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The Effect of Substituent on Molecules That Contain a **Triple Bond Between Arsenic and Group 13 Elements: Theoretical Designs and Characterizations**

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Additional information is available at the end of the chapter

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Abstract

The effect of substitution on the potential energy surfaces of RE_{13} =AsR (E_{13} = group 13 elements; R = F, OH, H, CH₃, and SiH₃) is determined using density functional theory (M06-2X/Def2-TZVP,B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp). The computational studies demonstrate that all triply bonded RE₁₃≡AsR species prefer to adopt a bent geometry that is consistent with the valence electron model. The theoretical studies also demonstrate that RE₁₃=AsR molecules with smaller substituents are kinetically unstable, with respect to the intramolecular rearrangements. However, triply bonded $R'E_{13} = AsR'$ species with bulkier substituents ($R' = SiMe(SitBu_3)_{\gamma}$, SiiPrDis₂, and NHC) are found to occupy the lowest minimum on the singlet potential energy surface, and they are both kinetically and thermodynamically stable. That is to say, the electronic and steric effects of bulky substituents play an important role in making molecules that feature an $E_{1,\Xi}$ As triple bond as viable synthetic target.

Keywords: arsenic, group 13 elements, triple bond, density functional theory, multiple bond

1. Introduction

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In the past two decades, studies that have been performed by many synthetic chemists have successfully synthesized and characterized homonuclear heavy alkyne-like $RE_{14} = E_{14}R$ ($E_{14} = Si$, Ge, Sn, and Pb) molecules [1–23]. Recently, heteronuclear ethyne-like compounds, $RC = E_{14}R$, have also been experimentally studied [24, 25, 26] and theoretically predicted [27, 28, 29].

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However, from the valence electron viewpoint, $RE_{13} = E_{15}R$ ($E_{13} = \text{group 13}$ elements and $E_{15} = \text{group 15}$ elements) is isoelectronic with the $RE_{14} = E_{14}R$ species. Therefore, triply bonded $RE_{13} = E_{15}R$ is the next synthetic challenge. To the best of the authors' knowledge, only R_2BN molecules that contain a B = N triple bond have been experimentally demonstrated to exist [30–40].

2. Theoretical methods

This chapter reports the possible existence of triply bonded RE_{13} =AsR molecules, from the viewpoint of the effect of substituents, using density functional theories (DFT): M06-2X/Def2-TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp for small substituents and B3LYP/LANL2DZ+dp//RHF/3-21G* for large substituents. It is hoped that this theoretical study will stimulate further research into the synthetic chemistry of triply bonded RE₁₃=AsR species.

3. Results and discussion

3.1. Small ligands on substituted RE₁₃=AsR

The effect of the electronegativity of six types of small substituents (R = F, OH, H, CH₃, and SiH₃) on the stability of the triply bonded RE₁₃=AsR molecules is determined using the three DFT methods. The molecular properties (geometrical parameters, singlet-triplet energy splitting, natural charge densities, binding energies (BE), and the Wiberg Bond Index (WBI)) are all listed in **Tables 1–5**. The reaction profiles for the unimolecular rearrangement reactions for the RE₁₃=AsR compounds are also given in **Figures 1–5**.

There are three noteworthy features of **Tables 1–5** and **Figures 1–5**.

- From the tables, the three DFT calculations show that the triple bond distances (Å) for B=As, Al=As, Ga=As, In=As, and Tl=As are estimated to be 1.835–1.908 (Table 1), 2.218–2.358 (Table 2), 2.239–2.364 (Table 3), 2.404–2.546 (Table 4), and 2.426–2.570 (Table 5). As previously mentioned, no experimental values for these triple bond lengths have been reported, so these computational data are a prediction.
- 2. In Tables 1–5, these DFT computations all demonstrate that the triply bonded RE_{13} =AsR molecules favor a bent structure, rather than a linear structure. This is explained by the bonding model, as shown in Figure 6. Because there is a significant difference between the sizes of the valence s and p atomic orbitals in the As atom, hybrid orbitals between the valence s and p orbitals are not easily formed (the so-called orbital non-hybridization effect or the inert s-pair effect) [41–44]. Therefore, RE_{13} =AsR molecules that have a heavier As center are predicted to favor a bent angle $\angle E_{13}$ -As-R (close to 90°). The DFT computational data that are shown in Tables 1–5 confirm this prediction.

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R	F	ОН	Н	CH3	SiH3
B≡As (Å)	1.901	1.892	1.837	1.839	1.814
	(1.898)	(1.888)	(1.835)	(1.839)	(1.820)
	[1.908]	[1.906]	[1.849]	[1.861]	[1.839]
∠R–B–As (°)	177.2	179.5	178.1	175.1	175.3
	(177.8)	(179.5)	(174.6)	(175.1)	(172.4)
	[177.0]	[179.1]	[177.5]	[174.3]	[174.8]
∠B–As–R (°)	93.03	92.73	81.22	94.69	68.92
	(92.71)	(92.21)	(89.39)	(94.69)	(68.98)
	[92.39]	[92.95]	[78.37]	[96.15]	[72.25]
∠R–B–As–R (°)	180.0	179.8	180.0	179.8	148.7
	(180.0)	(180.0)	(180.0)	(179.8)	(180.0)
	[180.0]	[176.2]	[179.0]	[176.3]	[179.4]
Q_{B}^{1}	0.354	0.184	-0.017	-0.007	0.037
	(0.262)	(0.108)	(-0.028)	(-0.057)	(0.036)
	[0.232]	[0.070]	[-0.106]	[-0.160]	[-0.407]
$Q_{\rm As}^{2}$	0.243	0.080	-0.152	-0.073	-0.085
	(0.255)	(0.097)	(-0.134)	(-0.040)	(-0.017)
	[0.238]	[0.086]	[0.034]	[-0.035]	[0.030]
BE (kcal mol ⁻¹) ³	63.56	56.97	114.7	94.39	79.90
	(63.34)	(60.28)	(120.1)	(137.6)	(74.75)
	[57.45]	[55.28]	[113.7]	[132.6]	[73.79]
WBI ⁴	1.800	1.830	2.141	2.027	2.204
	(1.813)	(1.823)	(2.158)	(2.029)	(2.168)
	[1.835]	[1.836]	[2.135]	[2.041]	[2.185]

1 The natural charge density on the B atom.

2 The natural charge density on the As atom.

3 BE = E(triplet state for R–B) + E(triplet state for R–As) – E(singlet state for RB \equiv AsR).

4 The Wiberg bond index (WBI) for the B–As bond: see [45, 46].

Table 1. The main geometrical parameters, the singlet-triplet energy splitting (ΔE_{sT}), the natural charge densities (Q_B and Q_{As}), the binding energies (BE), and the Wiberg Bond Index (WBI) for RB=AsR using the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP (in round brackets), and B3LYP/LANL2DZ+dp (in square brackets) levels of theory.

