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# Triple Bonds between Bismuth and Group 13 Elements: Theoretical Designs and Characterization

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#### Abstract

The effect of substitution on the potential energy surfaces of RE<sub>13</sub>=BiR (E<sub>13</sub> = B, Al, Ga, In, and Tl; R = F, OH, H, CH<sub>3</sub>, SiH<sub>3</sub>, Tbt, Ar\*, SiMe(SitBu<sub>3</sub>)<sub>2</sub>, and SitPrDis<sub>2</sub>) is investigated using density functional theories (M06-2X/Def2-TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp). The theoretical results suggest that all of the triply bonded RE<sub>13</sub>=BiR molecules prefer to adopt a bent geometry (i.e.,  $\angle$ RE<sub>13</sub>Bi  $\approx$  180° and  $\angle$ E<sub>13</sub>BiR  $\approx$  90°), which agrees well with the bonding model (model (B)). It is also demonstrated that the smaller groups, such as R = F, OH, H, CH<sub>3</sub>, and SiH<sub>3</sub>, neither kinetically nor thermodynamically stabilize the triply bonded RE<sub>13</sub>=BiR compounds, except for the case of H<sub>3</sub>SiB=BiSiH<sub>3</sub>. Nevertheless, the triply bonded R'E<sub>13</sub>=BiR' molecules that feature bulkier substituents (R' = Tbt, Ar\*, SiMe(SitBu<sub>3</sub>)<sub>2</sub>, and SitPrDis<sub>2</sub>) are found to have the global minimum on the singlet potential energy surface and are both kinetically and thermodynamically stable. In other words, both the electronic and the steric effects of bulkier substituent groups play an important role in making triply bonded RE<sub>13</sub>=BiR (Group 13–Group 15) species synthetically accessible and isolable in a stable form.

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

## 1. Introduction

Triply bonded molecules are of great interest in structural and synthetic inorganic chemistry as well as in fundamental science. Molecules that have triple bonds, however, pose a more difficult challenge than analogous doubly bonded molecules from a synthetic viewpoint [1–8]. Acetylene is one of the most commonly triply bonded molecules in traditional organic chemistry. Thanks to Kira, Power, Sekiguchi, Tokitoh, Wiberg and many coworkers, the stable

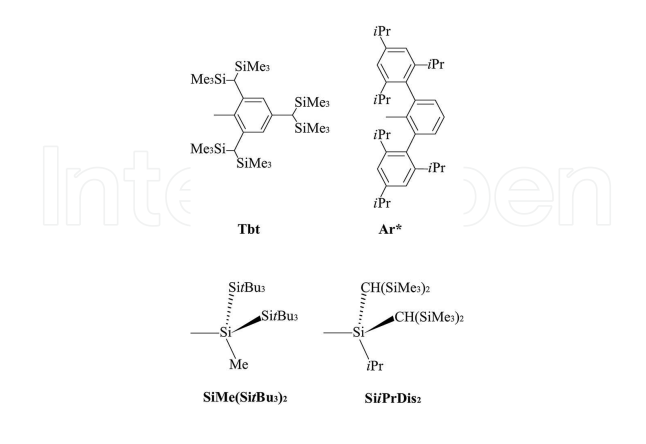


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homonuclear alkyne analogues of all of the heavier group 14 elements have now been isolated and characterized [9–19]. Recently, heteronuclear ethyne-like molecules that possess C=Ge [20, 21], C=Sn [22], and C=Pb [23] triple bonds have also been theoretically predicted and have been published elsewhere.

Nevertheless, to the authors' best knowledge, neither experimental nor theoretical studies have been performed on acetylene-like compounds that feature an  $E_{13}$ =Bi (E = B, Al, Ga, In, and Tl) triple bond. It is surprising how little is known about the stability and molecular properties of  $E_{13}$ =Bi, considering the importance of bismuth compounds [24] that contain group 13 elements in inorganic chemistry [25–35] and material chemistry [36–45].

The aim of this study is to theoretically determine the existence and relative stability of  $RE_{13}$ =BiR triply bonded molecules, which can be synthesized as stable compounds when they are properly substituted. For the first time, the structures of  $RE_{13}$ =BiR with various substituents are reported. That is, theoretical calculations of  $RE_{13}$ =BiR are performed, using both smaller ligands (such as, R = F, OH, H, CH<sub>3</sub>, and SiH<sub>3</sub>) and larger ligands with bulky aryl and silyl groups (i.e., R' = Tbt, Ar\*, SiMe(SitBu<sub>3</sub>)<sub>2</sub>, and SitPrDis<sub>2</sub>; Dis = CH(SiMe<sub>3</sub>)<sub>2</sub>; **Scheme 1**) [46–51]. As a result, the effect of substituents on these bismuth-group-13-element triple bonds is systematically investigated using density functional theory (DFT) calculations. It is expected that the theoretical interpretations of the effect of substituents, presented in this work, will help in the experimental preparation of the many precursors of  $RE_{13}$ =BiR.



# 2. Theoretical methods

Geometries were fully optimized using hybrid density functional theory at M06-2X, B3PW91, and B3LYP levels, using the Gaussian 09 program package [52]. It has been reported that M06-2X is proven to have excellent performance for main group chemistry [53]. In both the B3LYP and B3PW91 calculations, Becke's three-parameter nonlocal exchange functional (B3) [54, 55] is used, together with the exact (Hartree-Fock) exchange functional, in conjunction with the nonlocal correlation functional of Lee, Yang, Parr (LYP) [56] and Perdew and Wang (PW91) [57]. Therefore, the geometries of all of the stationary points were fully optimized at the M06-2X, B3PW91, and B3LYP levels of theory. For comparison, the geometries and energetics of the stationary points on the potential energy surface were calculated using the M06-2X, B3PW91, and B3LYP methods, in conjunction with the Def2-TZVP [58] and LANL2DZ+dp [59–62] basis sets. Consequently, these DFT calculations are denoted as M06-2X/Def2-TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp, respectively.

The spin-unrestricted (UM06-2X, UB3PW91, and UB3LYP) formalisms are used for the openshell (triplet) species. The  $\langle S^2 \rangle$  expectation values for the triplet state for the calculated species all have an ideal value (2.00), after spin annihilation, so their geometries and energetics are reliable for this study. Frequency calculations were performed on all structures, in order to confirm that the reactants and products have no imaginary frequencies, and that the transition states possess only one imaginary frequency. Thermodynamic corrections to 298 K, heat capacity corrections, and entropy corrections ( $\Delta$ S) are applied at the three DFT levels. Therefore, the relative free energy ( $\Delta$ G) at 298 K is also calculated at the same levels of theory.

Sequential conformation analyses were performed for each stationary point, for species containing bulky ligands (R' = Tbt, Ar\*, SiMe(SitBu<sub>3</sub>)<sub>2</sub>, and SiiPrDis<sub>2</sub>) using Hartree-Fock calculations (RHF/3-21G\*). The TbtE<sub>13</sub>=Bi=Tbt, Ar\*=E<sub>13</sub>=Bi=Ar\*, SiMe(SitBu<sub>3</sub>)<sub>2</sub>=E<sub>13</sub>=Bi=SiMe (SitBu<sub>3</sub>)<sub>2</sub>, and SiiPrDis<sub>2</sub>=E<sub>13</sub>=Bi=SiiPrDis<sub>2</sub> (E = B, Al, Ga, In, and Tl) are used as model reactants in this work. It is known that the Hartree-Fock level of theory is insufficient for even a qualitative description of the chemical potential energy surface, so these stationary points were then further calculated at the B3LYP/LANL2DZ+dp level, using the OPT=READFC keyword with a tight convergence option (maximum gradient convergence tolerance =  $5.0 \times 10^{-5}$  hartree/ bohr). Because of the limitations of the available CPU time and memory size, frequencies were not calculated for the triply bonded R'E<sub>13</sub>=BiR' systems with bulky ligands (R') at the B3LYP/LANL2DZ+dp level of theory. As a result, the zero-point energies and the Gibbs free energies for B3LYP/LANL2DZ+dp cannot be applied to these systems.

## 3. Results and discussion

### 3.1. Theoretical models for RE<sub>13</sub>≡BiR

In order to understand the bonding interactions in the  $R=E_{13}=Bi=R$  molecule,  $R=E_{13}=Bi=R$  is divided into one  $E_{13}=R$  and one Bi=R fragment. The theoretical calculations for these two

fragments indicate that the ground states of  $E_{13}$ =R and Bi=R are singlet and triplet states, respectively (vide infra). Therefore, there are two possible interaction modes (A and B) between the  $E_{13}$ =R and Bi=R moieties in the formation of the triply bonded R= $E_{13}$ =Bi=R species, as schematically illustrated in **Figure 1**. In model (A), both E=R and Bi=R units exist as triplet monomers. In this way, the combination between the group 13 element and bismuth can be considered as a triple bond, since it consists of 2  $\pi$  bonds and 1 donor-acceptor  $\sigma$  bond, for these 2 triplet fragments. As a result, this bonding model allows a linear structure, as shown in **Figure 1(A)**. In model (B), both  $E_{13}$ =R and Bi=R units still exist as triplets, so this bonding scheme contains one  $\sigma$  bond and one p- $\pi$  bond (indicated by two dashed lines), plus one donor-acceptor  $\pi$ -bond because of coupling between the lone pair in Bi=R and the empty p orbital at the  $E_{13}$  atom (indicated by the arrow). Accordingly, this bonding pattern results in a bent structure, as shown in **Figure 1(B)**. The importance of the RE<sub>13</sub>=BiR donor-acceptor interaction is emphasized, as it is essential for the stabilization of the nonlinear structure. These analyses are used to explain the geometrical structures of triply bonded RE<sub>13</sub>=BiR species in the following sections.

