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A wealth of structures for the Ge_2H_2^+ radical cation: comparison of theory and experiment[†]

Ethan J. Poncelet, ^a Henry F. Mull, ^a Yohannes Abate, ^b Gregory H. Robinson, ^c Gary E. Doublerly, ^c Justin M. Turney^a and Henry F. Schaefer III ^a

Five structures of Ge_2H_2 and Ge_2H_2^+ are investigated in this study. Optimized geometries at the CCSD(T)/cc-pwCVQZ-PP level of theory were obtained. Focal point analyses were performed on these optimized geometries to determine relative energies using the CCSD(T) method with polarized basis sets up to quintuple-zeta. Energy corrections include full T and perturbative (Q) coupled-cluster effects plus anharmonic corrections to the zero-point vibrational energy. Relative ordering in energy from lowest to highest of the five Ge_2H_2^+ structures is butterfly, germylidene, monobridged, *trans*, then linear. In neutral Ge_2H_2 , the monobridged structure lies lower in energy than the germylidene structure. Fundamental vibrational frequencies and IR intensities were computed for the minima at the CCSD(T)/cc-pwCVTZ-PP level of theory to compare with experimental research. Partial atomic charges and natural bonding orbital analyses indicated that the positive charge of Ge_2H_2^+ is contained in the region of the Ge–Ge bond.

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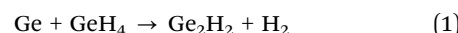
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Introduction

Germanium, as well as other group 14 elements, continues to be studied for its use in the production of high-purity nanoparticles,^{1–5} nanowires,^{6–8} and thin films.^{9–14} Use of germanium in the electronics industry is advantageous, as it has a greater carrier mobility than commonly-used silicon.^{14,15} As a result, thin-film and field-effect germanium transistors have the potential to increase performance speed while maintaining acceptable operating temperatures.^{10,11,16} Further benefits include nonhazardous, electrochemically-stable biological imaging and therapeutics using germanium nanocrystals instead of the frequently used Cd, Hg, and Pb.^{1,2,14}

GeH_4 and Ge_2H_6 are common germanium-hydride precursors used in the production of the aforementioned nanomaterials.^{3–5,7,12,13} In 2002, Wang, Andrews, and Kushto observed the formation of GeH_4 , Ge_2H_6 , and Ge_2H_2 from reactions of atomic germanium and hydrogen gas.¹⁷ These germanium hydrides were isolated in argon and neon matrices and identified using IR spectroscopy. Ge_2H_2 was identified from bands at 964.3 cm^{-1} and 972.2 cm^{-1} in argon and neon, respectively. They proposed

these bands to be the absorption of the Ge_2H_2 global minimum, butterfly geometry. Gas-phase formation of another Ge_2H_2 structure, the atypical monobridged geometry, was observed by Kaiser and coworkers in 2022. Time-of-flight mass spectrometry (TOF-MS) was used to determine an experimental reaction enthalpy of $-16 \pm 4\text{ kcal mol}^{-1}$ for the formation of the monobridged Ge_2H_2 structure from atomic germanium and GeH_4 (eqn (1)). Computationally, Kaiser and coworkers determined this reaction enthalpy to be -15 kcal mol^{-1} . Harmonic CCSD/cc-pVTZ ZPVE corrected CCSD(T)/CBS//CCSD/cc-pVTZ relative energies for the neutral Ge_2H_2 species were computed by Kaiser and coworkers, which were butterfly, monobridged (8.4 kcal mol⁻¹), germylidene (10.5), and *trans* (15.8).¹⁸



Si_2H_2 was the first molecule of interest in the group 14 M_2H_2 acetylene analogues with theoretical studies dating back to 1972.¹⁹ In 1983, Lischka and Kohler were the first to correctly propose the Si_2H_2 global minimum to be the butterfly geometry, also identifying the disilavinylidene geometry to be a local minimum.²⁰ Surprisingly, another Si_2H_2 stationary point was found to have an unusual, low-lying, monobridged equilibrium geometry in 1990 by Colgrove and Schaefer.²¹ 1990 was also the year of the first theoretical study of Ge_2H_2 by Grev and Schaefer, though it was absent of a monobridged equilibrium geometry. Stationary points of Ge_2H_2 predicted in 1990 included the germylidene, butterfly, planar dibridged, linear, and *trans*

