B, Al, and Ga) and Ng@A₁₂H₁₂²⁻ Endo Clusters (ppm)^a

		σ		NICS^b				
anion	A	Ng	Н	(c)	(0)	(0.5)	(1.0)	
$B_{12}H_{12}^{2-}$ $He@B_{12}H_{12}^{2-}$	127.0 122.4	82.0	30.8 30.8	25.9	-48.4 -47.0	-30.7 -33.6	-9.5 -9.4	
Ne@ $B_{12}H_{12}^{2-}$ A $l_{12}H_{12}^{2-}$ He@ $Al_{12}H_{12}^{2-}$	110.1 575.2 572.9	38.1 78.2	29.7 29.8 29.8	-20.5	-40.2 -28.1 -27.6	-25.7 -21.9 -20.7	-8.0 -11.8 -10.7	
Ne@Al ₁₂ H ₁₂ ²⁻ Ar@Al ₁₂ H ₁₂ ²⁻	575.6 588.2	400.2 752.5	29.8 29.7 29.5		-27.6 -26.5 -23.7	-20.7 -19.9 -18.1	-10.7 -10.3 -9.3	
$Kr@Al_{12}H_{12}^{2-}$ $Ga_{12}H_{12}^{2-}$	602.9 2191.0	2259.3	29.3 30.0	-25.5	-21.6 -36.3	-16.9 -26.8	-8.7 -13.0	
${ m He@Ga_{12}H_{12}^{2-}} \\ { m Ne@Ga_{12}H_{12}^{2-}} \\ { m Ar@Ga_{12}H_{12}^{2-}} \\ \\$	2199.7 2236.1 2313.0	82.9 332.0 581.4	30.0 29.8 29.2		-35.4 -32.9 -27.1	-26.0 -24.4 -20.7	-12.7 -12.0 -10.3	

^a Calculated at the GIAO/B3LYP/6-31G* level. Absolute shieldings (o) are presented. Calculations at the same level of theory give the following magnetic shielding constants (in ppm) for AlH_4^- ($\sigma(Al) = 541$), GaH_4^- ($\sigma(Ga) = 2008$), and B_2H_6 ($\sigma(B) = 28.1$), for which the accepted experimental NMR chemical shifts are $\delta(Al) = 100-103$, $\delta(Ga) = 682$, and $\delta(B) = 16.6$ ppm. Ng isotropic shielding constants are 59.8, 552.0, 1237.0, and 3236.0 ppm for He, Ne, Ar, and Kr, respectively. b The NICS values were calculated at the centers of the icosahedron (c) and of a face (0) as well as at the 0.5 and 1.0 Å distances above a face center ((0.5) and (1.0), respectively).

three-dimensional aromaticity due to the multicenter skeletal bonding. 10a,13 The computed NMR chemical shifts, δ (He), for He@A₁₂H₁₂²⁻ clusters depend on the size and nature of the $A_{12}H_{12}^{2-}$ cage. Nucleus independent chemical shifts (NICS)²² were also computed as an aromaticity criteria and are included in Table 5. NICS were calculated in the centers of the cages, and of the faces, as well as 0.5 and 1.0 Å above the face centers of both the empty $A_{12}H_{12}^{2-}$ dianions and $He@A_{12}H_{12}^{2-}$ clusters.

Akin to the fullerenes, He@A₁₂H₁₂²⁻ icosahedron He chemical shifts (δ (He)) are negative with respect to free He (i.e., shielding in all three cages) but are 2-2.5 times larger compared with $\delta(\text{He})$ for He@C₆₀. Interestingly, $\delta(\text{He})$ does not vary regularly along the $A_{12}H_{12}^{2-}$ (A = B-Al-Ga) series; the δ (He) values are similar (-22.2 and -23.1 ppm) in the borane and gallane complexes but only -18.4 ppm in the alane. A comparable trend is observed for NICS values in the center of the free $A_{12}H_{12}^{2-}$ dianions (-25.9 and -25.5 ppm for $B_{12}H_{12}^{2-}$ and $Ga_{12}H_{12}^{2-}$, respectively, vs -20.5 ppm for $Al_{12}H_{12}^{2-}$). Note that other properties of the closo systems (documented above) exhibited similar irregular changes down group 13. The alanes have the largest cages and are the most flexible with respect to deformations. The magnetic criteria indicate that alane clusters are less aromatic than both the boranes and the gallanes.