R	F	ОН	Н	CH3	SiH3
Al≡As (Å)	2.327	2.321	2.218	2.253	2.227
	(2.325)	(2.323)	(2.221)	(2.256)	(2.236)
	[2.355]	[2.358]	[2.269]	[2.285]	[2.292]
∠R–Al–As (°)	178.6	174.4	172.5	172.8	168.4
	(179.5)	(174.3)	(172.2)	(172.0)	(167.3)
	[178.8]	[173.9]	[177.5]	[171.1]	[173.7]
∠Al–As–R (°)	93.07	91.08	66.95	98.77	91.93
	(93.51)	(92.45)	(67.45)	(100.7)	(95.83)
	[90.64]	[90.97]	[75.97]	[100.5]	[90.36]
∠R–Al–As–R (°)	180.0	180.0	180.0	174.2	174.7
	(179.8)	(178.5)	(179.6)	(176.8)	(175.7)
	[180.0]	[179.0]	[178.0]	[174.5]	[176.8]
$oldsymbol{Q}_{ ext{Al}}^{-1}$	0.555	0.4574	0.2401	0.293	0.291
	(0.530)	(0.443)	(0.234)	(0.280)	(0.313)
	[0.784]	[0.540]	[0.504]	[0.353]	[0.245]
$Q_{\rm As}^{2}$	0.158	0.015	-0.276	-0.170	-0.262
	(0.142)	(-0.007)	(-0.246)	(-0.156)	(-0.209)
	[0.056]	[-0.032]	[-0.209]	[-0.284]	[-0.290]
BE (kcal mol ⁻¹) ³	33.90	28.23	71.86	56.47	53.22
	(38.90)	(31.24)	(77.42)	(60.57)	(54.98)
	[33.89]	[25.68]	[69.27]	[52.63]	[67.74]
WBI ⁴	1.532	1.523	1.714	1.649	1.647
	(1.567)	(1.553)	(1.742)	(1.679)	(1.675)
	[1.557]	[1.545]	[1.714]	[1.690]	[1.550]

2 The natural charge density on the As atom.

3 BE = E(triplet state for R−Al) + E(triplet state for R−As) – E(singlet state for RAl≡AsR).

4 The Wiberg bond index (WBI) for the Al–As bond: see [45, 46].

Table 2. The main geometrical parameters, the singlet-triplet energy splitting (ΔE_{ST}), the natural charge densities (Q_{AI} and Q_{AS}), the binding energies (BE), and the Wiberg Bond Index (WBI) for RAI=AsR using the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP (in round brackets), and B3LYP/LANL2DZ+dp (in square brackets) levels of theory.

3. In terms of the stability of the RE₁₃≡AsR species, the three DFT computations are used to study the energy surfaces for the RE₁₃≡AsR systems, and the theoretical results are shown in Figures 1–5. These figures show three local minima (i.e., R₂E₁₃=As, RE₁₃≡AsR, and E₁₃=AsR₂) and two saddle points that connect them. It is seen that regardless of the type of small substituent, triply bonded RE₁₃≡AsR molecules are unstable on the potential energy surfaces, so they easily undergo a 1,2-migration reaction to produce the most stable doubly bonded isomers. There is strong theoretical evidence that there is no possibility of observing triply bonded RE₁₃≡AsR compounds in transient intermediates or even in a matrix.

3.2. Large ligands on substituted $R'E_{13} \equiv AsR'$

Bulky substituents are used to determine the possible existence of triply bonded R'E₁₃=AsR' (R'= SiMe(SitBu₃)₂, SiiPrDis₂, and NHC; (**Scheme 1**)) molecules. The molecular properties, the natural bond orbital (NBO) [45, 46], and the natural resonance theory (NRT) [47, 48, 49] analyses of R'E₁₃=AsR' are computed at the B3LYP/LANL2DZ+dp//RHF/3-21G* level of theory, and the results are shown in **Tables 6**, **7** (R'B=AsR'), **8**, **9** (R'Al=AsR'), **10**, **11** (R'Ga=AsR'), **12**, **13** (R'In=AsR'), and **14** and **15** (R'Tl=AsR').

R	F	ОН	Н	СНЗ	SiH3
Ga≡As (Å)	2.261	2.339	2.239	2.330	2.243
	(2.319)	(2.314)	(2.224)	(2.243)	(2.242)
	[2.364]	[2.364]	[2.263]	[2.285]	[2.270
∠R–Ga–As (°)	179.5	173.2	176.2	169.9	168.5
	(178.5)	(177.4)	(178.6)	(173.6)	(179.1)
	[179.3]	[176.2]	[179.1]	[171.1]	[179.2]
∠Ga-As-R (°)	92.80	93.16	76.00	103.0	93.43
	(94.36)	(94.54)	(79.18)	(99.37)	(73.64)
	[91.81]	[93.68]	[80.30]	[100.4]	[76.86]
∠R–Ga–As–R (°)	180.0	175.6	179.6	175.7	173.5
	(180.0)	(178.1)	(179.1)	(178.4)	(175.6)
	[173.1]	[177.4]	[178.2]	[174.5]	[178.1]
$Q_{\rm Ga}^{-1}$	0.7067	0.592	0.310	0.4451	0.3352
	(0.554)	(0.410)	(0.215)	(0.260)	(0.241)
	[0.706]	[0.474]	[0.435]	[0.295]	[0.174]
$Q_{\rm As}^{2}$	0.0899	-0.047	-0.374	-0.256	-0.3697
	(0.154)	(0.023)	(-0.262)	(-0.151)	(-0.222)
	[0.133]	[0.006]	[-0.184]	[-0.246]	[-0.284]
BE (kcal mol ⁻¹) ³	28.56	23.82	67.79	53.57	49.26
	(30.61)	(25.96)	(71.91)	(58.12)	(51.77)
	[27.65]	[90.75]	[65.14]	[50.32]	[62.24]
WBI ⁴	1.476	1.498	1.691	1.648	1.646
	(1.486)	(1.503)	(1.717)	(1.652)	(1.596)
	[1.487]	[1.495]	[1.707]	[1.668]	[1.615]

1 The natural charge density on the Ga atom.

2 The natural charge density on the As atom.

3 BE = E(triplet state for R–Ga) + E(triplet state for R–As) – E(singlet state for RGa=AsR).

4 The Wiberg bond index (WBI) for the Ga–As bond: see [45, 46].

Table 3. The main geometrical parameters, the singlet-triplet energy splitting (ΔE_{st}), the natural charge densities (Q_{Ga} and Q_{As}), the binding energies (BE), and the Wiberg Bond Index (WBI) for RGa=AsR using the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP (in round brackets), and B3LYP/LANL2DZ+dp (in square brackets) levels of theory.