^a Center for Computational Quantum Chemistry, University of Georgia, Athens, Georgia 30602, USA. E-mail: ccq@uga.edu

^b Department of Physics and Astronomy, University of Georgia, Athens, Georgia

^c Department of Chemistry, University of Georgia, Athens, Georgia 30602, USA

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structures. The planar dibridged and linear structures were found to be transition states and the butterfly, germylidene, and *trans* structures were identified as minima. Using the most accurate yet computationally feasible *ab initio* methods at the time, they predicted the relative energies of these structures to be butterfly, germylidene (9.5 kcal mol⁻¹), planar dibridged (11.0), *trans* (17.7), and linear (45.8). Stationary point geometries were optimized by Grev at the Hartree–Fock level of theory using basis set “B1” with the contraction scheme Ge(13s9p6d/7s5p4d), H(4s/2s); relative energies were computed at the CISD+Q level of theory with basis set “B1” augmented with f-type polarization functions on germanium.²²

In 1993, Pálágyi and Schaefer were the first to predict a monobridged local minimum for Ge₂H₂.²³ In that paper, they provided geometries, energies, and vibrational frequencies for the butterfly, monobridged, and germylidene structures. Due to computational limitations of the time, there was disagreement between theory and the subsequent matrix isolation experiments of Wang, Andrews, and Kushto in 2002; the b₂ Ge–H stretching mode of the butterfly structure was computed to have a frequency of 1056 cm⁻¹ compared to the experimental value of 972.2 cm⁻¹. Vibrational frequencies for the three structures were evaluated at the CCSD/DZP level of theory. Although the the energy of the monobridged structure computed by Pálágyi cannot be compared to the enthalpy of formation in the gas phase experiment observed by Kaiser and coworkers, the respective 8.9 and 8.4 kcal mol⁻¹ relative computational energies compare well. Energetics were evaluated by Pálágyi at the CCSD(T) level of theory with a TZP basis set and corrected with harmonic ZPVEs. While the germylidene structure of Ge₂H₂ has yet to be observed experimentally, there are similar relative energy predictions of 11.0 kcal mol⁻¹ by Pálágyi and 10.5 kcal mol⁻¹ by Kaiser and coworkers.^{18,23}

Methods

Our work involving Ge₂H₂ and Ge₂H₂⁺ examines five of the previously presented structures of Ge₂H₂: butterfly, monobridged, germylidene, *trans*, and linear.^{22–25} Each structure considered here was investigated using high-level *ab initio* quantum chemical methods. The correlation consistent small core pseudopotential-containing basis set family cc-pwCVXZ-PP (X = D,T,Q,5)^{26–30} was used to account for relativistic effects on the core electrons of the germanium atoms.³¹ Ten of the 32 electrons in germanium were contained in the pseudopotential, having the noble gas configuration of neon (1s² 2s² 2p⁶). Hydrogen was assigned the basis set family of cc-pVXZ.³²

A restricted open shell Hartree–Fock (ROHF) reference wavefunction was used for calculations involving the cationic species and restricted Hartree–Fock (RHF) was used for the neutral species. Geometries were optimized at the coupled cluster singles, doubles, and perturbative triple excitations [CCSD(T)]³³ level of theory with the respective quadrupole-zeta (QZ) polarized basis sets on hydrogen and germanium. Fundamental vibrational frequencies were computed at the

CCSD(T)/TZ level of theory. Single point energies were computed at Hartree–Fock (HF), Møller–Plesset second-order perturbation theory (MP2), coupled cluster singles and doubles (CCSD), and CCSD(T), with double, triple, quadruple, and quintuple-zeta polarized basis sets. Focal point analysis^{34–37} was used to extrapolate these energies to the CBS limit using a three-point fitting equation for HF (eqn (2)) and a two-point fitting equation for correlated levels of theory (eqn (3)).

$$E_{\text{HF}} = A + B e^{-cx} \quad (2)$$

$$E_{\text{corr}} = A + B X^{-3} \quad (3)$$

Energy corrections incorporated coupled cluster singles, doubles, and triple excitations (CCSDT)³⁸ and coupled cluster singles, doubles, triples, and perturbative quadruple excitations (CCSDT(Q)),³⁹ both with double-zeta polarized basis sets. In addition, CCSD(T)/TZ harmonic ZPVEs with VPT2 anharmonic corrections at the same level of theory were used to obtain the final energies of the minimum structures. All computations done on Ge₂H₂⁺ were replicated for its neutral counterpart, Ge₂H₂.