In spacious fullerene cages like C₆₀ the chemical shifts of endohedral He and Ne atoms are nearly identical. le In contrast, in dense cages like $Ne@B_{12}H_{12}^{2-}$, $Ar@Al_{12}H_{12}^{2-}$, and Kr@Al₁₂H₁₂²⁻, where the internal pressure is high and cage deformation approaches the threshold, the $\delta(Ne)$, $\delta(Ar)$, and $\delta(Kr)$ chemical shifts change sign from negative to positive and increase sharply (+513.9, +484.5, and +976.7 ppm, respectively). In principle, one can think that the use of heavier rare gas atoms to probe dense endo clusters would be more informative than the use of light Ng's. For instance, our calculations indicate that in alanes and gallanes the changes of $\delta(He)$ from the He@ $A_{12}H_{12}^{2-}$ anion to Li⁺[He@ $A_{12}H_{12}^{2-}$] with a face-coordinated external cation Li⁺ do not exceed few parts per million, while in their Ne analogues the changes are several times larger.

Transition States and Potential Barriers for Endohedral Clusters. Since all Ng@A₁₂H₁₂²⁻ clusters considered here are higher in energy than dissociated Ng + $A_{12}H_{12}^{2-}$, the barrier heights $(H_{\text{exit}}^{\ddagger})$ for Ng expulsion from the cage provide

estimates of $H_{\text{exit}}^{\dagger}$ provided general indications of stability but only employed partially optimized (B3LYP/6-31G*) "surface" structures 2, C_2 , and 3, $C_{3\nu}$, where the Ng's were positioned at the center of an edge or of a face, respectively.¹² The search for the real transition states (TS) is far more demanding due to the complexity of the opening of the Ng@A₁₂H₁₂²⁻ cages and the motions of the exiting heteroatom. For instance, for the He@B₁₂H₁₂²⁻ cluster, which dissociates into He and B₁₂H₁₂²⁻ with a moderate exothermicity, the usual saddle point optimization procedures with only initial force constant calculations [i.e., opt = (ts,calcfc)] as well as the quadratic synchronous transit (opt = qst2) procedures proved to be inefficient. Neither approach converged after more than 100 optimization cycles due to oscillations of the He atom inside the cage. The most successful procedure was the time-demanding optimization with recalculation of force constants at each step [opt = (ts, calcall)]. Although all these searches were carried out without any symmetry constraints, the optimized transition structure has C_2 symmetry (TS-1 in Figure 3). The single imaginary frequency eigenvector is directed along the 2-fold molecular axis toward the "active" B-B edge 4-6. The cage in **TS-1** is deformed and its geometry differs substantially from the initial icosahedron symmetry. The "active" edge 4-6 is stretched to over 3.3 Å, and the attached hydrogen atoms H₁₆ and H₁₈ are twisted strongly in opposite directions. In the transition state the He atom, located on a 2-fold axis, has moved only ~0.6 Å from its location in the former center of the icosahedron. NBO population analysis indicates that this He shift is accompanied by significant redistribution of electron density from atoms B₄ and B₆ of the "active" edge (by 0.45 e from each of them) to the neighbor cage atoms, mostly B_1 , B_5 , B_7 , and B_{10} . Intrinsic reaction coordinate (IRC) calculations at B3LYP/ 6-31G* confirmed that **TS-1** lies on the minimal energy pathway for expulsion of He from the cage and connects the endohedral structure 1 with the decomposition products He + $B_{12}H_{12}^{2-}$.

information about their kinetic stability. Our preliminary

The barrier height in $He@B_{12}H_{12}^{2-}$ relative to structure 1 is calculated to be 63.5 kcal/mol at B3LYP/6-31G* with ZPE correction. This result confirms our earlier conclusion¹² concerning the high kinetic stability of icosahedral clusters with an internal He atom. However, the energy of structure $2(C_2)$, where He was fixed in the center of the broken 4-6 edge, is