R	F	ОН	Н	CH3	SiH3
In≡As (Å)	2.511	2.512	2.412	2.431	2.411
	(2.495)	(2.497)	(2.399)	(2.418)	(2.404)
	[2.535]	[2.546]	[2.432]	[2.459]	[2.444]
∠R–In–As (°)	179.9	178.8	179.3	173.6	170.9
	(179.9)	(176.9)	(179.9)	(173.3)	(168.4)
	[177.8]	[175.2]	[179.8]	[172.5]	[167.4]
∠In-As-R (°)	92.32	95.31	81.43	99.72	93.85
	(93.86)	(96.11)	(82.67)	(100.4)	(99.59)
	[91.08]	[94.22]	[82.28]	[100.5]	[102.0]
∠R–In–As–R (°)	180.0	169.3	177.3	174.7	177.1
	(180.0)	(166.8)	(175.9)	(173.0)	(177.4)
	[180.0]	[163.8]	[179.6]	[179.8]	[178.2]
Q_{In}^{1}	1.288	1.233	1.012	1.144	0.8840
	(1.196)	(1.123)	(0.912)	(1.037)	(0.7881)
	[1.343]	[1.287]	[1.076]	[1.121]	[0.9682]
$Q_{\rm As}^{2}$	0.138	0.036	-0.624	-0.388	-0.767
	(0.146)	(0.047)	(-0.571)	(-0.335)	(-0.703)
	[0.077]	[-0.005]	[-0.591]	[-0.367]	[-0.748]
BE (kcal mol ⁻¹) ³	22.14	18.30	55.63	53.87	57.82
	(19.72)	(20.13)	(60.95)	(50.24)	(57.34)
	[24.06]	[16.22]	[57.18]	[53.36]	[54.39]
WBI ⁴	1.536	1.551	1.773	1.719	1.726
	(1.546)	(1.554)	(1.798)	(1.738)	(1.749)
	[1.572]	[1.562]	[1.780]	[1.729]	[1.710]

1 The natural charge density on the In atom.

2 The natural charge density on the As atom.

3 BE = E(triplet state for R–In) + E(triplet state for R–As) – E(singlet state for RIn=AsR).

4 The Wiberg bond index (WBI) for the In-As bond, see [45, 46].

Table 4. The main geometrical parameters, the singlet-triplet energy splitting (ΔE_{sT}), the natural charge densities (Q_{ln} and Q_{As}), the binding energies (BE), and the Wiberg Bond Index (WBI) for RIn=AsR using the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP (in round brackets), and B3LYP/LANL2DZ+dp (in square brackets) levels of theory.

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R	F	ОН	Н	CH3	SiH3
Tl≡As (Å)	2.535	2.531	2.426	2.446	2.431
	(2.533)	(2.536)	(2.428)	(2.450)	(2.432)
	[2.558]	[2.570]	[2.429]	[2.459]	[2.433]
∠R–T1–As (°)	179.9	178.2	180.0	176.6	176.5
	(179.9)	(175.8)	(179.5)	(175.0)	(173.4)
	[179.2]	[177.0]	[179.5]	[173.8]	[177.7]
∠Tl–As–R (°)	91.49	94.88	84.22	97.14	90.08
	(93.64)	(96.73)	(84.51)	(99.33)	(93.68)
	[92.21]	[96.20]	[84.07]	[99.33]	[89.37]
∠R–T1–As–R (°)	180.0	175.5	173.0	178.0	179.2
	(179.3)	(176.7)	(178.1)	(178.2)	(178.5)
	[180.0]	[172.9]	[179.6]	[177.6]	[177.2]
Q_{Π}^{-1}	0.736	0.640	0.3883	0.482	0.3051
	(0.656)	(0.538)	(0.352)	(0.428)	(0.382)
	[0.817]	[0.549]	[0.472]	[0.361]	[0.244]
$Q_{\rm As}^{2}$	0.190	0.035	-0.4169	-0.251	-0.3290
	(0.163)	(0.013)	(-0.351)	(-0.208)	(-0.291)
	[0.139]	[0.021]	[-0.204]	[-0.273]	[-0.336]
BE (kcal mol ⁻¹) ³	13.48	10.36	50.28	38.25	29.93
	(16.73)	(13.88)	(55.13)	(43.44)	(30.60)
	[15.13]	[8.720]	[49.40]	[37.22]	[45.10]
WBI ⁴	1.109	1.148	1.456	1.382	1.409
	(1.143)	(1.174)	(1.492)	(1.416)	(1.407)
	[1.168]	[1.175]	[1.484]	[1.413]	[1.411]

1 The natural charge density on the Tl atom.

2 The natural charge density on the As atom.

3 BE = E(triplet state for R–TI) + E(triplet state for R–As) – E(singlet state for RTI=AsR).

4 The Wiberg bond index (WBI) for the Tl-As bond, see [45, 46].

Table 5. The main geometrical parameters, the singlet-triplet energy splitting (ΔE_{sT}), the natural charge densities (Q_{TI} and Q_{As}), the binding energies (BE), and the Wiberg Bond Index (WBI) for RTI=AsR using the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP (in round brackets), and B3LYP/LANL2DZ+dp (in square brackets) levels of theory.