CFOUR 2.0⁴⁰ was used for the aforementioned computations. Natural bond order (NBO) analyses were performed using NBO 7.0⁴¹ to determine natural bonding orders and partial charges. The NBO results were computed using the HF/def2-TZVP method and the CCSD(T)/QZ optimized structures.⁴²

Results and discussion

Geometries

CCSD(T)/QZ optimized geometries of five Ge₂H₂ and Ge₂H₂⁺ structures are presented in Table 1. The butterfly (C_{2v}), monobridged (C_s), germylidene (C_{2v}), and *trans* (C_{2h}) structures were found to be minima while the linear (D_{∞h}) structure was determined to be a second-order transition state (Fig. 1).

Table 1 Optimized geometries at the CCSD(T)/cc-pwCVQZ-PP level of theory for five structures of the neutral and cationic Ge₂H₂ species are presented. Bond lengths are reported in Ångströms and angles in degrees

Geometry	Internal coordinate	Cation	Neutral
Butterfly	Ge–Ge	2.437	2.346
	Ge–H	1.764	1.750
	τ(H–Ge–Ge–H)	108.5	105.0
Monobridged	Ge–Ge	2.326	2.216
	Ge ₍₁₎ –H ₍₁₎	1.812	1.791
	Ge ₍₂₎ –H ₍₁₎	1.747	1.712
	Ge ₍₂₎ –H ₍₂₎	1.528	1.529
	H–Ge–H	105.3	105.8
Germylidene	Ge–Ge	2.417	2.280
	Ge–H	1.516	1.521
	Ge–Ge–H	123.2	124.7
<i>trans</i>	Ge–Ge	2.302	2.186
	Ge–H	1.535	1.533
	Ge–Ge–H	122.5	124.2
Linear	Ge–Ge	2.092	2.037
	Ge–H	1.490	1.480



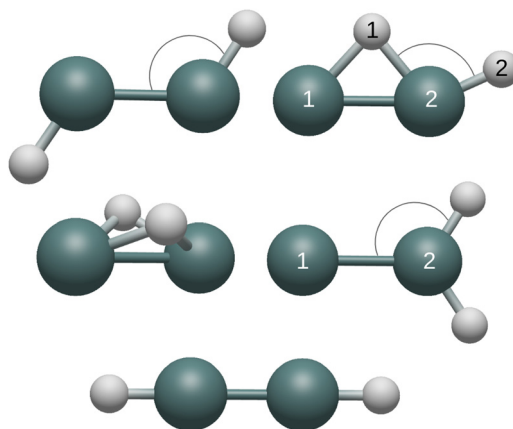


Fig. 1 From left to right and top to bottom, the five Ge_2H_2 structures of interest are shown: *trans*, monobridged, butterfly, germylidene, and linear. Structures containing non-equivalent atoms and bonds are henceforth referenced with the numbers in the figure. (ex: the terminal Ge–H bond of the monobridged structure would be $\text{Ge}_{(2)}-\text{H}_{(2)}$).

Classification of these five stationary points is in agreement with previous theoretical work on Ge_2H_2 .^{22,23} Furthermore, the same minima were identified by Schueller and coworkers for the isovalent Si_2H_2^+ . They found the Si–Si bond distances to be larger in Si_2H_2^+ compared to neutral Si_2H_2 .⁴³ Similarly, this trend is also seen here in Ge_2H_2^+ , as all Ge–Ge bond distances are larger than in neutral Ge_2H_2 .

As a triply-bonded structure, linear Ge_2H_2 and Ge_2H_2^+ have the shortest Ge–Ge and Ge–H bonds. Consequently, the linear Ge_2H_2^+ had the smallest increase in Ge–Ge bond length from its neutral counterpart at 0.055 Å. Bridged hydrogens appear to hamper the increase of the Ge–Ge bond lengths when comparing the cationic and the neutral species. Therefore, the butterfly and monobridged structures exhibit limited increases in Ge–Ge bond lengths from the neutral to cationic structures at 0.091 and 0.110 Å, respectively. Containing a single bridged hydrogen compared to the doubly-bridged butterfly structure, the monobridged structure has a modestly larger increase than the butterfly structure of 0.110 Å from the neutral to cationic structure. Containing no bridged hydrogen, the *trans* and germylidene structures have the greatest increase in Ge–Ge bond distance between the neutral and cationic structures. Having the weaker Ge–Ge bond of the two, the germylidene structure also has a larger increase in Ge–Ge bond length than the *trans* structure at 0.137 Å compared to 0.116 Å.