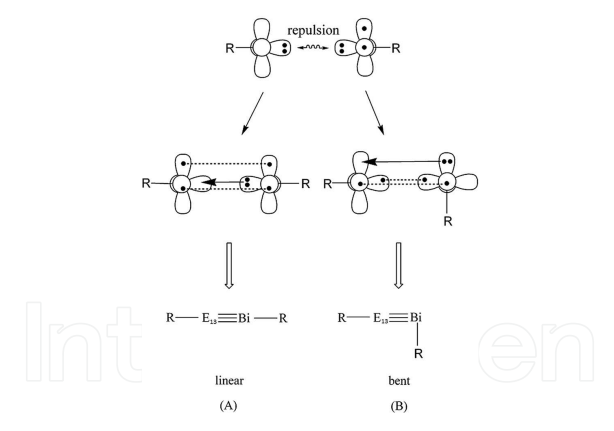


Figure 1. Two interaction models, A and B, in forming triply bonded RE<sub>13</sub>=BiR species.

#### 3.2. Small ligands on substituted RE<sub>13</sub>≡BiR

Small ligands, such as R = F, OH, H, CH<sub>3</sub>, and SiH<sub>3</sub>, are firstly chosen to study the geometries of the RE<sub>13</sub>=BiR (E<sub>13</sub> = B, Al, Ga, In, and Tl) species. As mentioned in the Introduction, neither experimental nor theoretical results for the triply bonded RE<sub>13</sub>=BiR species are available to allow a definitive comparison. As a result, three DFT methods were used (i.e., M06-2X/Def2-

TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp) to examine their molecular properties. The selected geometrical parameters, natural charge densities ( $Q_{E13}$  and  $Q_{Bi}$ ), binding energies (BE), and Wiberg bond order (BO) [63, 64] are shown in **Table 1** (RB=BiR), **Table 2** (RAl=BiR), **Table 3** (RGa=BiR), **Table 4** (RIn=BiR), and **Table 5** (RTl=BiR).

R	F	ОН	H	CH <sub>3</sub>	SiH <sub>3</sub>
B=Bi (Å)	2.218	2.202	2.091	2.141	2.075
	-2.210	-2.199	-2.083	-2.137	-2.084
	[2.196]	[2.196]	[2.083]	[2.140]	[2.085]
∠R−B−Bi (°)	177.6	176	163.6	176.9	170.9
	-178.8	-175.9	-163.6	-176.9	-169.4
	[178.1]	[176.1]	[163.9]	[174.7]	[171.9]
$\angle B-Bi-R$ (°)	80.89	91.96	34.78	90.62	58.89
	-88.58	-90.21	-34.83	-90.26	-58.47
	[87.52]	[89.53]	[38.88]	[99.39]	[59.00]
$\angle R - B - Bi - R$ (°)	179.8	77.43	180	173.3	179.7
	-179.1	-75.70	-180.0	-173.5	-179.5
	[179.5]	[76.80]	[180.0]	[179.9]	[180.0]
$Q_{\rm B}$ <sup>(1)</sup>	0.1096	-0.0543	-0.3684	-0.3023	-0.5098
	-0.124	(-0.0372)	(-0.2262)	(-0.1390)	(-0.4100)
	[0.2303]	[0.0511]	[-0.1623]	[-0.01010]	[-0.4101]
Q <sub>Bi</sub> <sup>(2)</sup>	0.4759	0.3569	0.2189	0.1881	0.1784
	-0.4975	-0.3431	-0.1331	-0.1602	-0.103
	[0.4983]	[0.3552]	[0.1870]	[0.1430]	[0.1520]

Several important conclusions can be found in **Tables 1–5**, which are shown as follows:

Notes: (1) The natural charge density on the central boron atom. (2) The natural charge density on the central bismuth atom. (3) BE = E (triplet state of R=B) + E (triplet state of R=Bi) – E(RB=BiR). (4) Wiberg bond orders for the B=Bi bonds, see Ref. [18].

**Table 1.** Selected geometrical parameters, natural charge densities ( $Q_B$  and  $Q_{Bi}$ ), binding energies (BE), and Wiberg bond orders (BO) of RB=BiR at the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP (in round bracket), and B3LYP/LANL2DZ+dp (in square bracket) levels.

R	F	ОН	Н	CH <sub>3</sub>	SiH <sub>3</sub>
Al≡Bi (Å)	2.604	2.606	2.439	2.529	2.501
	(2.601)	(2.601)	(2.463)	(2.539)	(2.532)
	[2.621]	[2.624]	[2.483]	[2.561]	[2.542]
∠R−Al−Bi (°)	176.8	173.4	170.7	177.7	172.5
	(175.4)	(172.9)	(168.1)	(177.6)	(170.4)
	[177.3]	[174.2]	[166.9]	[177.1]	[174.5]

R	F	ОН	Н	CH <sub>3</sub>	SiH <sub>3</sub>
∠Al−Bi−R (°)	83.16	84.20	48.09	92.52	62.57
	(84.59)	(85.35)	(49.85)	(92.93)	(61.82)
	[87.00]	[88.32]	[51.00]	[93.77]	[64.03]
$\angle R - Al - Bi - R$ (°)	180.0	177.3	180.0	179.9	179.9
	(180.0)	(176.0)	(180.0)	(179.6)	(179.8)
	[180.0]	178.0]	[180.0]	[179.6]	[180.0]
$Q_{\rm Al}$ <sup>(1)</sup>	0.5031	0.3942	0.1493	0.2692	0.1841
	(0.4904)	(0.3918)	(0.1417)	(0.2544)	(0.2145)
	[0.6664]	[0.4315]	[0.3786]	[0.2414]	[0.1517]
Q <sub>Bi</sub> <sup>(2)</sup>	0.3947	0.2709	-0.05788	0.03761	-0.1384
	(0.3196)	(0.1834)	(-0.04954)	(0.02100)	(-0.07446)
	[0.3044]	[0.1982]	[0.03410]	[-0.05262]	[-0.1074]
BE (kcal mol <sup>-1</sup> ) $^{(3)}$	22.61	20.28	50.55	38.69	53.41
	(30.36)	(31.77)	(85.64)	(63.54)	(57.96)
	[25.47]	[20.51]	[53.65]	[42.77]	[53.47]
Wiberg BO <sup>(4)</sup>	1.393	1.403	1.746	1.634	1.602
	(1.509)	(1.511)	(1.798)	(1.690)	(1.615)
	[1.521]	[1.516]	[1.787]	[1.706]	[1.653]

Notes: (1) The natural charge density on the central aluminum atom. (2) The natural charge density on the central bismuth atom. (3) BE = E (triplet state of R=Al) + E (triplet state of R=Bi) – E(RAl=BiR). (4) Wiberg bond orders for the Al=Bi bonds, see Refs. [63, 64].

**Table 2.** Selected geometrical parameters, natural charge densities ( $Q_{Al}$  and  $Q_{Bi}$ ), binding energies (BE), and Wiberg bond orders (BO) of RAl=BiR at the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP (in round bracket), and B3LYP/LANL2DZ +dp (in square bracket) levels.

R	F	ОН	Н	CH <sub>3</sub>	SiH <sub>3</sub>
Ga≡Bi (Å)	2.639	2.625	2.463	2.543	2.512
	(2.602)	(2.621)	(2.465)	(2.524)	(2.510)
	[2.632]	[2.629]	[2.487]	[2.550]	[2.520]
∠R-Ga-Bi (°)	179.7	175.0	166.5	178.9	178.7
	(178.3)	(173.3)	(166.2)	(177.8)	(177.6)
	[177.3]	[175.5]	[167.0]	[177.3]	[177.0]
$\angle Ga - Bi - R(^{\circ})$	86.32	86.85	52.56	91.27	65.56
	(88.49)	(88.52)	(56.24)	(92.86)	(66.28)
	[88.18]	[90.75]	[59.49]	[93.37]	[69.82]
$\angle R-Ga-Bi-R$ (°)	179.5	157.1	180.0	179.2	175.8
	(180.0)	(159.8)	(180.0)	(178.8)	(179.9)

Triple Bonds between Bismuth and Group 13 Elements: Theoretical Designs and Characterization 77 http://dx.doi.org/10.5772/67220

R	F	OH	Н	CH <sub>3</sub>	SiH <sub>3</sub>
	[180.0]	[158.0]	[180.0]	[179.3]	[180.0]
$Q_{Ga}^{(1)}$	0.6246	0.5224	0.2127	0.2266	0.1845
	(0.5012)	(0.3464)	(0.1135)	(0.2356)	(0.1507)
	[0.5700]	[0.3813]	[0.3031]	[0.1984]	[0.07925]
Q <sub>Bi</sub> <sup>(2)</sup>	0.3696	0.2435	-0.1367	0.03356	-0.2002
	(0.3574)	(0.2743)	(-0.04212)	(0.05503)	(-0.05512)
	[0.4000]	[0.2395]	[0.06523]	[-0.01620]	[-0.08931]
BE (kcal mol <sup><math>-1</math></sup> ) <sup>(3)</sup>	18.03	16.41	45.28	46.24	46.10
	(22.92)	(26.32)	(79.05)	(60.77)	(49.96)
	[20.80]	[16.61]	[49.49]	[40.12]	[48.61]
Wiberg BO <sup>(4)</sup>	1.286	1.335	1.718	1.578	1.653
	(1.382)	(1.393)	(1.787)	(1.633)	(1.646)
	[1.403]	[1.431]	[1.758]	[1.656]	[1.673]

Notes: (1) The natural charge density on the central gallium atom. (2) The natural charge density on the central bismuth atom. (3) BE = E (triplet state of R=Ga) + E (triplet state of R=Bi) – E(RGa=BiR). (4) Wiberg bond orders for the Ga=Bi bonds, see Refs. [63, 64].