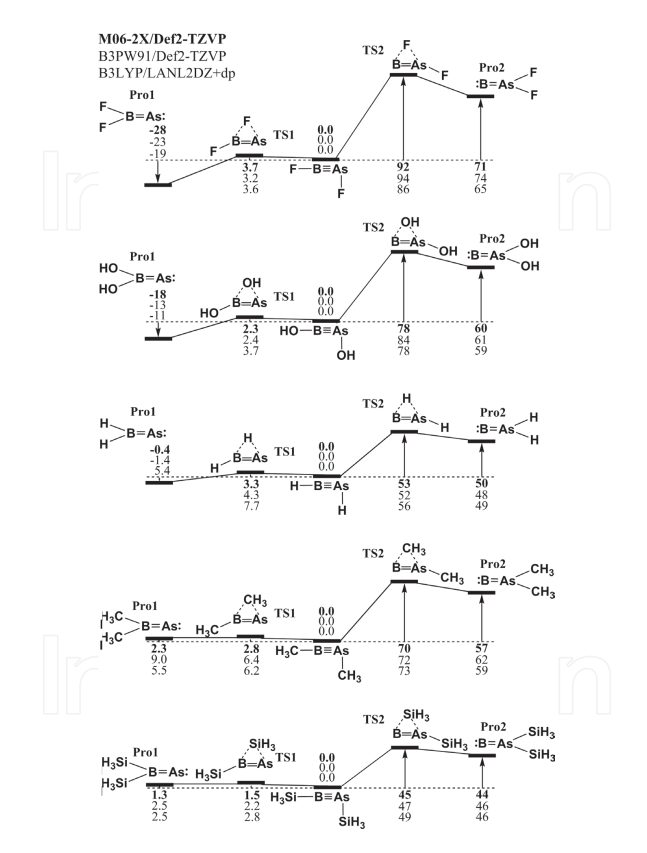


Figure 1. The relative Gibbs free energies for RB = AsR (R = F, OH, H, CH₃, and SiH₃). All energies are in kcal/mol and are calculated at the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp levels of theory.

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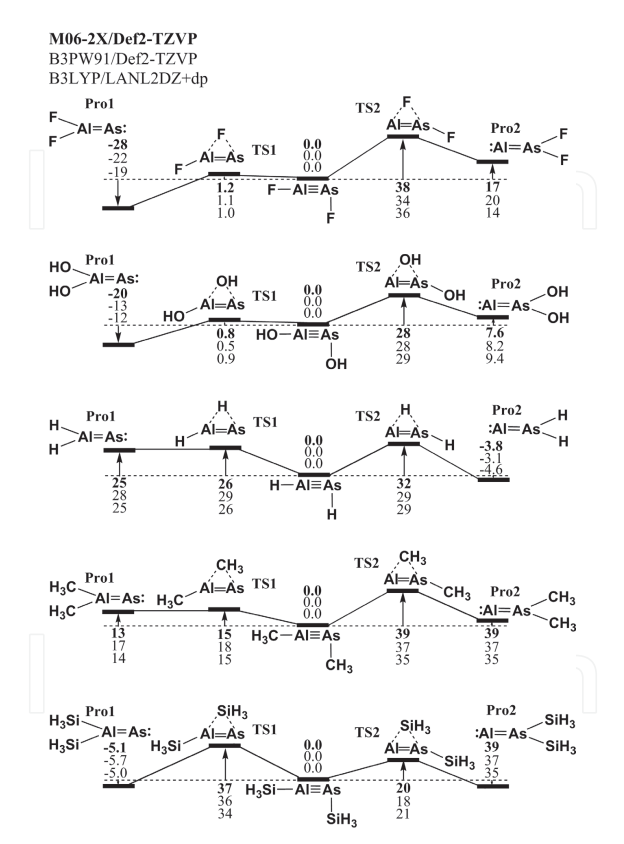


Figure 2. The relative Gibbs free energies for RAl=AsR (R = F, OH, H, CH₃, and SiH₃). All energies are in kcal/mol and are calculated at the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp levels of theory.

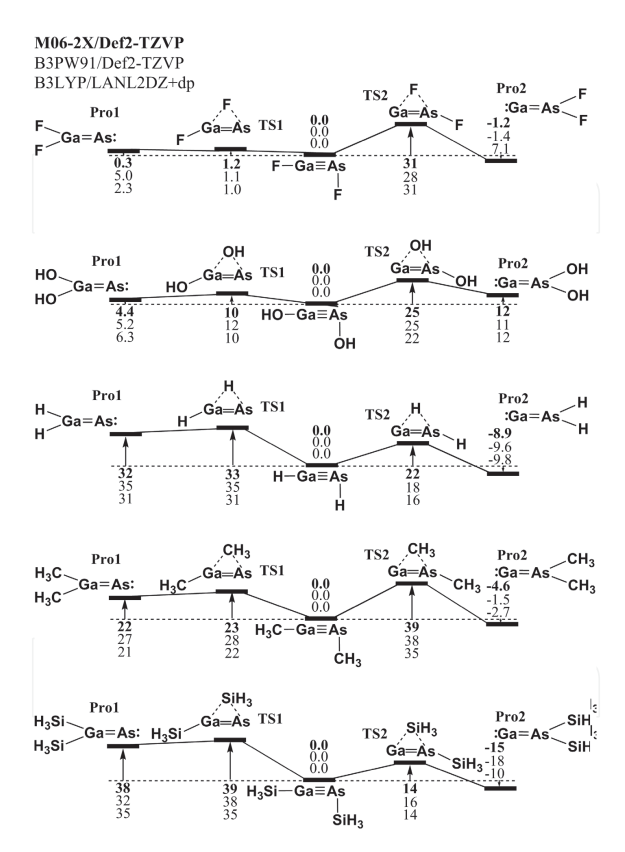


Figure 3. The relative Gibbs free energies for RGa = AsR (R = F, OH, H, CH₃, and SiH₃). All energies are in kcal/mol and are calculated at the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp levels of theory.

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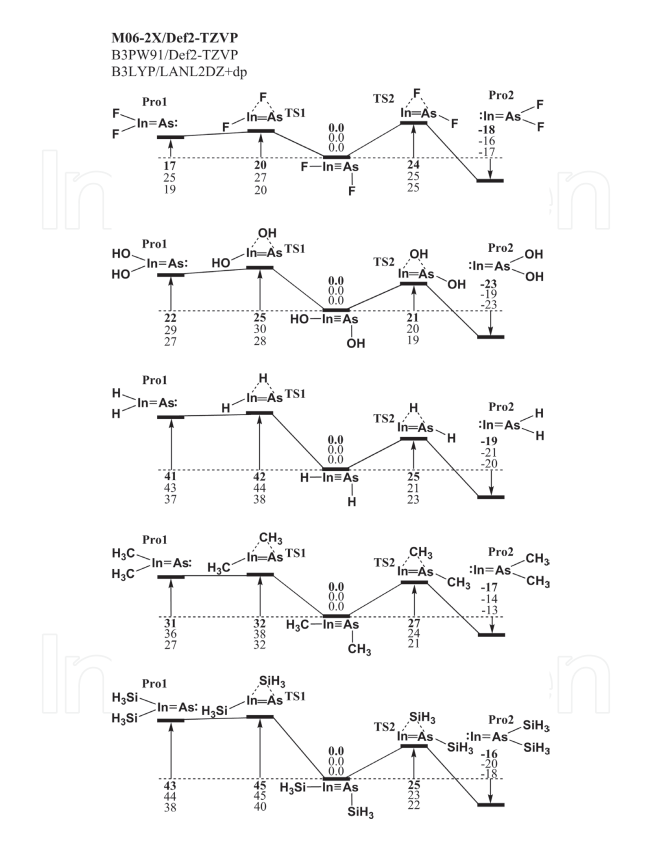


Figure 4. The relative Gibbs free energies for RIn=AsR (R = F, OH, H, $CH_{3^{\prime}}$ and SiH₃). All energies are in kcal/mol and are calculated at the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp levels of theory.