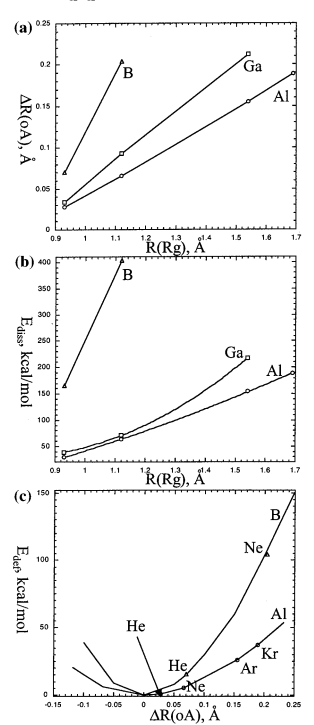


Figure 2. Dependence of endohedral Ng@A₁₂H₁₂²⁻ cluster structure and energy parameters on the Ng atomic radius (R(Ng)): (a) $\Delta R(oA)$ / R(Ng), Å; (b) $E_{diss}/R(Ng)$, kcal/mol; (c) $E_{def}/\Delta R(oA)$, kcal/mol. R(oA)is the distance between the center and a vertex of the A₁₂H₁₂² icosahedron, $E_{\rm diss}$ is the exothermicity of the Ng@A₁₂H₁₂²⁻ \rightarrow Ng + $A_{12}H_{12}^{2-}$ reaction, and E_{def} is the cage strain energy due to Ng insertion. The last was based on the energy difference between the relaxed and deformed cage (without the Ng atom).

 \sim 38 kcal/mol less than **TS-1**. Hence, although **2** can easily be calculated by partial optimization, its relative energy provides only a crude estimate of $H_{\text{exit}}^{\ddagger}$ in boranes.

The Ne@ $B_{12}H_{12}^{2-}$ cluster is extremely strained, and the endohedral atom expulsion reaction is highly exothermic. However, we were able to locate the TS easily using the quadratic synchronous transit method (also without symmetry constraints).

The calculated TS structure (TS-2 in Figure 3) essentially converged to $C_{3\nu}$ symmetry. The Ne atom is located on the 3-fold axis and shifted from the icosahedron center toward a face by only ~ 0.10 Å. However, the B-B edges in the "breaking" face have elongated from 1.86 Å (structure 1) to \sim 2.4 Å in **TS-2**. IRC calculations showed that when the Ne atom moves from its position in TS-2 toward the cage face by 0.1 Å along the minimal energy reaction path, the B-B edges of this face lengthen by an additional 0.25 Å. In the optimized C_{3v} model structure 3, where Ne was fixed in the center of the broken face, the distances between atoms 1-2-3 are longer, \sim 3.3 Å, but the energy of **TS-2** is about 90 kcal/mol higher than 3. The barrier height $H_{\text{exit}}^{\ddagger}$ calculated relative to structure 1 is only 2.6 kcal/mol with the ZPE correction. Thus, the $Ne@B_{12}\ddot{H}_{12}^{2-}$ cluster stability is predicted to be very low.

The character of the potential energy surface (PES) and that of the expulsion TS's change drastically for the less rigid alane clusters. The initial geometry in the He@Al₁₂H₁₂²⁻ TS search was based on 2, similar to **TS-1** for He@B₁₂H₁₂²⁻. The structure changed dramatically, and the optimized (also without symmetry restrictions) TS acquired an unusual open shape with large deformations in the "active" hemisphere of the cluster (see TS-3 in Figure 3). **TS-3** possesses C_s symmetry and resembles a "pot with open lid" where the "capping" Al₆–H₁₈ group is strongly shifted away from the 5-fold C₅ axis, and among five former bonds of Al₆ with neighbor cage atoms, only two, Al₆-Al₁ and Al₆-Al₅, remain, while the other edges including the base of the "lid" Al₁₋Al₅ are broken. In contrast to TS-1 (for $He@B_{12}H_{12}^{2-}$) and **TS-2** (for Ne@B₁₂H₁₂²⁻) where the heteroatoms are still located inside the cages, the He atom has nearly exited the cage in **TS-3** (for He@Al₁₂H₁₂ $^{2-}$). The most favorable exit pathway through a pentagonal aperture with a larger number of broken and elongated Al-Al bonds²³ can be described as a "decapping" mechanism. $H_{\text{exit}}^{\ddagger}$ for He@Al₁₂H₁₂²⁻ (58.5 kcal/mol including ZPE), only ~8 kcal/mol lower than $H_{\rm exit}$ [‡] for He@B₁₂H₁₂²⁻, is large enough to ensure a large kinetic stability of the He@Al₁₂H₁₂²⁻ cluster predicted earlier. ¹²