**Table 3.** Selected geometrical parameters, natural charge densities ( $Q_{Ga}$  and  $Q_{Bi}$ ), binding energies (BE), and Wiberg bond orders (BO) of RGa=BiR at the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP (in round bracket), and B3LYP/LANL2DZ +dp (in square bracket) levels.

R	F	ОН	Н	CH <sub>3</sub>	SiH <sub>3</sub>
In≡Bi (Å)	2.804	2.790	2.659	2.696	2.667
	(2.802)	(2.700)	(2.691)	(2.719)	(2.692)
	[2.790]	[2.795]	[2.673]	[2.712]	[2.683]
∠R—In—Bi (°)	179.6	173.9	168.3	179.1	173.8
	(178.0)	(172.2)	(175.2)	(177.3)	(174.3)
	[177.0]	[174.5]	[174.1]	[177.6]	[174.7]
∠In−Bi−R (°)	84.16	85.14	67.00	92.20	70.37
	(87.83)	(89.71)	(77.82)	(94.83)	(75.05)
	[87.43]	[90.79]	[74.35]	[94.16]	[74.37]
$\angle R$ -In-Bi-R (°)	179.9	176.1	180.0	178.3	178.9
	(180.0)	(177.3)	(180.0)	(179.1)	(177.5)
	[180.0]	[176.5]	[180.0]	[179.4]	[179.8]
Q <sub>In</sub> <sup>(1)</sup>	0.7021	0.6352	0.3755	0.3297	0.3650
	(0.5692)	(0.4640)	(0.2561)	(0.3403)	(0.2872)
	[0.7571]	[0.5053]	[0.4055]	[0.3044]	[0.1756]
Q <sub>Bi</sub> <sup>(2)</sup>	0.3973	0.2452	-0.2474	-0.01187	-0.2599

R	F	ОН	Н	CH <sub>3</sub>	SiH <sub>3</sub>
	(0.4000)	(0.2511)	(-0.08703)	(0.04410)	(-0.08023)
	[0.3468]	[0.2141]	[0.02620]	[-0.05735]	[-0.1365]
BE (kcal mol <sup><math>-1</math></sup> ) <sup>(3)</sup>	14.66	13.17	39.19	36.87	39.28
	(15.06)	(12.88)	(42.01)	(35.00)	(40.94)
	[18.80]	[13.70]	[44.04]	[35.54]	[41.83]
Wiberg BO <sup>(4)</sup>	1.312	1.403	1.590	1.543	1.553
	(1.308)	(1.334)	(1.601)	(1.539)	(1.546)
	[1.323]	[1.336]	[1.615]	[1.548]	[1.549]

Notes: (1) The natural charge density on the central indium atom. (2) The natural charge density on the central bismuth atom. (3) BE = E (triplet state of R=In) + E (triplet state of R=Bi) – E(RIn=BiR). (4) Wiberg bond orders for the In=Bi bonds, see Refs. [63, 64].

**Table 4.** Selected geometrical parameters, natural charge densities ( $Q_{In}$  and  $Q_{Bi}$ ), binding energies (BE), and Wiberg bond orders (BO) of RIn=BiR at the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP (in round bracket), and B3LYP/LANL2DZ+dp (in square bracket) levels.

R	F	ОН	Н	CH <sub>3</sub>	SiH <sub>3</sub>
Tl≡Bi (Å)	2.859	2.843	2.713	2.742	2.707
	(2.812)	(2.803)	(2.698)	(2.725)	(2.705)
	[2.819]	[2.822]	[2.679]	[2.713]	[2.682]
∠R−Tl−Bi (°)	175.9	173.5	175.7	179.6	174.4
	(178.8)	(172.5)	(176.6)	(178.5)	(174.7)
	[177.2]	[174.4]	[176.3]	[178.5]	[175.9]
$\angle Tl-Bi-R$ (°)	81.34	86.72	78.87	91.91	76.03
	(87.73)	(89.82)	(79.00)	(93.54)	(76.50)
	[87.92]	[92.34]	[78.86]	[93.25]	[80.00]
$\angle R-Tl-Bi-R(^{\circ})$	180.0	143.7	180.0	172.1	178.9
	(179.9)	(132.9)	(179.9)	(178.6)	(179.9)
	[180.0]	[130.6]	[180.0]	[180.0]	[179.2]
$Q_{\mathrm{TI}}^{(1)}$	0.6481	0.5672	0.3014	0.4162	0.2665
	(0.6614)	(0.5879)	(0.2284)	(0.2746)	(0.3510)
	[0.7100]	[0.4812]	[0.3536]	[0.2734]	[0.1601]
$Q_{\mathrm{Bi}}$ <sup>(2)</sup>	0.4752	0.3214	-0.1836	0.008121	-0.2245
	(0.3615)	(0.2201)	(-0.05213)	(-0.1282)	(-0.09637)
	[0.3854]	[0.2455]	[0.04813]	[-0.03131]	[-0.1282]
BE (kcal mol <sup>-1</sup> ) <sup>(5)</sup>	7.90	6.61	30.74	26.94	30.56
	(2.98)	(8.17)	(48.12)	(35.39)	(29.13)

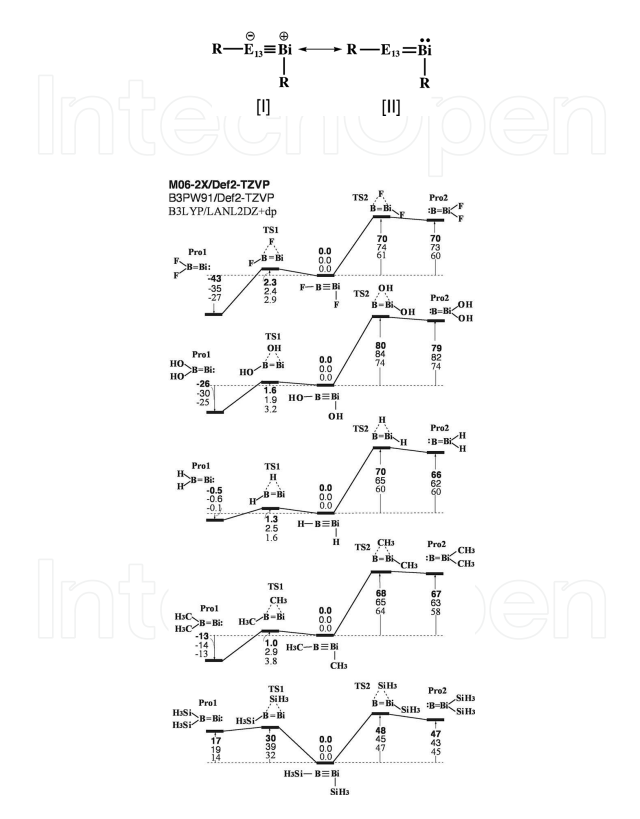
R	F	ОН	Н	CH <sub>3</sub>	SiH <sub>3</sub>
	[11.34]	[7.38]	[37.14]	[31.09]	[35.03]
Wiberg BO <sup>(6)</sup>	1.000	1.132	1.652	1.448	1.792
	(1.121)	(1.254)	(1.765)	(1.514)	(1.824)
	[1.112]	[1.212]	[1.756]	[1.568]	[1.652]

Notes: (1) The natural charge density on the central thallium atom. (2) The natural charge density on the central bismuth atom. (3) BE = E (triplet state of R=TI) + E (triplet state of R=Bi) – E(RTI=BiR). (4) Wiberg bond orders for the TI=Bi bonds, see Refs. [63, 64].

**Table 5.** Selected geometrical parameters, natural charge densities ( $Q_{TI}$  and  $Q_{Bi}$ ), binding energies (BE), and Wiberg bond orders (BO) of RTI=BiR at the M06-2X/Def2-TZVP, B3PW91/Def2-TZVP (in round bracket), and B3LYP/LANL2DZ+dp (in square bracket) levels.