Changes in Ge–H bond lengths between the cation and the neutral structures were modest. An observed trend was the increase in Ge–H distances from the neutral to cationic species, with exceptions in the $\text{Ge}_{(2)}-\text{H}_{(2)}$ bond of the monobridged structure and the Ge–H bonds of the germylidene structure. Likewise, changes in the angles between the germanium and hydrogen atoms were also quite small when comparing the cation and neutral structures. Nonetheless, the most significant change was in the butterfly τ (H–Ge–Ge–H) angle, which increased from 105.0 in the neutral structure to 108.5 in the cation.

Energetics

Focal point analysis with harmonic ZPVE and aZPVE corrections for the neutral Ge_2H_2 species is shown in Table 2 and for the cationic species in Table 3. CCSD(T)/cc-pwCV5Z-PP single point energies were extrapolated to the CBS limit with CCSDT and CCSDT(Q) corrections. Tables 4–7 contain the incremented focal point energies for each structure, relative to its respective cationic or neutral butterfly structure, considered in this study.

As seen in previous theoretical work on Si_2H_2 ,^{20,21} Si_2H_2^+ ,⁴³ and Ge_2H_2 ,^{22,23} the butterfly structure is likewise the Ge_2H_2^+ global minimum geometry. Unlike the neutral Ge_2H_2 species, the Ge_2H_2^+ germylidene structure lies very slightly lower in energy than the monobridged structure. This is also seen in Si_2H_2^+ compared to neutral Si_2H_2 . In Si_2H_2^+ , the relative energy of the vinylidene-like structure is 6.61 kcal mol^{−1} and the monobridged structure is 10.31 kcal mol^{−1}.⁴³ The germylidene and monobridged structures have respective relative energies of 9.83 and 9.87 kcal mol^{−1} for Ge_2H_2^+ , while the neutral Ge_2H_2 germylidene and monobridged structures have respective relative energies of 13.47 and 9.67 kcal mol^{−1}. An enthalpy of formation for the monobridged structure was determined by Kaiser and coworkers using TOF-MS. However, because they did not also measure the butterfly structure, the relative energy of the monobridged structure cannot be obtained. Nonetheless, a relative energy of 8.4 kcal mol^{−1} was predicted by Kaiser and coworkers at the CCSD(T)/CBS//CCSD/cc-pVTZ level of theory. The geometry obtained by Kaiser and coworkers has little deviation from our work, therefore, the disagreement is ascribed to our higher-order corrections and pseudopotential-containing basis set family. We include quintuple-zeta polarized basis functions in our extrapolation to the CBS limit along with full T , perturbative Q , and anharmonic corrections to the ZPVE whereas Kaiser and coworkers do not. Considering these additional corrections, located in Tables 2 and 4, have little effect on the total energy, the difference in energy between our

Table 2 CCSD(T)/CBS energies with CCSD(T)/cc-pwCVTZ-PP harmonic ZPVE and anharmonic corrections computed for the neutral Ge_2H_2 species in kcal mol^{−1}. See text for higher-level theoretical corrections

Structure	CCSD(T)/CBS	δZPVE	δaZPVE	Total energy
Monobridged	10.44	−0.80	0.03	9.67
Germylidene	13.77	−0.32	0.02	13.47
<i>trans</i>	20.23	−0.93	0.14	19.44
Linear	49.70	—	—	49.70

Table 3 CCSD(T)/CBS energies with CCSD(T)/cc-pwCVTZ-PP harmonic ZPVE and anharmonic corrections computed for the cationic Ge_2H_2^+ species in kcal mol^{−1}. See text for higher-level theoretical corrections

Structure	CCSD(T)/CBS	δZPVE	δaZPVE	Total energy
Germylidene	9.78	0.04	0.01	9.83
Monobridged	10.77	−0.93	0.03	9.87
<i>trans</i>	20.73	−0.98	−0.07	19.68
Linear	56.31	—	—	56.31



Table 4 Incremented focal point energies in kcal mol⁻¹ for the cationic and neutral monobridged structures