The Ne@Al₁₂H₁₂²⁻ TS search (at B3LYP/6-31G*) used three different initial geometries corresponding to TS-1, TS-2, and TS-3, but only the opened TS-3a resulted (Figure 3). In contrast to He@Al₁₂H₁₂²⁻ TS-3, in TS-3a the Ne atom remains closer to the icosahedron center despite the significant deformations of Al-Al bonds seen throughout the skeleton. The barrier $H_{\text{exit}}^{\ddagger}$ for Ne exit from Ne@Al₁₂H₁₂²⁻ via TS-3a is 44.4 kcal/mol relative to structure 1. This barrier is \sim 17 kcal/mol lower than that for $He@Al_{12}H_{12}^{2-}$, but still high enough to guarantee the kinetic stability of Ne@Al₁₂H₁₂²⁻.

Like the He and Ne analogues, the exit transition state for Ar@Al₁₂H₁₂²⁻ has a **TS-3** type structure (see **TS-3b** in Figure 3). In this TS, Ar is located even closer to the center of the icosahedron than Ne in TS-3a, and the cage deformations are severe. $H_{\text{exit}}^{\ddagger}$ is 15.2 kcal/mol and Ar@Al₁₂H₁₂²⁻ is kinetically much less stable than its lighter analogues. Its existence apparently would be possible only in low-temperature inert matrixes, in combination with bulky cations. Because the cage strain is so high in $Ar@Al_{12}H_{12}^{2-}$ (as in $Ne@B_{12}H_{12}^{2-}$), even small 0.3 Å displacements of Ar from the cage center in TS-3b result in very strong deformation of the skeleton and rupture of a number of Al-Al bonds. The structure of the

⁽²³⁾ Assignment of the number of "ruptured" A-A bonds in the TS of elastic clusters like Ng@Al₁₂H₁₂²⁻ often is not simple since the entire cage is strained and various edges are elongated considerably. Therefore, the boundary between "completely ruptured" and "strongly stretched" bonds is not clear.

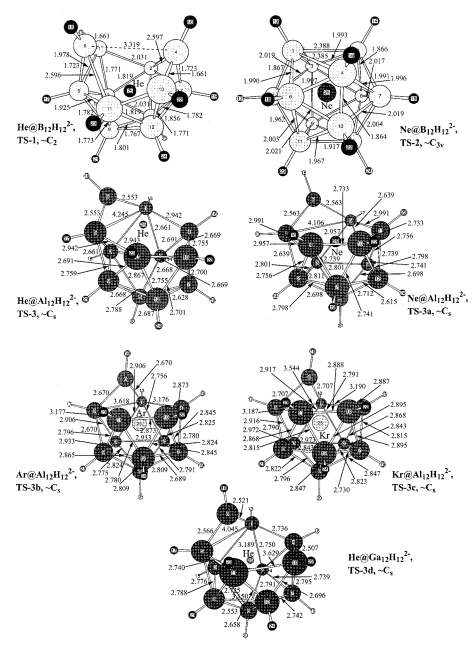


Figure 3. B3LYP/6-31G* optimized transition structures for the Ng@A₁₂H₁₂² $^ ^-$ Ng + A₁₂H₁₂² $^-$ reaction. The B3LYP/SDD method was employed for He@Ga₁₂H₁₂² $^-$, **TS-3d.** In **TS-3**, He@Al₁₂H₁₂² $^-$, the He atom is roughly located in the middle of a planar pentagon with average $R(\text{He-Al}) \approx 2.6 \text{ Å}$ and displacement above the plane of $\sim 0.1 \text{ Å}$.

transition state **TS-3c** for Kr@Al₁₂H₁₂²⁻ is quite similar to that of **TS-3b**, with the barrier $H_{\rm exit}^{\ddagger}$ of 10.5 kcal/mol, so this species is expected to be the least stable kinetically among the alane clusters.