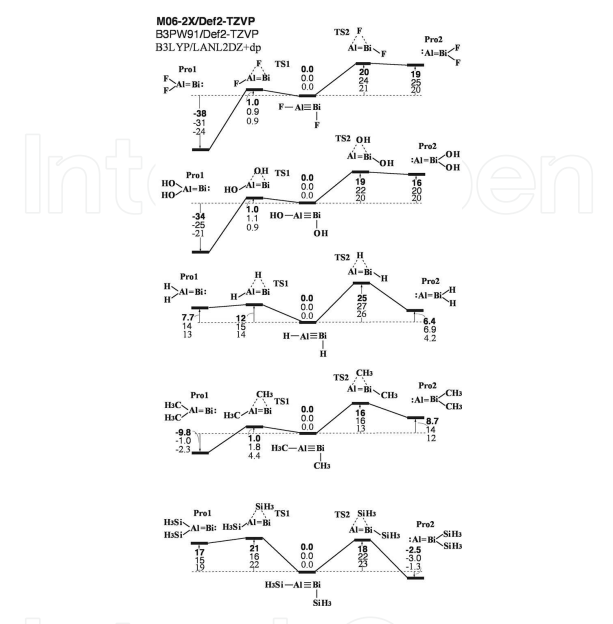
- As can be seen from Tables 1–5, the geometrical parameters of the RE<sub>13</sub>=BiR triple bond species are quite analogous at the three levels employed. For instance, the predicted triple bond length for small ligands on RB=BiR is 2.075–2.218 Å (M06-2X/Def2-TZVP), 2.083–2.210 Å (B3PW91/Def2-TZVP), and 2.083–2.196 Å (B3LYP/LANL2DZ+dp). For RAI=BiR, the AI=Bi triple bond length for small ligands is predicted to be 2.439–2.606 Å (M06-2X/Def2-TZVP), 2.463–2.601 Å (B3PW91/Def2-TZVP), and 2.483–2.624 Å (B3LYP/LANL2DZ+dp). For RGa=BiR, the Ga=Bi triple bond length for small ligands is predicted to be 2.463–2.639 Å (M06-2X/Def2-TZVP), 2.465–2.621 Å (B3PW91/Def2-TZVP), and 2.487–2.632 Å (B3LYP/LANL2DZ+dp). For RIn=BiR, the In=Bi triple bond length for small ligands is predicted to be 2.659–2.804 Å (M06-2X/Def2-TZVP), 2.691–2.802 Å (B3PW91/Def2-TZVP), and 2.673–2.795 Å (B3LYP/LANL2DZ+dp). For RTI=BiR, the TI=Bi triple bond length for small ligands is predicted to be 2.707–2.859 Å (M06-2X/Def2-TZVP), 2.698–2.812 Å (B3PW91/Def2-TZVP), and 2.679–2.822 Å (B3LYP/LANL2DZ+dp).
- 2. It is apparent from Tables 1–5 that an acute bond angle ∠E<sub>13</sub>=Bi=R (close to 90°) in the triply bonded molecule RE<sub>13</sub>=BiR is favored. The reason for this can be attributed to the "orbital nonhybridization effect," also known as the "inert *s*-pair effect" [65–68], as discussed previously. Accordingly, these phenomena strongly indicate that mode (B) (Figure 1) is preferred in the RE=BiR molecule, for which bent geometry is favored.
- 3. The Wiberg bond orders (WBOs) [63, 64] on the substituted RE<sub>13</sub>=BiR compounds are also given in **Tables 1–5**. For all the triply bonded RE<sub>13</sub>=BiR molecules with small substituents, their WBOs were computed to be less than 2.0, except for the cases of HB=BiH and (SiH<sub>3</sub>) B=Bi(SiH<sub>3</sub>). These WBO values imply that the bonding structure of RE<sub>13</sub>=BiR may be due to the resonance structures, [I] and [II]. That is to say, the E<sub>13</sub>=Bi bond could be either double or triple bonds. From **Tables 1–5**, it seems that the resonance structure [II] prevails for the small ligands on the substituted RE<sub>13</sub>=BiR species studied in this work. Indeed, since it is known that the electronegativities decrease in the order B (2.051) > Bi (2.01) > TI (1.789) > Ga (1.756) > In (1.656) > Al (1.613) [69], the bonding mode of RE<sub>13</sub>=BiR should prefer to adopt resonance structure [II] (**Scheme 2**).

With regard to the stability of  $RE_{13}BiR$ , the results of theoretical calculations on the energy surface of the model  $RE_{13}BiR$  (R = F, OH, H, CH<sub>3</sub>, and SiH<sub>3</sub>) system are depicted in **Figures 2–6**.



**Figure 2.** Relative Gibbs free energy surfaces for RB=BiR (R = F, OH, H, CH<sub>3</sub>, and SiH<sub>3</sub>). Energies are in kcal/mol, calculated at M06-2X/Def2-TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp levels of theory. For details see the text and **Table 1**.

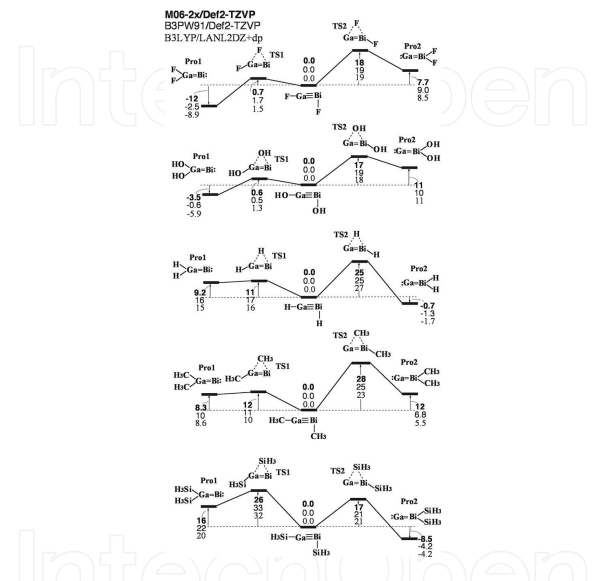
Triple Bonds between Bismuth and Group 13 Elements: Theoretical Designs and Characterization 81 http://dx.doi.org/10.5772/67220



**Figure 3.** Relative Gibbs free energy surfaces for RAl≡BiR (R = F, OH, H, CH<sub>3</sub>, and SiH<sub>3</sub>). Energies are in kcal/mol, calculated at M06-2X/Def2-TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp levels of theory. For details see the text and **Table 2**.

This system exhibits a number of stationary points, including local minima that correspond to  $RE_{13}$ =BiR,  $R_2E_{13}$ =Bi:,  $:E_{13}$ =Bi $R_2$ , and the saddle points connecting them. The transition structures that separate the three stable molecular forms involve a successive unimolecular 1,2-shift TS1 (from  $RE_{13}$ =BiR to  $R_2E_{13}$ =Bi:) and a 1,2-shift TS2 (from  $RE_{13}$ =BiR to  $:E_{13}$ =Bi $R_2$ ). As shown in **Figures 1–5**, these theoretical studies using the M06-2X, B3PW91, and B3LYP levels show that the  $RE_{13}$ =BiR species are local minima on the singlet potential energy surface, but they are neither kinetically nor thermodynamically stable for small substituents, except for the case of (SiH\_3)B=Bi(SiH\_3). As a result, these triply bonded structures  $RE_{13}$ =BiR seem to be unstable on the singlet energy surface and undergo unimolecular rearrangement to the doubly bonded isomer. In brief, these triply bonded molecules ( $RE_{13}$ =BiR) possessing the small substituents are

predicted to be a kinetically unstable isomer, so these could not be isolated in a matrix or even as transient intermediates.

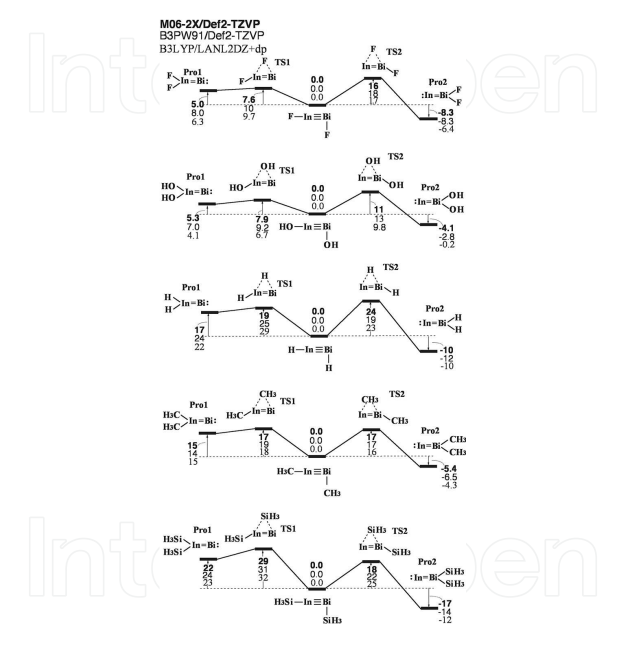


**Figure 4.** Relative Gibbs free energy surfaces for RGa=BiR (R = F, OH, H, CH<sub>3</sub>, and SiH<sub>3</sub>). Energies are in kcal/mol, calculated at M06-2X/Def2-TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp levels of theory. For details see the text and **Table 3**.

#### 3.3. Large ligands on substituted R'E<sub>13</sub>≡BiR'

According to the above conclusions for the cases of small substituents, it is necessary to determine whether bulky substituents can destabilize  $R_2E_{13}$ =Bi: and : $E_{13}$ =Bi $R_2$  relative to  $RE_{13}$ =BiR ( $E_{13}$  = B, Al, Ga, In, and Tl), due to severe steric overcrowding. From **Figure 7**, it is easily anticipated that the presence of extremely bulky substituents at both ends of the  $RE_{13}$ =BiR compounds protects its triple bond from intermolecular reactions, such as polymerization. In order to examine the effect of bulky substituents, the structures of  $R'E_{13}$ =BiR'

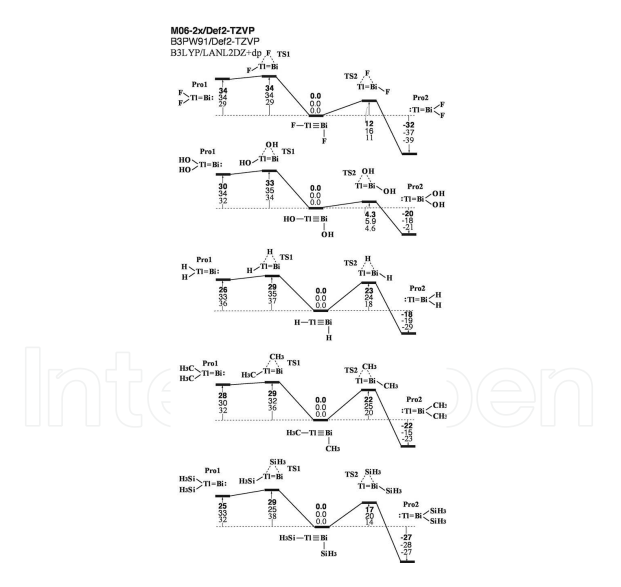
optimized for R' = Tbt, Ar\*, SiMe(SitBu<sub>3</sub>)<sub>2</sub>, and SitPrDis<sub>2</sub> (Scheme 1) at the B3LYP/LANL2DZ +dp level. Selected geometrical parameters, natural charge densities on the central group 13 elements and bismuth ( $Q_{E13}$  and  $Q_{Bi}$ ), binding energies (BE), and Wiberg bond order (BO) [69, 70] are summarized in **Tables 6–10**.