Monobridged	HF	+δ MP2	+δ CCSD	+δ CCSD(T)	+δ CCSDT	+δ CCSDT(Q)	Net
Cation							
cc-pwCVDZ-PP	+8.54	+2.90	+0.26	-0.39	-0.03	-0.05	[+11.23]
cc-pwCVTZ-PP	+8.74	+2.91	-0.32	-0.27	[−0.03]	[−0.05]	[+10.98]
cc-pwCVQZ-PP	+8.73	+2.82	-0.36	-0.26	[−0.03]	[−0.05]	[+10.85]
cc-pwCV5Z-PP	+8.74	+2.74	-0.32	-0.25	[−0.03]	[−0.05]	[+10.82]
CBS limit	[+8.74]	[+2.66]	[−0.29]	[−0.25]	[−0.03]	[−0.05]	[+10.77]
Neutral							
cc-pwCVDZ-PP	+14.70	-4.49	+1.84	-1.51	+0.10	-0.14	[+10.50]
cc-pwCVTZ-PP	+14.58	-4.30	+1.67	-1.52	[+0.10]	[−0.14]	[+10.40]
cc-pwCVQZ-PP	+14.57	-4.29	+1.72	-1.55	[+0.10]	[−0.14]	[+10.41]
cc-pwCV5Z-PP	+14.61	-4.33	+1.77	-1.56	[+0.10]	[−0.14]	[+10.44]
CBS limit	[+14.61]	[−4.37]	[+1.81]	[−1.57]	[+0.10]	[−0.14]	[+10.44]

Table 5 Incremented focal point energies in kcal mol⁻¹ for the cationic and neutral germylidene structures

Germylidene	HF	+δ MP2	+δ CCSD	+δ CCSD(T)	+δ CCSDT	+δ CCSDT(Q)	Net
Cation							
cc-pwCVDZ-PP	+4.45	+14.25	-2.54	+1.55	-0.05	-0.00	[+9.26]
cc-pwCVTZ-PP	-3.81	+15.60	-3.98	+2.02	[−0.05]	[−0.00]	[+9.78]
cc-pwCVQZ-PP	-3.82	+15.62	-4.15	+2.14	[−0.05]	[−0.00]	[+9.74]
cc-pwCV5Z-PP	-3.81	+15.58	-4.14	+2.18	[−0.05]	[−0.00]	[+9.76]
CBS limit	[−3.81]	[+15.54]	[−4.12]	[+2.22]	[−0.05]	[−0.00]	[+9.78]
Neutral							
cc-pwCVDZ-PP	+6.70	+9.04	-3.31	+0.27	-0.12	-0.08	[+12.50]
cc-pwCVTZ-PP	+6.92	+10.50	-4.40	+0.67	[−0.12]	[−0.08]	[+13.49]
cc-pwCVQZ-PP	+6.88	+10.66	-4.50	+0.76	[−0.12]	[−0.08]	[+13.61]
cc-pwCV5Z-PP	+6.90	+10.68	-4.47	+0.78	[−0.12]	[−0.08]	[+13.70]
CBS limit	[+6.91]	[+10.69]	[−4.44]	[+0.80]	[−0.12]	[−0.08]	[+13.77]

Table 6 Incremented focal point energies in kcal mol⁻¹ for the cationic and neutral *trans* structures

<i>trans</i>	HF	+δ MP2	+δ CCSD	+δ CCSD(T)	+δ CCSDT	+δ CCSDT(Q)	Net
Cation							
cc-pwCVDZ-PP	+18.47	+3.78	-0.35	-1.15	-0.17	-0.13	[+20.46]
cc-pwCVTZ-PP	+19.06	+4.22	-1.29	-0.94	[−0.17]	[−0.13]	[+20.76]
cc-pwCVQZ-PP	+19.02	+4.21	-1.31	-0.92	[−0.17]	[−0.13]	[+20.71]
cc-pwCV5Z-PP	+19.03	+4.16	-1.25	-0.92	[−0.17]	[−0.13]	[+20.72]
CBS limit	[+19.03]	[+4.11]	[−1.20]	[−0.92]	[−0.17]	[−0.13]	[+20.73]
Neutral							
cc-pwCVDZ-PP	+26.11	-5.86	+1.96	-2.66	+0.02	-0.30	[+19.28]
cc-pwCVTZ-PP	+26.22	-5.02	+1.51	-2.50	[+0.02]	[−0.30]	[+19.94]
cc-pwCVQZ-PP	+26.24	-4.95	+1.59	-2.53	[+0.02]	[−0.30]	[+20.07]
cc-pwCV5Z-PP	+26.28	-4.95	+1.65	-2.55	[+0.02]	[−0.30]	[+20.16]
CBS limit	[+26.30]	[−4.95]	[+1.72]	[−2.56]	[+0.02]	[−0.30]	[+20.23]

work and the work of Kaiser and coworkers should be attributed to the difference in basis set family.