The alanes with He, Ne, Ar, and Kr show a clear trend, based on simple steric considerations, which characterize the **TS-3** type transition states: with increasing noble gas atom radius the barrier height $H_{\text{exit}}^{\ddagger}$ drops rapidly, the torsion angle between the planes of triangular "lid" and pentagonal aperture and the Ng atom shift from the icosahedron center toward the attacked vertex in TS also decrease (in this sense, TS geometry develops sooner), and the rupture and elongation of the Al–Al bond is less localized to the active hemisphere and involves the entire cage. Similar trends can be expected for the gallanes and related electron-deficient clusters with dense, elastic cages (e.g., indanes, $X@In_{12}H_{12}^{2-}$, thallanes, $X@Tl_{12}H_{12}^{2-}$, and "naked" aluminides, $X@Al_{12}^{4-}$). The largest changes may be anticipated in going

from Ne to Ar because the largest change in the size of Ng occurs (N.B. the effect of Ne/Ar interchange in structure 1, above).

Our gallane studies were less extensive. B3LYP/SDD transition state optimization of He@Ga $_{12}$ H $_{12}$ ²⁻ used the Stuttgart—Dresden effective core potential for Ga atoms. ²⁴ The He exit transition state similar to **TS-3** resulted in **TS-3d** (see Figure 3). The He atom lies below the plane of the uncovered pentagonal aperture, and deformations of all the cage bonds are large. The **TS-3d** barrier height is about 32 kcal/mol. However, the effective core potential approach is less reliable than the all-electron B3LYP/6-31G* method. Note that our parallel B3LYP/6-31G* and B3LYP/SDD calculations on He and Ne

^{(24) (}a) Dolg, M.; Wedig, U.; Stoll, H.; Preuss, H. J. Chem. Phys. 1987, 86, 866. (b) Dolg, M.; Stoll, H.; Preuss, H. J. Chem. Phys. 1989, 90, 1730. (c) Kaupp, M.; Schleyer, P. v. R.; Stoll, H.; Preuss, H. J. Chem. Phys. 1991, 94, 1360.

alanes and gallanes show systematic differences between the two methods. The Ga-Ga bond lengths in the B3LYP/SDD optimized icosahedral structure are 0.20-0.25 Å larger than at B3LYP/6-31G*, vibrational frequencies are underestimated, and the alane transition states, although of the TS-3 type, are strongly deformed and lose C_s symmetry. Therefore, we can only conclude qualitatively that gallane endohedral clusters should be less stable kinetically than the corresponding alane clusters.

Since the transition states for all alanes and gallanes considered here belong to the TS-3 type, one can expect that the "decapping" mechanism for expulsion of the endohedral atom will be a feature of other related electron-deficient clusters formed by the third row atoms and heavier elements. In contrast, clusters of light second row elements, with strong σ - and π -bonds, should find this pathway significantly less favorable as compared, for instance, with an exit via the cage edge.

Influence of External Cations. Because of the close relationship to the Ng@ $A_{12}H_{12}^{2-}$ dianions, we performed similar calculations on Li[Ng@A₁₂H₁₂]⁻ anions and neutral Li₂-[Ng@A₁₂H₁₂] molecules. The influence of external cations on cluster structure and stability was assessed with the structures **4**, C_{3v} (one Li⁺ cation located above a cage face) and **5**, D_{3d} (two Li⁺ cations located above opposite cage faces) (Figure 1). These structures are local minima. The related empty anion systems L[A₁₂H₁₂]⁻, with H⁺, Al⁺, Al³⁺, alkali metal, and alkaline earth metal cations coordinated, were investigated carefully earlier.¹² We compare the results for empty and Ngcentered salts here and concentrate on the peculiarities caused by the presence of the internal atom Ng.