**Figure 5.** Relative Gibbs free energy surfaces for RIn=BiR (R = F, OH, H, CH<sub>3</sub>, and SiH<sub>3</sub>). Energies are in kcal/mol, calculated at M06-2X/Def2-TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp levels of theory. For details see the text and **Table 4**.

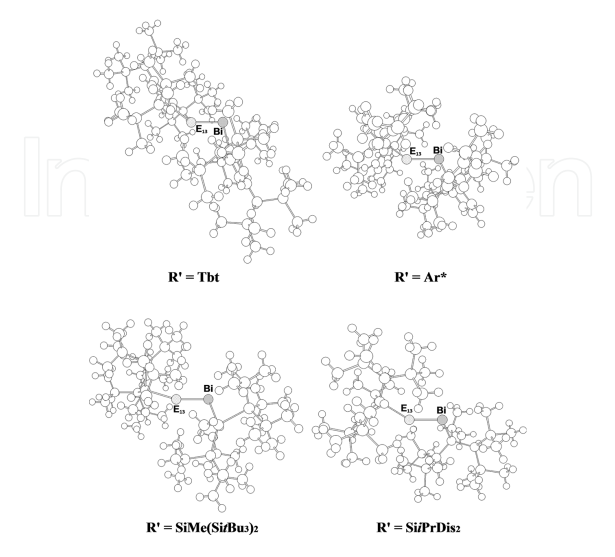
The computational results given in **Tables 6–10** estimate that the  $E_{13}$ =Bi triple bond distances (Å) are about 2.117–2.230 ( $E_{13}$  = B), 2.461–2.562 ( $E_{13}$  = Al), 2.576–2.580( $E_{13}$  = Ga), 2.615–2.779 ( $E_{13}$  = In), and 2.789–2.833 ( $E_{13}$  = Tl), respectively. Again, these theoretically predicted values are much shorter than the available experimentally determined  $E_{13}$ =Bi single bond lengths

[36, 70–72]. This strongly implies that the central group 13 element ( $E_{13}$ ) and bismuth in the R' $E_{13}$ =BiR' (R' = Tbt, Ar\*, SiMe(SitBu<sub>3</sub>)<sub>2</sub>, and SitPrDis<sub>2</sub>) species are triply bonded. Indeed, as shown in **Tables 6–10**, the R' $E_{13}$ =BiR' molecules accompanied by bulky ligands can effectively produce the triply bonded species. That is, the WBOs in **Tables 6–10** (with larger ligands) are apparently larger than those in **Tables 1–5** (with smaller ligands). Additionally, from **Tables 6–10**, the central  $E_{13}$ =Bi bond lengths calculated for R' = Tbt and Ar\* are an average 0.095Å longer than those calculated for R' = SiMe(SitBu<sub>3</sub>)<sub>2</sub> and SitPrDis<sub>2</sub>, respectively. The reason for these differences is that the Tbt and Ar\* groups are electronegative, but the SiMe (SitBu<sub>3</sub>)<sub>2</sub> and SitPrDis<sub>2</sub> ligands are electropositive. Further, the short length of the  $E_{13}$ =Bi bond in the R'E=BiR' species can be understood by noting that both SiMe(SitBu<sub>3</sub>)<sub>2</sub> and SitPrDis<sub>2</sub> are more electropositive than the small substituents, as mentioned earlier.



**Figure 6.** Relative Gibbs free energy surfaces for RTI=BiR (R = F, OH, H, CH<sub>3</sub>, and SiH<sub>3</sub>). Energies are in kcal/mol, calculated at M06-2X/Def2-TZVP, B3PW91/Def2-TZVP, and B3LYP/LANL2DZ+dp levels of theory. For details see the text and **Table 5**.

Triple Bonds between Bismuth and Group 13 Elements: Theoretical Designs and Characterization 85 http://dx.doi.org/10.5772/67220



**Figure 7.** The optimized structures of  $R'E_{13}$ =BiR' ( $E_{13}$  = B, Al, Ga, In, and Tl; R' = Tbt, Ar\*, SiMe(SitBu<sub>3</sub>)<sub>2</sub>, and SiiPrDis<sub>2</sub>) at the B3LYP/LANL2DZ+dp level of theory. For details see the text and **Tables 6–10**.

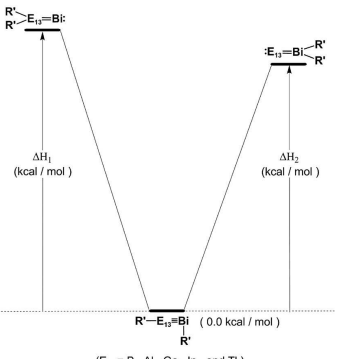
Similar to the small ligands, these DFT results demonstrate that all the R'E<sub>13</sub>BiR' molecules that possess bulky substituents (R') adopt a bent geometry, as illustrated in **Figure 7**. Our theoretical computations show that model (B), given in **Figure 1**, still predominates and can be used to interpret the geometries of the R'E<sub>13</sub>=BiR' systems that bear bulky substituents.

As shown in **Tables 6–10**, the R'E<sub>13</sub>=BiR' molecules can be separated into two fragments in solution, when the substituent R' becomes bulkier. The BE that is essential to break the central  $E_{13}$ =Bi bond was computed to be at least > 32 kcal/mol for R' = Tbt, Ar\*, SiMe(SitBu<sub>3</sub>)<sub>2</sub>, and SiiPrDis<sub>2</sub>, for the B3LYP/LANL2DZ+dp method, as given in **Tables 6–10**. These BE values show that the central  $E_{13}$  and bismuth elements are strongly bonded and R'E<sub>13</sub>=BiR' molecules that contain bulky substituents do not dissociate in solution. Namely, the larger the dissociation energy of the  $E_{13}$ =Bi bond, the shorter and stronger the  $E_{13}$ =Bi triple bond.

As predicted previously, bulky groups destabilize the 1,2-R' migrated isomers because they crowd around one end of the central  $E_{13}$ =Bi bond. As a consequence, the bulky substituents (R') can prevent the isomerization of R'E<sub>13</sub>=BiR' compounds, as outlined in **Scheme 3** and

**Tables 6–10.** The B3LYP/LANL2DZ+dp calculations indicate that the R'E<sub>13</sub>=BiR' species with Tbt, Ar\*, SiMe(SitBu<sub>3</sub>)<sub>2</sub>, and SitPrDis<sub>2</sub> substituents ( $\Delta$ H<sub>1</sub> and  $\Delta$ H<sub>2</sub>) are at least 56 kcal/mol more stable than the 1,2-R' shifted isomers, respectively. These theoretical results suggest that both doubly bonded R'<sub>2</sub>E<sub>13</sub>=Bi: and :E<sub>13</sub>=BiR'<sub>2</sub> isomers are kinetically and thermodynamically unstable, so they rearrange spontaneously to the global minimum R'E<sub>13</sub>=BiR' triply bonded molecules, provided that significantly bulky groups are employed.

Theoretical values from the natural bond orbital (NBO) [63, 64] and natural resonance theory (NRT) [73–75] analyses of the R'E<sub>13</sub>=BiR' molecules, computed at the B3LYP/LANL2DZ+dp level of theory, are summarized in **Table 11** (E<sub>13</sub> = B), **Table 12** (E<sub>13</sub> = Al), **Table 13** (E<sub>13</sub> = Ga), **Table 14** (E<sub>13</sub> = In), and **Table 15** (E<sub>13</sub> = Tl).



(E<sub>13</sub> = B , Al , Ga , In , and TI ) (R' = Tbt , Ar\* , SiMe(Si $tBu_3)_2$  , and Si $tPrDis_2$  )

R'	Tbt	Ar*	SiMe(SitBu <sub>3</sub> ) <sub>2</sub>	Si <i>i</i> PrDis <sub>2</sub>
B≡Bi (Å)	2.230	2.214	2.117	2.131
$\angle R' - B - Bi (^{\circ})$	177.3	110.3	112.9	113.6
∠B-Bi-R' (°)	115.6	115.5	112.5	114.3
$\angle R' - B - Bi - R'$ (°)	173.7	172.3	170.4	175.3
$Q_{\rm B}^{\ (1)}$	-0.4310	-0.1711	-0.3251	-0.4742
$Q_{\mathrm{Bi}}$ <sup>(2)</sup>	0.2915	0.3004	0.1426	0.1071
BE (kcal mol <sup><math>-1</math></sup> ) <sup>(3)</sup>	37.58	41.68	36.25	51.07

<b>R</b> '	Tbt	Ar*	SiMe(SitBu <sub>3</sub> ) <sub>2</sub>	Si <i>i</i> PrDis <sub>2</sub>
Wiberg BO <sup>(4)</sup>	2.385	2.249	2.640	2.701
$\Delta H_1$ (kcal mol <sup>-1</sup> ) <sup>(5)</sup>	62.6	61.04	76.52	77.24
$\Delta H_2$ (kcal mol <sup>-1</sup> ) <sup>(6)</sup>	98.51	88.38	68.72	78.13

Notes: (1) The natural charge density on the central boron atom. (2) The natural charge density on the central bismuth atom. (3) BE = E (triplet state of B–R') + E (triplet state of Bi–R') – E(R'B=BiR'). (4) Wiberg Bond Orders for the B–Bi bond, see Refs. [63, 64]. (5)  $\Delta H_1 = E(:B=BiR'_2) - E(R'B=BiR')$ ; see Scheme 3. (6)  $\Delta H_2 = E(R'_2B=Bi) - E(R'B=BiR')$ ; see Scheme 3.