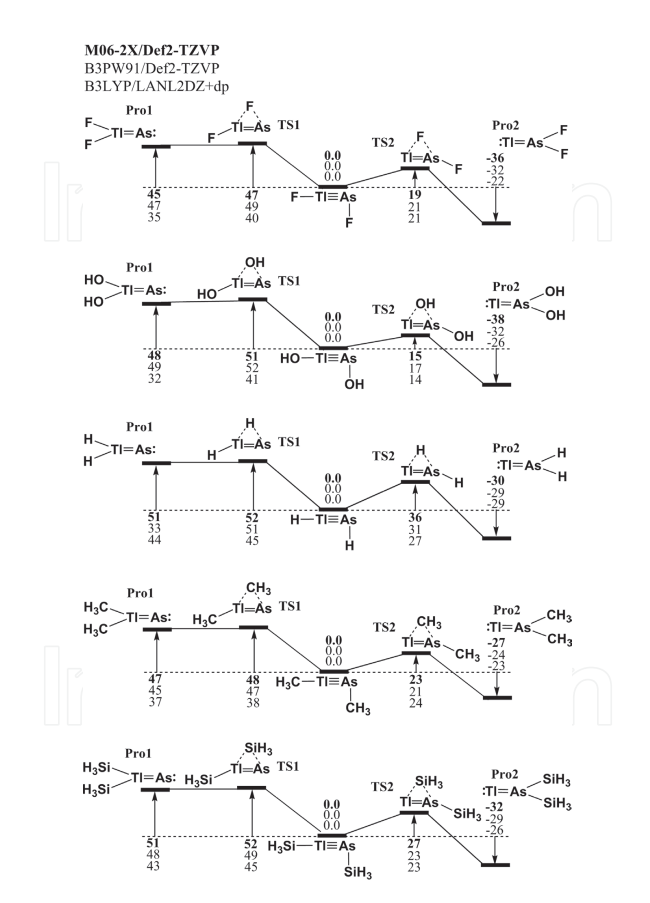
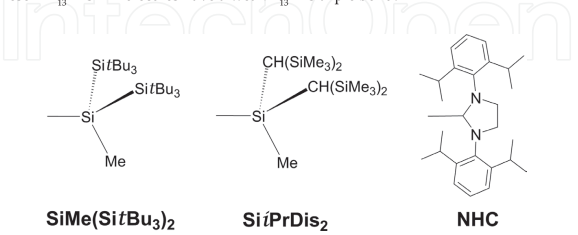


Figure 5. The relative Gibbs free energies for RTI=AsR (R = F, OH, H, CH₃, and SiH₃). All energies are in kcal/mol and are calculated at the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp levels of theory.

The results in Tables 6–15 allow three conclusions to be drawn.

- The calculations that are shown in Tables 6 (B), 8 (Al), 10 (Ga), 12 (In), and 14 (Tl) show that the computed E₁₃≡As triple bond distances (Å) for these bulkily substituted species (R'E₁₃≡AsR') are estimated to be 1.821–1.837 (B≡As), 2.257–2.307 (Al≡As), 2.252–2.316 (Ga≡As), 2.430–2.482 (In≡As), and 2.565–2.653 (Tl≡As). The values for the WBO that are shown in Tables 6–10 (for bulky ligands) are obviously greater than those that are shown in Tables 1–5 (for smaller ligands). These WBO values show that bulkier substituents increase the bond order for the E₁₃≡As triple bond length.
- 2. Similarly to the results for small ligands, the computational results show that R'E₁₃≡AsR' species that feature large substituents all adopt a bent conformation. This phenomenon is explained by bonding model (II), which is shown in **Figure 6**.
- 3. The NBO values that are shown in Tables 7 (B=As), 9 (Al=As), 11 (Ga=As), 13 (In=As), and 15 (Tl=As) show that the acetylene-like $R'E_{13}$ =AsR' compounds feature a weak triple bond. For example, the B3LYP/LANL2DZ+dp data for the NBO analyses of the B=As π bonding in $(SiiPrDis_2-B=As-SiiPrDis_2)$, which shows that NBO $(B=As) = 0.5880(2s2p^{99.99})$ $B + 0.8089(4s4p^{1.00})$ As, provide strong evidence that the predominant bonding interaction between the B-SiiPrDis, and the As-SiiPrDis, units results from $2p(B) \leftarrow 4p(As)$ donation, whereby boron's electron deficiency and π bond polarity are partially balanced by the donation of the arsenic lone pair into the empty boron *p* orbital to develop a hybrid π bond. The polarization analyses using the NBO model again demonstrate the presence of the B=As π bonding orbital, 34.58% of which is composed of natural B orbitals and 65.42% of which is natural As orbitals. **Table 7** also shows that the B=As triple bond in (Si*i*PrDis₂- $B=As-SiiPrDis_2$) has a shorter single bond character (6.04%) and a shorter triple bond character (36.74%), but a greater double bond character (57.2%), because the ionic part of the NRT bond order (0.53) is shorter than its covalent part (1.71). The same theoretical observations are also seen for the other two differently substituted R'B=AsR' compounds, as shown in **Table** 7, and in the data for the other $R'E_{13} \equiv BiR'$ compounds that is shown in Tables 9 (Al), 11 (Ga), 13 (In), and 15 (Tl). These computational data demonstrate that these $R'E_{13} = AsR'$ molecules have a weak $E_{13} = As$ triple bond.



Scheme 1. Three bulky ligands: SiMe(Si*t*Bu₃)₂, Si*i*PrDis₂, and N-heterocyclic carbine.