As expected, the cation influences differ substantially and correlate with the exothermicities of the Ng@ $A_{12}H_{12}^{2-} \rightarrow Ng$ $+ A_{12}H_{12}^{2-}$ reactions. In He@Al₁₂H₁₂²⁻ or He@Ga₁₂H₁₂²⁻, where skeletal strain is small, the presence of He does not change the deformation of the $A_{12}H_{12}^{2-}$ anion in the field of a cation significantly. The coordinated face in 4 expands, the opposite face contracts, the bridging H_b hydrogen atoms are shifted toward the Li⁺ cation, and there is a parallel decrease in the H_bALi angle and the elongation in the $R(A-H_b)$ distance. Deformations of the less rigid alanes are much larger than the boranes, especially when the cation radii are small and the cations are multiply charged. Degenerate vibrational frequencies and orbital energies split when the cation field reduces the symmetry from I_h to $C_{3\nu}$ or D_{3d} . The exothermicity of He expulsion from the alane and the borane cages increases for $Li[He@A_{12}H_{12}]^-$ compared with $He@A_{12}H_{12}^{2-}$ by just a few kilocalories per mole. The increase in exothermicity for boranes is 10 kcal/mol, but this change does not exceed 6% $E_{\rm diss}$.

The cation-anion interaction can drastically change the structure of systems with very exothermic E_{diss} and with a large Ng atom inside. The cage strain is large and the influence of the cation may even be sufficient to overcome the dissociation threshold and lead to expulsion of Ng from the cage. For example, Li⁺ coordination to Ne@B₁₂H₁₂²⁻ causes the Ne atom to exit from the cage without barrier: structure 4 dissociates into Ne and LiB₁₂H₁₂⁻. However, 4 corresponds to a local minima for Ne@Al₁₂H₁₂²⁻ and for Ne@Ga₁₂H₁₂²⁻. The aluminum and gallium cages in 4 are deformed by the Li⁺ cations to approximately the same extent as in the C_{3v} LiAl₁₂H₁₂⁻ and $LiGa_{12}H_{12}^{-}$ anions. In other words, the metastable Ne@B₁₂H₁₂²⁻ becomes unstable even when it is disturbed by one Li⁺ cation. Our calculations on the Na[Ne@B₁₂H₁₂]⁻ and Al[Ne@B₁₂H₁₂]⁻ systems showed that replacement of Li⁺ with the larger Na⁺ and Al⁺ cations does not change the result; in both cases Ne leaves the cage without a barrier.

The combined effects of two Li+ cations coordinated to opposite faces of the Ng@A₁₂H₁₂²⁻ icosahedrons in boranes strongly differ from those in alanes and gallanes due to the differences in the strengths of the A-A bonds. For instance, the closo structure 5 is maintained in Li₂[He@B₁₂H₁₂] where the borane cage is only slightly deformed as compared to the He@B₁₂H₁₂²⁻ anion. The B-B bond lengths in this salt are in the 1.822–1.879 Å range, close to 1.861 Å in He@B₁₂H₁₂^{2–}, and the B-H bond lengths change within 0.01-0.02 Å. The geometry deformation from the empty Li₂B₁₂H₁₂ salt to endohedral Li₂[He@B₁₂H₁₂] is rather similar to that discussed for the anions $B_{12}H_{12}^{2-}$ and $He@B_{12}H_{12}^{2-}$. In contrast, the closo structures of the less rigid alane Li₂[He@Al₁₂H₁₂] and gallane Li₂[He@Ga₁₂H₁₂] salts are destroyed: the equatorial two-center Al-H bonds transform into three-center bridging bonds, and the D_{3d} structures 5 optimize in D_{3d} symmetry without barriers into new, unusual structures 6 where only some of the of Al—Al and Ga—Ga bonds remain. Structure 6 for both the alane and the gallane have one a_{1u} imaginary frequency which corresponds to the transition state for complex intramolecular rearrangements. Apparently, even the kinetically stable Ng alane and gallane dianions would be stable in salts only with bulky countercations.