**Table 6.** Geometrical parameters, nature charge densities ( $Q_B$  and  $Q_{Bi}$ ), binding energies (BE), and Wiberg bond order (BO) of R'B=BiR' at the B3LYP/LANL2DZ+dp level of theory. Also see **Figure 7**.

R'	Tbt	Ar*	SiMe(SitBu <sub>3</sub> ) <sub>2</sub>	Si <i>i</i> PrDis <sub>2</sub>
Al≡Bi (Å)	2.562	2.561	2.463	2.461
∠R'−Al−Bi (°)	178.0	113.6	115.4	113.3
$\angle Al - Bi - R' (^{\circ})$	113.4	115.5	112.2	109.0
$\angle R' - Al - Bi - R' (^{\circ})$	167.1	165.8	173.2	174.7
$Q_{Al}$ <sup>(1)</sup>	0.4171	0.4111	0.2173	0.1585
$Q_{\mathrm{Bi}}^{(2)}$	0.0773	0.2208	-0.1031	-0.1862
BE (kcal mol <sup><math>-1</math></sup> ) <sup>(3)</sup>	38.62	66.78	33.65	31.76
Wiberg BO <sup>(4)</sup>	2.092	2.023	2.204	2.259
$\Delta H_1$ (kcal mol <sup>-1</sup> ) <sup>(5)</sup>	64.52	63.77	77.71	73.68
$\Delta H_2$ (kcal mol <sup>-1</sup> ) <sup>(6)</sup>	57.43	67.81	71.86	76.62

Notes: (1) The natural charge density on the central aluminum atom. (2) The natural charge density on the central bismuth atom. (3) BE = E (triplet state of Al=R') + E (triplet state of Bi=R') – E(R'Al=BiR'). (4) Wiberg Bond Orders for the Al=Bi bond, see Refs. [63, 64]. (5)  $\Delta H_1 = E(:Al=BiR'_2) - E(R'Al=BiR')$ ; see Scheme 3. (6)  $\Delta H_2 = E(R'_2Al=Bi:) - E(R'Al=BiR')$ ; see Scheme 3.

**Table 7.** Geometrical parameters, nature charge densities ( $Q_{A1}$  and  $Q_{Bi}$ ), binding energies (BE), and Wiberg bond order (BO) of R'Al=BiR' at the B3LYP/LANL2DZ+dp level of theory. Also see **Figure 7**.

All the NBO values listed in **Tables 11–15** demonstrate that there exists a weak triple bond, or perhaps a bond between a double and a triple, in the ethyne-like R'E<sub>13</sub>=BiR' molecule. For instance, the B3LYP/LANL2DZ+dp data for the NBO [63, 64] analyses of the B=Bi bonding in SiMe(SitBu<sub>3</sub>)<sub>2</sub>=B=Bi=SiMe(SitBu<sub>3</sub>)<sub>2</sub>, which shows that NBO(B=Bi) =  $0.615(2s2p^{52.48})B + 0.789$  (6s6p<sup>19.73</sup>)Bi, strongly suggests that the predominant bonding interaction between the B=SiMe (SitBu<sub>3</sub>)<sub>2</sub> and the Bi=SiMe(SitBu<sub>3</sub>)<sub>2</sub> fragments originates from 2p(B)  $\leftarrow$  6p(Bi) donation. In other words, boron's electron deficiency and  $\pi$  bond polarity are partially balanced by the donation of the bismuth lone pair into the empty boron p orbital. This, in turn, forms a hybrid  $\pi$  bond. Again, the polarization analyses using the NBO model indicate the presence of the B=Bi  $\pi$  bonding orbital, 38% of which is composed of natural boron orbitals and 62% of natural bismuth orbitals. There is supporting evidence in **Table 11** that reveals that the B=Bi triple bond in SiMe(SitBu<sub>3</sub>)<sub>2</sub>=B=Bi=SiMe(SitBu<sub>3</sub>)<sub>2</sub> has a shorter single bond character (5.8%) and a

R'	Tbt	Ar*	SiMe(SitBu <sub>3</sub> ) <sub>2</sub>	Si <i>i</i> PrDis <sub>2</sub>
Ga≡Bi (Å)	2.578	2.576	2.580	2.579
∠R'−Ga−Bi (°)	178.1	113.4	115.2	112.1
$\angle Ga-Bi-R'$ (°)	113.4	115.7	112.0	110.1
$\angle R'-Ga-Bi-R'$ (°)	167.9	164.1	175.4	178.5
Q <sub>Ga</sub> <sup>(1)</sup>	0.240	0.196	0.069	0.012
Q <sub>Bi</sub> <sup>(2)</sup>	0.120	0.261	-0.055	-0.140
BE (kcal mol <sup>-1</sup> ) <sup>(3)</sup>	43.73	39.28	35.54	32.92
Wiberg BO <sup>(4)</sup>	2.091	2.181	2.262	2.313
$\Delta H_1$ (kcal mol <sup>-1</sup> ) <sup>(5)</sup>	68.10	69.08	61.74	58.83
$\Delta H_2$ (kcal mol <sup>-1</sup> ) <sup>(6)</sup>	78.07	71.28	77.43	64.13

Notes: (1) The natural charge density on the central gallium atom. (2) The natural charge density on the central bismuth atom. (3) BE = E (triplet state of Ga=R') + E (triplet state of Bi=R') – E(R'Ga=BiR'). (4) Wiberg bond orders for the Ga=Bi bond, see Refs. [63, 64]. (5)  $\Delta H_1$  = E(:Ga = BiR'\_2) – E(R'Ga=BiR'); see Scheme 3. (6)  $\Delta H_2$  = E(R'\_2Ga=Bi:) – E(R'Ga=BiR'); see Scheme 3.

**Table 8.** Geometrical parameters, nature charge densities ( $Q_{Ga}$  and  $Q_{Bi}$ ), binding energies (BE), and Wiberg bond order (BO) of R'Ga=BiR' at the B3LYP/LANL2DZ+dp level of theory. Also see **Figure 7**.

R'	Tbt	Ar*	SiMe(SitBu <sub>3</sub> ) <sub>2</sub>	Si <i>i</i> PrDis <sub>2</sub>
In≡Bi (Å)	2.737	2.779	2.615	2.678
∠R'−In−Bi (°)	178.7	111.7	110.0	110.9
$\angle$ In-Bi-R' (°)	112.5	113.0	110.6	111.7
$\angle R'$ -In-Bi-R' (°)	170.7	174.6	164.3	162.0
$Q_{\mathrm{In}}^{(1)}$	0.299	0.345	0.179	0.101
$Q_{\mathrm{Bi}}^{(2)}$	0.066	0.293	-0.126	-0.132
BE (kcal mol <sup><math>-1</math></sup> ) <sup>(3)</sup>	63.45	45.97	36.20	37.05
Wiberg BO <sup>(4)</sup>	2.052	2.153	2.211	2.304
$\Delta H_1$ (kcal mol <sup>-1</sup> ) <sup>(5)</sup>	64.06	60.17	55.72	62.99
$\Delta H_2$ (kcal mol <sup>-1</sup> ) <sup>(6)</sup>	79.38	61.44	56.03	67.61

Notes: (1) The natural charge density on the central indium atom. (2) The natural charge density on the central bismuth atom. (3) BE = E (triplet state of In=R') + E (triplet state of Bi=R') – E(R'In=BiR'). (4) Wiberg Bond Orders for the In=Bi bond, see Refs. [63, 64]. (5)  $\Delta H_1 = E(:In = BiR'_2) - E(R'In=BiR')$ ; see Scheme 3. (6)  $\Delta H_2 = E(R'_2In=Bi:) - E(R'In=BiR')$ ; see Scheme 3.

**Table 9.** Geometrical parameters, nature charge densities ( $Q_{In}$  and  $Q_{Bi}$ ), binding energies (BE), and Wiberg bond order (BO) of R'In=BiR' at the B3LYP/LANL2DZ+dp level of theory. Also see **Figure 7**.

shorter triple bond character (40.1%) but a larger double bond character (54.1%), because the covalent part of the NRT bond order (1.49) is shorter than its ionic part (0.78). The same can also be said of the other three R'B≡BiR' molecules, as shown in **Table 11** as well as other

 $R'E_{13}$ =BiR' compounds represented in **Tables 12–15**. These theoretical evidences strongly suggest that these  $R'E_{13}$ =BiR' species have a weak triple bond.