R′	SiMe(SitBu ₃) ₂	SiiPrDis ₂	NHC
B≡As (Å)	1.837	1.821	1.819
∠R′–B–As (°)	177.2	172.9	174.5
$\angle B$ –As–R' (°)	128.2	121.6	111.2
$\angle R'$ –B–As– R' (°)	179.6	177.4	171.5
$Q_{\scriptscriptstyle m B}{}^{\scriptscriptstyle 1}$	-0.280	-0.397	-0.205
$Q_{\rm As}^{2}$	-0.228	-0.134	0.061
ΔE_{ST} (kcal mol ⁻¹) ³	94.42	75.22	83.64
Wiberg BO ⁴	2.327	2.395	2.254

1 The natural charge density on the central B atom.

2 The natural charge density on the central As atom.

3 BE = E (triplet state for B-R') + E (triplet state for As-R') – E (singlet state for R'B=AsR').

4 The Wiberg bond index (WBI) for the B-As bond.

 $5 \Delta H_1 = E (:B=AsR'_2) - E (R'B=AsR');$ see Scheme 2.

6 ΔH_2 = E (R'₂B=As:) − E (R'B≡AsR'); see Scheme 2.

Table 6. The geometrical parameters, natural charge densities (Q_B and Q_{As}), Binding Energies (BE), the HOMO-LUMO Energy Gaps, the Wiberg Bond Index (WBI), and some reaction enthalpies for R'B=AsR' at the B3LYP/LANL2DZ+dp// RHF/3-21G* Level of Theory.

R′B≡AsR′	WBI	NBO analysis	NBO analysis			NRT analysis	
		Occupancy	hybridization	Polarization	total/covalent/ ionic	Resonance weight	
R' = SiMe(SitBu ₃) ₂	2.31	σ = 1.98	$\sigma: 0.6627 \text{ B} \\ (\text{sp}^{1.46}) + 0.7489 \\ \text{As (sp}^{1.07})$	43.91% (B) 56.09% (As)	2.35/1.66/0.69	B–As: 5.68% B=As: 60.70% B≡As: 33.62%	
		π = 1.94	$\pi: 0.5941 \text{ B} \ (\text{sp}^{1.00}) + 0.8044 \ \text{As} \ (\text{sp}^{99.99})$	35.29% (B) 64.71% (As)			
R' = Si <i>i</i> PrDis2 2.27	2.27	2.27 σ = 1.98	$\sigma: 0.6630 \text{ B} \ (\text{sp}^{1.54}) + 0.7486 \ \text{As} \ (\text{sp}^{1.22})$	43.96% (B) 56.04% (As)	2.24/1.71/0.53	B–As: 6.04% B=As : 57.2% B=As : 36.74%	
		π = 1.94	π : 0.5880 B (sp ^{99.99}) + 0.8089 As (sp ^{99.99})	34.58% (B) 65.42% (As)			
R' = NHC	2.26	σ = 1.98	$\sigma: 0.6918 \text{ B} \ (\text{sp}^{0.90}) + 0.7221 \ \text{As} \ (\text{sp}^{2.66})$	47.86% (B) 52.14% (As)	2.23/1.52/0.71	B–As : 7.05% B=As : 69.13% B≡As : 23.82%	
		π = 1.94	π : 0.5899 B (sp ^{99.99}) + 0.8075 As (sp ^{99.99})	34.80% (B) 65.20% (As)			

Table 7. Selected results for the natural bond orbital (NBO) and natural resonance theory (NRT) analyses at the B3LYP/ LANL2DZ+dp level of theory for R'B=AsR' compounds that have large substituents.

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R′	SiMe(SitBu ₃) ₂	SiiPrDis ₂	NHC
Al≡As (Å)	2.285	2.257	2.307
∠R′–Al–As (°)	179.4	176.0	174.5
\angle Al-As-R' (°)	116.4	118.7	113.0
$\angle R'$ –Al–As–R' (°)	176.1	170.6	176.4
	0.3771	0.3120	0.4392
$Q_{\rm As}^{2}$	-0.5579	-0.4907	-0.3144
ΔE_{ST} (kcal mol ⁻¹) ³	44.64	54.23	34.53
Wiberg BO ⁴	2.171	2.184	2.185

1 The natural charge density on the central Al atom.

2 The natural charge density on the central As atom.

3 BE = E(triplet state for Al–R') + E(triplet state for As–R') – E(singlet state for R'Al=AsR').

4 The Wiberg bond index (WBI) for the Al–As bond.

5 $\Delta H_1 = E(:Al=AsR'_2) - E(R'Al=AsR')$; see Scheme 2.

6 $\Delta H_2 = E(R'_2Al=As:) - E(R'Al=AsR');$ see Scheme 2.

Table 8. The geometrical parameters, natural charge densities (Q_{Al} and Q_{As}), binding energies (BE), the HOMO-LUMO energy gaps, the Wiberg Bond Index (WBI), and some reaction enthalpies for R'Al=AsR' at the B3LYP/LANL2DZ+dp// RHF/3-21G* level of theory.

R'Al=AsR'	WBI	NBO analy	NBO analysis			NRT analysis	
		Occupancy	hybridization	Polarization	total/covalent/ ionic	Resonance weight	
$R' = SiMe(SitBu_3)_2$	2.21	σ = 1.92	$ \begin{aligned} \sigma &: 0.5080 \ Al \\ (sp^{1.59}) &+ 0.8614 \\ As \ (sp^{1.14}) \end{aligned} $	25.81% (Al) 74.19% (As)	2.24/1.66/0.58	Al-As : 6.51% Al=As : 70.32% Al=As : 23.17%	
		π = 1.92	$\pi: 0.4437 \text{ Al} \ (\text{sp}^{99.99}) + 0.8962 \ \text{As} \ (\text{sp}^{99.99})$	19.69% (Al) 80.31% (As)			
R' = Si <i>i</i> PrDis2	2.29	σ = 1.92	$\sigma: 0.4956 \text{ Al} \\ (sp^{1.84}) + 0.8685 \\ \text{As} (sp^{1.06})$	24.57% (Al) 75.43% (As)	2.27/1.73/0.54	Al–As: 4.52% Al≡As : 57.55% Al≡As : 37.93%	
		π = 1.91	π : 0.4383 Al (sp ^{99.99}) + 0.8988 As (sp ^{99.99})	19.21% (Al) 80.79% (As)			
R' = NHC	2.36	σ = 1.87	σ : 0.5834 Al (sp ^{0.99}) + 0.8122 As (sp ^{10.87})	34.04% (Al) 65.96% (As)	2.30/1.59/0.71	Al-As : 6.61% Al=As : 74.90% Al=As : 18.49%	
		π = 1.94	$\pi: 0.4408 \text{ Al}$ (sp ^{90.78}) + 0.8976 As (sp ^{99.99})	19.43% (Al) 80.57% (As)			

Table 9. Selected results for the natural bond orbital (NBO) and natural resonance theory (NRT) analyses at the B3LYP/ LANL2DZ+dp level of theory for R'Al=AsR' compounds that have large substituents.