Conclusions

We have computed related closo-boranes, -alanes, and -gallanes Ng@ $A_{12}H_{12}^{2-}$ with noble gas atoms inside the [A₁₂] cages. The icosahedral endohedral structures 1 are local minima, significantly higher in energy than the Ng + $A_{12}H_{12}^{2-}$ dissociation limits for $\text{He@B}_{12}\text{H}_{12}^{2-}$, $\text{Ne@B}_{12}\text{H}_{12}^{2-}$, $\text{He@Al}_{12}\text{H}_{12}^{2-}$, $\text{Ne@Al}_{12}\text{H}_{12}^{2-}$, $\text{Ar@Al}_{12}\text{H}_{12}^{2-}$, $\text{Kr@Al}_{12}\text{H}_{12}^{2-}$, $\text{He@Ga}_{12}\text{H}_{12}^{2-}$, and Ne@Ga₁₂ H_{12}^{2-} . The exothermicity of the expulsion of Ng from a cage increases rapidly with increased noble gas atomic radius. $E_{\rm diss}$ is very large for boranes (166 and 403 kcal/mol with Ng = He and Ne, respectively) and much smaller for alanes (29, 63, 154, and 189 kcal/mol for Ng = He, Ne, Ar, and Kr,respectively), but the values for the gallanes (39 and 71 kcal/ mol with Ng = He and Ne, respectively) are somewhat larger than those for the alanes.

There are three types of Ng exit transition state: via an edge (TS-1); through a face (TS-2); and via a pentagonal "neck" of the cage, much like a "pot with open lid" (TS-3). These favored TS types in different clusters depend on the cage rigidity, the expulsion reaction exothermicity, and the relation between the size of cage internal cavity and the internal Ng atomic radius. When dissociation is only moderately exothermic (e.g., alanes and gallanes with He and Ne inside), Ng exit via a pentagonal "neck" is preferred (**TS-3**). In He@B₁₂H₁₂ $^{2-}$ exit occurs via a cage edge (TS-1). In Ne@B₁₂H₁₂²⁻, which has the largest dissociation exothermicity, the exit occurs via a triangular face (TS-2). $H_{\text{exit}}^{\ddagger}$ is quite high (30–60 kcal/mol) for all heliumcontaining clusters, but the barriers are lower for the Ne@Al₁₂H₁₂²⁻ and Ne@Ga₁₂H₁₂²⁻ systems (e.g., Ar@Al₁₂H₁₂²⁻ and Kr@Al₁₂H₁₂²⁻ barriers are about 15 and 10 kcal/mol, respectively) and the kinetic stability of these systems is low. The Ne@B₁₂H₁₂²⁻ cluster has a very low barrier and is not expected to be observable.

The Ng@A₁₂H₁₂²⁻ cage geometry exhibits a linear dependence on the atomic radius of the internal noble gas. NMR chemical shifts for He atoms in the clusters are negative, while for the other Ng's NMR chemical shifts are positive and have much larger values. NICS values indicate that all three A₁₂H₁₂² anions are highly aromatic with the alanes somewhat less so. Charge transfer between the internal atom and cage is insignificant in clusters with He and Ne, so that steric and electronic repulsion between the closed electronic shells of Ng and $A_{12}H_{12}{}^{2-}$ play a crucial role in the heteroatom—cage interactions. A medium intense t_{1u} band in the 925–1030 cm $^{-1}$ region for boranes and 340–560 cm $^{-1}$ for alanes and gallanes corresponding mainly to vibrations of the internal atom should be the most characteristic feature of IR spectra of Ng@ $A_{12}H_{12}{}^{2-}$ systems.

The effect of one external Li⁺ located above a cage face of the lithium salts Li[Ng@ $A_{12}H_{12}$]⁻ is not large in the helium clusters. However, in systems such as Li[Ne@ $B_{12}H_{12}$]⁻, which have extremely large expulsion exothermicities, the internal atom exits from the cage and the salt dissociates into Ne + Li $B_{12}H_{12}$ ⁻ without a barrier. Systems with two Li⁺ ions located above opposite cage faces, (Li₂[Ng@ $A_{12}H_{12}$]), undergo complex

intramolecular rearrangements which destroy the skeletal closo structure.

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Supporting Information Available: Optimized Cartesian coordinates of various local minima and transition states. This material is available free of charge via the Internet at http://pub.acs.org.

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