R'	Tbt	Ar*	SiMe(SitBu <sub>3</sub> ) <sub>2</sub>	Si <i>i</i> PrDis <sub>2</sub>
Tl≡Bi (Å)	2.816	2.833	2.820	2.789
$\angle R' - Tl - Bi (^{\circ})$	148.2	146.7	164.4	152.4
$\angle Tl - Bi - R' (^{\circ})$	116.5	121.4	114.1	125.1
$\angle R' - Tl - Bi - R(^{\circ})$	169.5	171.4	176.3	175.8
Q <sub>TI</sub> <sup>(1)</sup>	0.3323	0.4221	0.1504	0.1282
$Q_{\mathrm{Bi}}^{(2)}$	-0.0921	-0.0652	-0.3925	-0.3491
BE (kcal mol <sup>-1</sup> ) <sup>(3)</sup>	41.66	37.59	36.66	51.07
Wiberg BO <sup>(4)</sup>	2.081	2.052	2.105	2.128
$\Delta H_1$ (kcal mol <sup>-1</sup> ) <sup>(5)</sup>	68.66	64.52	63.67	72.12
$\Delta H_2$ (kcal mol <sup>-1</sup> ) <sup>(6)</sup>	61.54	58.19	68.73	64.61

Notes: (1) The natural charge density on the central thallium atom. (2) The natural charge density on the central bismuth atom. (3) BE = E (triplet state of Tl=R') + E (triplet state of Bi=R') – E(R'Tl=BiR'). (4) Wiberg bond orders for the Tl=Bi bond, see Refs. [63, 64]. (5)  $\Delta H_1 = E(:Tl = BiR'_2) - E(R'Tl=BiR')$ ; see Scheme 3. (6)  $\Delta H_2 = E(R'_2Tl = Bi:) - E(R'Tl=BiR')$ ; see Scheme 3.

**Table 10.** Geometrical parameters, nature charge densities ( $Q_{TI}$  and  $Q_{Bi}$ ), binding energies (BE), and Wiberg bond order (BO) of R'TI=BiR' at the B3LYP/LANL2DZ+dp level of theory. Also see **Figure 7**.

R′B≡BiR′	WBI	NBO analys	sis		NRT analysis	
		Occupancy	Hybridization	Polarization	Total/covalent/ ionic	Resonance weight
R'=Tbt	2.39	σ = 1.95	σ: 0.7870 B (sp <sup>0.90</sup> ) + 0.6170 Bi	61.93% (B)	2.01/1.24/0.77	B=Bi: 12.58%
			(sp <sup>9.90</sup> )	38.07% (Bi)		B=Bi: 53.81%
		π = 1.93	π: 0.5938 B (sp <sup>1.00</sup> ) + 0.8046 Bi	35.26% (B)		B≡Bi: 33.61%
			(sp <sup>1.00</sup> )	64.74% (Bi)		
$R' = Ar^*$	2.25	σ = 1.94	σ: 0.8058 B (sp <sup>0.74</sup> ) + 0.5922 Bi	64.93% (B)	1.95/1.30/0.65	B=Bi: 6.78%
			(sp <sup>19.46</sup> )	35.07% (Bi)		B=Bi: 62.97%
		$\pi = 1.90$	π: 0.5587 B (sp <sup>99.99</sup> ) + 0.8294 Bi	31.22% (B)		B≡Bi: 30.25%
			(sp <sup>64.69</sup> )	68.78% (Bi)		
R' = SiMe	2.64	σ = 1.96	σ: 0.7812 B (sp <sup>0.96</sup> ) + 0.6242 Bi	61.03% (B)	2.27/1.49/0.78	B=Bi: 5.77%
(SitBu <sub>3</sub> ) <sub>2</sub>			(sp <sup>10.68</sup> )	38.97% (Bi)		B=Bi: 54.1%

R'B≡BiR' WE	I NBO analys	sis		NRT analysis	
	Occupancy	Hybridization	Polarization	Total/covalent/ ionic	Resonance weight
	π = 1.89	π: 0.6146 B (sp <sup>62.48</sup> ) + 0.7889 Bi	37.77% (B)		B≡Bi: 40.11%
		(sp <sup>19.73</sup> )	62.23% (Bi)		
$R' = SiiPrDis_2$ 2.7		σ: 0.6502 B (sp <sup>4.29</sup> ) + 0.7598 Bi (sp <sup>1.07</sup> )	42.27% (B)	2.31/1.52/0.79	B=Bi: 6.01%
			57.73% (Bi)		B=Bi: 54.39%
	$\pi = 1.80$	π: 0.5606 B (sp <sup>1.73</sup> ) + 0.8281 Bi	31.43% (B)		B≡Bi: 39.96%
		(sp <sup>4.99</sup> )	68.57% (Bi)		

(1) The Wiberg bond index (WBI) for the B=Bi bond and occupancy of the corresponding  $\sigma$  and  $\pi$  bonding NBO: see Refs. [63, 64], and (2) the natural resonance theory (NRT): see Refs. [73–75].

**Table 11.** Selected results for the natural bond orbital (NBO) and natural resonance theory (NRT) analyses of R'B=BiR' compounds that have small substituents, at the B3LYP/LANL2DZ+dp level of theory [1–8, 76–80].

R'Al≡BiR'	WBI	NBO analys	NBO analysis					
		Occupancy	Hybridization	Polarization	Total/covalent/ ionic	Resonance weight		
R' = Tbt	2.09	σ = 1.98	σ: 0.7538 Al (sp <sup>0.15</sup> ) + 0.6571 Bi	56.83% (Al)	2.12/1.10/1.02	Al=Bi: 12.76%		
			(sp <sup>22.99</sup> )	43.17% (Bi)		Al=Bi: 75.36%		
		π = 1.93	$\pi$ : 0.4709 Al (sp <sup>1.00</sup> ) + 0.8822 Bi	22.17% (Al)		Al≡Bi: 11.88%		
			(sp <sup>1.00</sup> )	77.83% (Bi)				
$R' = Ar^*$	2.02	σ = 1.84	σ: 0.7806 Al (sp <sup>0.15</sup> ) + 0.6250 Bi	60.93% (Al)	2.07/1.01/1.06	Al=Bi: 19.33%		
			(sp <sup>28.77</sup> )	39.07% (Bi)		Al=Bi: 74.20%		
				π = 1.94	$\pi$ : 0.4960 Al (sp <sup>46.09</sup> ) + 0.8673 Bi	24.60% (Al)		Al=Bi: 6.47%
			(sp <sup>15.43</sup> )	75.40% (Bi)				
R' = SiMe	2.2	σ = 1.96		24.70% (Al)	2.24/1.38/0.86	Al=Bi: 11.69%		
(SitBu <sub>3</sub> ) <sub>2</sub>			(sp <sup>21.26</sup> )	75.30% (Bi)		Al=Bi: 84.51%		
			π: 0.8678 Al (sp <sup>19.21</sup> ) + 0.4970 Bi	37.77% (Al)		Al≡Bi: 3.80%		
			(sp <sup>16.37</sup> )	62.23% (Bi)				
$R' = SiiPrDis_2$	2.26	σ = 1.86	σ: 0.7184 Al (sp <sup>0.93</sup> ) + 0.6956 Bi	51.61% (Al)	1.91/1.35/0.60	Al=Bi: 12.68%		
		(sp <sup>29.72</sup> )	48.39% (Bi)		Al=Bi: 83.75%			

R'Al≡BiR'	WBI	NBO analys	sis		NRT analysis	
		Occupancy	Hybridization	Polarization	Total/covalent/ ionic	Resonance weight
		$\pi = 1.90$	π: 0.4430 Al (sp <sup>59.07</sup> ) + 0.8965 Bi	19.63% (Al)		Al≡Bi: 3.57%
			(sp <sup>35.38</sup> )	80.37% (Bi)		

(1) The Wiberg bond index (WBI) for the Al=Bi bond and occupancy of the corresponding  $\sigma$  and  $\pi$  bonding NBO: see Refs. [63, 64], and (2) the natural resonance theory (NRT): see Refs. [73–75].

**Table 12.** Selected results for the natural bond orbital (NBO) and natural resonance theory (NRT) analyses of R'Al=BiR' compounds that have small substituents, at the B3LYP/LANL2DZ+dp level of theory [1–8, 76–80].

R′Ga≡BiR′	WBI	NBO analysis			NRT analysis	
		Occupancy	Hybridization	Polarization	Total/covalent/ ionic	Resonance weight
R' = Tbt	2.09	σ = 1.78	σ: 0.7338 Ga (sp <sup>1.04</sup> ) + 0.6794 Bi	53.85% (Ga)	2.09/1.36/0.73	Ga=Bi: 11.42%
			(sp <sup>31.81</sup> )	46.15% (Bi)		Ga=Bi: 87.53%
		$\pi = 1.94$	$\pi$ : 0.4586 Ga (sp <sup>1.00</sup> ) + 0.8886 Bi	21.04% (Ga)		Ga≡Bi: 1.05%
			(sp <sup>1.00</sup> )	78.96% (Bi)		
$R' = Ar^*$	2.18	σ = 1.93	σ: 0.8142 Ga (sp <sup>0.12</sup> ) + 0.5806 Bi	66.29% (Ga)	2.05/1.30/0.75	Ga=Bi: 10.87%
			(sp <sup>40.52</sup> )	33.71% (Bi)		Ga=Bi: 88.05%
		$\pi$ = 1.84	34 π: 0.4810 Ga (sp <sup>50.27</sup> ) + 0.8767 Bi (sp <sup>14.17</sup> )	23.13% (Ga)		Ga≡Bi: 1.08%
				76.87% (Bi)		
R' = SiMe	2.26	σ=1.80	σ: 0.7393 Ga (sp <sup>0.97</sup> ) + 0.6733 Bi	54.66% (Ga)	2.11/1.35/0.76	Ga=Bi: 11.84%
(SitBu <sub>3</sub> ) <sub>2</sub>			(sp <sup>27.39</sup> )	67.33% (Bi)		Ga=Bi: 82.11%
		π = 1.86	$\pi$ : 0.4937 Ga (sp <sup>22.70</sup> ) + 0.8696 Bi	24.38% (Ga)		Ga≡Bi: 6.05%
			(sp <sup>17.00</sup> )	75.62% (Bi)		
$R' = SiiPrDis_2$	2.31	σ = 1.90	σ: 0.7462 Ga (sp <sup>0.92</sup> ) + 0.6657 Bi	55.68% (Ga)	1.93/1.40/0.53	Ga=Bi: 12.93%
			(sp <sup>58.99</sup> )	44.32% (Bi)		Ga=Bi: 80.74%
		$\pi = 1.84$	$\pi$ : 0.4201 Ga (sp <sup>99.99</sup> ) + 0.9075 Bi	17.65% (Ga)		Ga≡Bi: 6.33%
			(sp <sup>99.99</sup> )	82.35% (Bi)		

(1) The Wiberg bond index (WBI) for the Ga=Bi bond and occupancy of the corresponding  $\sigma$  and  $\pi$  bonding NBO: see Refs. [63, 64], and (2) the natural resonance theory (NRT): see Refs. [73–75].