R'	SiMe(SitBu ₃) ₂	SiiPrDis ₂	NHC
Ga≡As (Å)	2.274	2.252	2.316
∠R′–Ga–As (°)	178.8	178.1	171.9
\angle Ga–As–R' (°)	119.3	122.8	114.5
$\angle R'$ –Ga–As–R' (°)	176.6	171.0	176.5
Q_{Ga}^{1}	0.1760	0.09195	0.2133
$Q_{\rm As}^{2}$	-0.4683	-0.3978	-0.2257
ΔE _{ST} (kcal mol ⁻¹) ³	40.67	31.52	33.97
Wiberg BO ⁴	2.125	2.174	2.154

1 The natural charge density on the central Ga atom.

2 The natural charge density on the central As atom.

3 BE = E(triplet state for Ga-R') + E(triplet state for As-R') – E(singlet state for R'Ga=AsR').

4 The Wiberg bond index (WBI) for the Ga–As bond.

5 $\Delta H_1 = E(:Ga=AsR'_2) - E(R'Ga=AsR');$ see Scheme 2.

6 $\Delta H_2 = E(R'_2Ga=As:) - E(R'Ga=AsR');$ see Scheme 2.

Table 10. The geometrical parameters, natural charge densities (Q_{Ga} and Q_{As}), binding energies (BE), the HOMO-LUMO energy gaps, the Wiberg Bond Index (WBI), and some reaction enthalpies for R'Ga=AsR' at the B3LYP/LANL2DZ+dp// RHF/3-21G* level of theory.

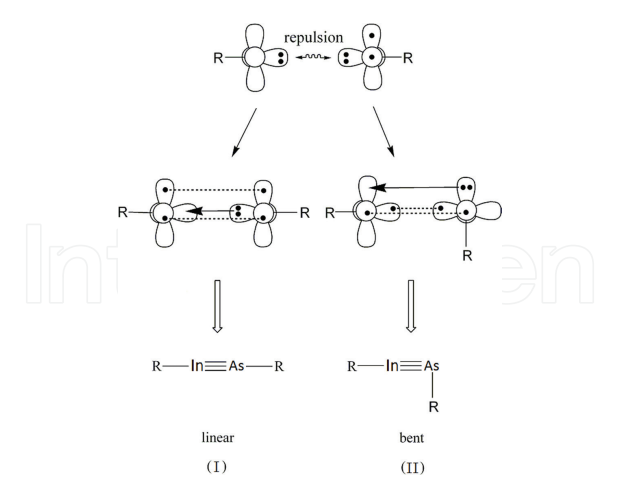


Figure 6. The bonding models (I) and (II) for the triply bonded RE_{13} =AsR molecule.

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R′Ga≡AsR′	WBI	NBO analysis	5		NRT analysis	
		Occupancy	Hybridization	Polarization	total/covalent/ ionic	Resonance weight
R' = SiMe(SitBu ₃) ₂	2.19	σ = 1.90	σ : 0.5320 Ga (sp ^{1.52}) + 0.8468 As (sp ^{1.32})	28.30% (Ga) 71.70% (As)	2.27/1.62/0.65	Ga–As : 4.72% Ga=As : 56.61%
		π = 1.93	π : 0.4467 Ga (sp ^{99.99}) + 0.8947 As (sp ^{99.99})	19.95% (Ga) 80.05% (As)		Ga=As : 38.67%
R' = Si <i>i</i> PrDis2	R' = SiiPrDis2 2.25	σ = 1.91	$\sigma: 0.5386 \text{ Ga}$ (sp ^{1.49}) + 0.8426 As (sp ^{1.46})	29.01% (Ga) 70.99% (As)	2.31/1.64/0.67	Ga=As : 7.03% Ga=As : 68.13% Ga=As : 24.84%
		π = 1.92	π : 0.4392 Ga (sp ^{99.99}) + 0.8984 As (sp ^{99.99})	19.29% (Ga) 80.71% (As)		
R' = NHC	2.33	σ = 1.85	σ : 0.6076 Ga (sp ^{0.98}) + 0.7942 As (sp ^{12.06})	36.92% (Ga) 63.08% (As)	2.14/1.71/0.43	Ga–As : 7.12% Ga=As : 75.34% Ga=As : 17.54%
	π	<i>π</i> = 1.93	π : 0.4370 Ga (sp ^{82.50}) + 0.8995 As (sp ^{99.99})	19.09% (Ga) 80.91% (As)		

Table 11. Selected results for the natural bond orbital (NBO) and natural resonance theory (NRT) analyses at the B3LYP/ LANL2DZ+dp level of theory for R'Ga=AsR' compounds that have large substituents.

R′	SiMe(SitBu ₃) ₂	SiiPrDis ₂	NHC
In≡As (Å)	2.446	2.430	2.482
∠R'–In–As (°)	155.9	168.4	171.3
\angle In–As–R' (°)	127.8	120.3	110.8
$\angle R'$ –In–As–R' (°)	173.9	162.0	168.3
Q_{\ln}^{1}	0.874	0.880	1.021
$Q_{\rm As}^{2}$	-0.783	-0.822	-0.359
ΔE _{ST} (kcal mol ⁻¹) ³	41.5	45.2	35.7
Wiberg BO ⁴	2.174	2.271	2.141

1 The natural charge density on the central In atom.

2 The natural charge density on the central As atom.

3 BE = E(triplet state for In-R') + E(triplet state for As-R') - E(singlet state for R'In=AsR').

4 The Wiberg bond index (WBI) for the In-As bond.

 $5 \Delta H_1 = E(:In=AsR'_2) - E(R'In=AsR');$ see Scheme 2.

6 ΔH_2 = E(R'_In=As:) – E(R'In=AsR'); see Scheme 2.

Table 12. The geometrical parameters, natural charge densities (Q_{In} and Q_{As}), Binding Energies (BE), the HOMO-LUMO Energy Gaps, the Wiberg Bond Index (WBI), and some reaction enthalpies for R'In=AsR' at the B3LYP/LANL2DZ+dp// RHF/3-21G* Level of Theory.

R'In≡AsR'	WBI	NBO analysis			NRT analysis	
		Occupancy	Hybridization	Polarization	total/covalent/ ionic	Resonance weight
R' = SiMe(SitBu ₃) ₂	1.50	σ = 1.87	σ : 0.4940 In (sp ^{1.58}) + 0.8695 As (sp ^{1.28})	24.41% (In) 75.59% (As)	2.31/1.55/0.76	In-As : 5.78% In=As :55.2 % In=As : 39.0%
		π = 1.85	π : 0.4411 In (sp ^{2.80}) + 0.8975 As (sp ^{4.33})	19.45% (In) 80.55% (As)		
R' = Si <i>i</i> PrDis2	1.48	σ = 1.87	σ : 0.4854 In (sp ^{1.71}) + 0.8743 As (sp ^{1.26})	23.56% (In) 76.44% (As)	2.18/1.62/0.56	In-As : 6.01% In=As : 56.29% In=As : 37.70%
		π = 1.83	π : 0.3873 In (sp ^{99.99}) + 0.9220 As (sp ^{1.00})	15.00% (In) 85.00% (As)		
R' = NHC	1.33	σ = 1.80	σ : 0.5709 In (sp ^{1.07}) + 0.8210 As (sp ^{8.66})	32.60% (In) 67.40% (As)	2.21/1.48/0.73	In–As : 7.72% In=As : 78.30% In=As : 13.98%
		π = 1.94	$\pi : 0.4805$ In (sp ^{37.19}) + 0.8770 As (sp ^{14.95})	23.09% (In) 76.91% (As)		

Table 13. Selected results for the natural bond orbital (NBO) and natural resonance theory (NRT) analyses at the B3LYP/ LANL2DZ+dp level of theory for R'In≡AsR' compounds that have large substituents.

R′	SiMe(SitBu ₃) ₂	SiiPrDis ₂	NHC
Tl≡As (Å)	2.615	2.565	2.653
∠R′–Tl–As (°)	176.9	177.6	178.7
\angle Tl–As–R' (°)	127.7	121.8	108.0
∠R′–T1–As–R′ (°)	172.2	170.4	175.2
\mathbf{Q}_{Π}^{-1}	0.310	0.246	0.262
$Q_{\rm As}^{2}$	-0.462	-0.440	-0.313
ΔE _{ST} (kcal mol ⁻¹) ³	45.07	32.71	34.83
Wiberg BO ⁴	2.157	2.214	2.209

1 The natural charge density on the central Tl atom.

2 The natural charge density on the central As atom.

3 BE = E(triplet state for Tl–R') + E(triplet state for As–R') – E(singlet state for R'Tl=AsR').

4 The Wiberg bond index (WBI) for the Tl–As bond.

5 $\Delta H_1 = E(:TI=AsR'_2) - E(R'TI=AsR');$ see Scheme 2.

 $6 \Delta H_2 = E(R'_2TI=As:) - E(R'TI=AsR');$ see Scheme 2.

Table 14. The geometrical parameters, natural charge densities (Q_{Π} and Q_{As}), Binding Energies (BE), the HOMO-LUMO Energy Gaps, the Wiberg Bond Index (WBI), and some reaction enthalpies for R'Tl=AsR' at the B3LYP/LANL2DZ+dp// RHF/3-21G* Level of Theory.

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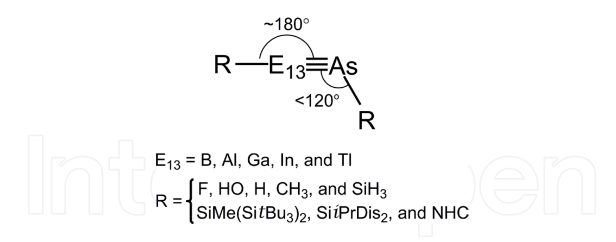
R′Tl≡AsR′	WBI	NBO analysis			NRT analysis	
		Occupancy	Hybridization	Polarization	total/covalent/ ionic	Resonance weight
$R' = SiMe(SitBu_3)_2$	2.15	σ = 1.74	σ : 0.5404 Tl (sp ^{1.51}) + 0.8414 As (sp ^{2.20})	29.20% (Tl) 70.80% (As)	2.24/1.68/0.56	TI-As : 6.11% TI=As : 57.27% TI=As : 36.62%
		π = 1.79	$\pi: 0.3968 \text{ Tl} \ (\text{sp}^{4.25}) + 0.9179 \ \text{As} \ (\text{sp}^{1.51})$	15.74% (Tl) 84.26% (As)		
R' = Si <i>i</i> PrDis2	2.21	σ = 1.90	$\sigma: 0.3627$ Tl (sp ^{38,20}) + 0.9318 As (sp ^{1.44})	13.16% (Tl) 86.84% (As)	2.16/1.73/0.43	TI–As : 7.01% TI=As : 66.48% TI=As : 26.51%
		π = 1.94	$\pi: 0.3315$ Tl (sp ^{99.99}) + 0.9435 As (sp ^{1.00})	10.99% (Tl) 89.01% (As)		
R' = NHC	2.11	σ = 1.97	σ : 0.7814 Tl (sp ^{0.07}) + 0.6240 As (sp ^{52.63})	61.06% (Tl) 38.94% (As)	2.14/1.71/0.43	Tl-As : 6.71% Tl=As : 75.51% Tl=As : 17.78%
		π = 1.97	$\pi : 0.4726$ Tl (sp ^{57.71}) + 0.8812 As (sp ^{17.96})	22.34% (Tl) 77.66% (As)		

Table 15. Selected results for the natural bond orbital (NBO) and natural resonance theory (NRT) analyses at the B3LYP/ LANL2DZ+dp level of theory for R'Tl=AsR' compounds that have large substituents.

4. Conclusion

This study of the effect of substituents on the possibility of the existence of triply bonded RE_{13} =AsR allows the following conclusions to be drawn (**Scheme 2**):

- 1. The theoretical observations provide strong evidence that bonding mode (B) is dominant in the triply bonded RE₁₃≡BiR species, because their structures are bent due to electron transfer (denoted by arrows in **Figure 1**) and the relativistic effect, which increases stability.
- 2. The theoretical evidence shows that both the electronic and the steric effects of substituents are crucial to rendering the E_{13} =As triple bond synthetically accessible. However, this theoretical study shows that these E_{13} =As triple bonds are weak. They are not as strong as the traditional C=C triple bond. The results of this theoretical study show that triply bonded R' E_{13} =AsR' molecules that feature bulky substituents are more stable because bulky substituents not only protect the central E_{13} =As triple bond because there is large steric hindrance but also prohibit polymerization reactions.



Scheme 2. The predicted structure for the triply bonded $RE_{13} \equiv AsR$ molecules based on the present theoretical computations.

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