**Table 13.** Selected results for the natural bond orbital (NBO) and natural resonance theory (NRT) analyses of R'Ga=BiR' compounds that have small substituents, at the B3LYP/LANL2DZ+dp level of theory [1–8, 76–80].

R'In≡BiR'	WBI	VBI NBO analysis				NRT analysis	
		Occupancy	Hybridization	Polarization	Total/covalent/ ionic	Resonance weight	
R' = Tbt	2.05	σ = 1.78	σ: 0.7626 In (sp <sup>0.09</sup> ) + 0.6469 Bi	58.15% (In)	2.10/1.15/0.95	In=Bi: 14.33%	
			(sp <sup>50.74</sup> )	41.85% (Bi)		In=Bi: 76.37%	
		π = 1.94	π: 0.4230 In (sp <sup>99.99</sup> ) + 0.9061 Bi	17.85% (In)		In≡Bi: 0.93%	
			(sp <sup>95.23</sup> )	82.11% (Bi)			
$R' = Ar^*$	2.15	σ = 1.97	$\sigma = 1.97$ $\sigma: 0.7145 \text{ In } (\text{sp}^{0.07}) + 0.6996 \text{ Bi}$ (sp <sup>21.13</sup> )	51.05% (In)	2.11/1.04/1.07	In=Bi: 11.10%	
				48.95% (Bi)		In=Bi: 85.07%	
		$\pi = 1.94$	= 1.94 $\pi$ : 0.4774 In (sp <sup>99.99</sup> ) + 0.8787Bi (sp <sup>99.99</sup> )	22.79% (In)		In≡Bi: 3.83%	
				77.21% (Bi)			
R' = SiMe	2.21	1 σ=1.74	$\sigma = 1.74$ $\sigma: 0.7433 \text{ In } (\text{sp}^{0.99}) + 0.6690 \text{ Bi}$ (sp <sup>35.62</sup> )	55.25% (In)	2.14/1.22/0.92	In=Bi: 14.20%	
(SitBu <sub>3</sub> ) <sub>2</sub>				44.75% (Bi)		In=Bi: 81.22%	
		$\pi = 1.88$	π: 0.4468 In (sp <sup>42.13</sup> ) + 0.8696 Bi	19.96% (In)		In≡Bi: 4.58%	
			(sp <sup>12.11</sup> )	80.04% (Bi)			
$R' = SiiPrDis_2$	2.3	σ = 1.92	σ: 0.7341 In (sp <sup>53.22</sup> ) + 0.6790 Bi	46.11% (In)	1.88/1.27/0.61	In=Bi: 15.31%	
			(sp <sup>0.98</sup> )	53.89% (Bi)		In=Bi: 81.02%	
		$\pi = 1.78$	$\pi$ : 0.4357 In (sp <sup>26.46</sup> ) + 0.9001 Bi	18.99% (In)		In≡Bi: 3.67%	
			(sp <sup>99.99</sup> )	81.01% (Bi)			

Notes: (1) The Wiberg bond index (WBI) for the In=Bi bond and occupancy of the corresponding  $\sigma$  and  $\pi$  bonding NBO: see Refs. [63, 64], and (2) the natural resonance theory (NRT): see Refs. [73–75].

**Table 14.** Selected results for the natural bond orbital (NBO) and natural resonance theory (NRT) analyses of R'In=BiR' compounds that have small substituents, at the B3LYP/LANL2DZ+dp level of theory [1–8, 76–80].

R'Tl≡BiR'	WBI	I NBO analysis			NRT analysis			
		Occupancy	Hybridization	Polarization	Total/covalent/ ionic	Resonance weight		
R' = Tbt 2	2.08	σ = 1.96	σ:0.8268 Tl (sp <sup>0.07</sup> ) + 0.5625 Bi	68.36% (Tl)	2.15/1.81/0.34	Tl=Bi: 18.71%		
			(sp <sup>99.99</sup> )	31.64% (Bi)		Tl = Bi: 71.54%		
		$\pi = 1.94$	π:0.4315 In (sp <sup>99.99</sup> ) + 0.9021 Bi	18.61% (In)		Tl≡Bi: 9.75%		
			(sp <sup>99.99</sup> )	81.37% (Bi)				
$R' = Ar^*$		2.05	2.05	σ = 1.97	σ: 0.8159 Tl (sp <sup>0.05</sup> ) + 0.5782 Bi	66.57% (Tl)	2.11/1.70/0.41	Tl=Bi: 21.11%
		(sp <sup>99.99</sup> )	(sp <sup>99.99</sup> )	33.43% (Bi)		Tl = Bi: 70.15%		
		$\pi = 1.95$	π: 0.4788 In (sp <sup>99.99</sup> ) + 0.8779 Bi	22.92% (In)		Tl≡Bi: 8.74%		
			(sp <sup>99.99</sup> )	77.08% (Bi)				

R′Tl≡BiR′	WBI	NBO analysis			NRT analysis	
		Occupancy	Hybridization	Polarization	Total/covalent/ ionic	Resonance weight
R' = SiMe 2.1 (SitBu3)2	2.1	σ = 1.98	σ: 0.7986 Tl (sp <sup>0.02</sup> ) + 0.6019 Bi	63.77% (Tl)	2.21/1.40/0.81	Tl=Bi: 16.13%
			(sp <sup>31.50</sup> )	36.23% (Bi)		Tl = Bi: 76.20%
		π = 1.92	π: 0.3890 Tl (sp <sup>99.99</sup> ) + 0.9212 Bi (sp <sup>1.00</sup> )	15.13% (Tl)		Tl≡Bi: 7.67%
				84.87% (Bi)		
R' = SiiPrDis2	2.13	σ = 1.98	σ: 0.7757 Tl (sp <sup>0.03</sup> ) + 0.6311 Bi (sp <sup>23.28</sup> )	60.17% (Tl)	1.85/1.23/0.62	Tl=Bi: 20.21%
				39.83% (Bi)		Tl=Bi: 74.69%
		$\pi$ = 1.91 $\pi$ : 0.4149 Tl (sp <sup>99.99</sup> ) + 0.9099 Bi (sp <sup>99.99</sup> )	$\pi$ : 0.4149 Tl (sp <sup>99.99</sup> ) + 0.9099 Bi	17.22% (Tl)		Tl≡Bi: 5.10%
			(sp <sup>99.99</sup> )	82.78% (Bi)		

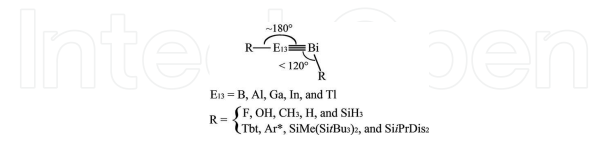
Notes: (1) The Wiberg bond index (WBI) for the TI=Bi bond and occupancy of the corresponding  $\sigma$  and  $\pi$  bonding NBO: see Refs. [63, 64], and (2) the natural resonance theory (NRT): see Refs. [73–75].

**Table 15.** Selected results for the natural bond orbital (NBO) and natural resonance theory (NRT) analyses of R'Tl=BiR' compounds that have small substituents, at the B3LYP/LANL2DZ+dp level of theory [1–8, 76–80].

### 4. Overview of RE<sub>13</sub>=BiR (E = B, Al, Ga, In, and Tl) systems

This study of the effect of substituents on the possibilities of the existence of triply bonded  $RE_{13}$ =BiR allows the following conclusions to be drawn (**Scheme 4**):

1. The theoretical observations strongly demonstrate that bonding mode (B) is dominant in the triply bonded RE<sub>13</sub>≡BiR species, since their structures are bent to increase stability, due to electron transfer (denoted by arrows in **Figure 1**) as well as the relativistic effect [65–68].



2. The theoretical evidence shows that both the electronic and the steric effects of substituents are crucial to making the  $E_{13}$ =Bi triple bond synthetically accessible. Based on the present theoretical study, however, these  $E_{13}$ =Bi triple bonds should be weak, not as strong as the traditional C=C triple bond. From our theoretical study, both bulky and electropositive substituents, such as the silvl groups demonstrated in Scheme 1, have a significant effect on the stability of  $E_{13}$ =Bi triply bonded compounds.

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