As noted in the 1993 Ge_2H_2 study by Pálágyi and Schaefer, the monobridged structure is favored by correlation effects whereas the germylidene structure is disfavored.²³ Our work is in agreement with their previous observations, and it can be seen in Table 4 for the monobridged structure and in Table 5 for the germylidene structure. Both the monobridged and germylidene structures of Ge_2H_2^+ are disfavored by correlation effects, although the monobridged structure less so. The Ge_2H_2^+ δMP2 energies at the CBS limit for the monobridged

and germylidene isomers are +2.66 and +15.54 kcal mol⁻¹, respectively. However, considering the CCSDT(Q) energy correction is 0.00 kcal mol⁻¹ for the germylidene structure and -0.05 kcal mol⁻¹ for the monobridged structure, it is unlikely that higher levels of correlation beyond the scope of this study would continue to affect the energies of the monobridged and germylidene structures of Ge_2H_2^+ .

Highest in relative energy of the minimum stationary points studied, the *trans* structure has respective relative energies of 20.23 and 20.73 kcal mol⁻¹ for the neutral and cationic species. Similar to the monobridged structure, the neutral *trans*



Table 7 Incremented focal point energies in kcal mol⁻¹ for the cationic and neutral linear structures

Linear	HF	+δ MP2	+δ CCSD	+δ CCSD(T)	+δ CCSDT	+δ CCSDT(Q)	Net
Cation							
cc-pwCVDZ-PP	+55.69	+5.24	+0.77	-1.15	+0.13	-0.24	[+60.45]
cc-pwCVTZ-PP	+55.09	+4.49	-0.75	-1.04	[+0.13]	[−0.24]	[+57.68]
cc-pwCVQZ-PP	+54.94	+3.98	-0.89	-1.02	[+0.13]	[−0.24]	[+56.89]
cc-pwCV5Z-PP	+54.92	+3.60	-0.81	-1.01	[+0.13]	[−0.24]	[+56.60]
CBS limit	[+54.92]	[+3.21]	[-0.72]	[-1.01]	[+0.13]	[−0.24]	[+56.31]
Neutral							
cc-pwCVDZ-PP	+55.85	-3.83	+3.07	-2.38	+0.22	-0.31	[+52.62]
cc-pwCVTZ-PP	+54.89	-4.09	+2.22	-2.38	[+0.22]	[−0.31]	[+50.55]
cc-pwCVQZ-PP	+54.90	-4.41	+2.16	-2.42	[+0.22]	[−0.31]	[+50.15]
cc-pwCV5Z-PP	+54.94	-4.72	+2.23	-2.42	[+0.22]	[−0.31]	[+49.94]
CBS limit	[+54.95]	[-5.04]	[+2.31]	[-2.43]	[+0.22]	[−0.31]	[+49.70]

structure is favored by correlation effects and the cationic structure is disfavored. This can be seen in the δ MP2 contribution, which is -4.95 kcal mol⁻¹ for the neutral structure but $+4.11$ kcal mol⁻¹ in the cationic structure at the CBS limit. In addition, the neutral *trans* structure has the largest energetic contribution due to including perturbative triple excitations in CCSD(T), which is -2.56 kcal mol⁻¹ at the CBS limit. In the cationic *trans* structure, the CCSD(T) contribution is a more modest -0.92 kcal mol⁻¹. The incremented focal point energies for the *trans* structures are contained in Table 6.

Although the acetylene linear structure is the commonly known C_2H_2 minimum geometry, it is a transition state in Si_2H_2 , Ge_2H_2 , and their cations. Not appearing to be influenced much by correlation effects until higher levels of electronic excitation, the linear Ge_2H_2 structures have the largest CCSDT(Q) correction in this study at -0.24 and -0.31 for the respective cationic and neutral structures. However, when considering the magnitude of the relative energies of these Ge_2H_2 linear structures, the (Q) corrections are not so important. Table 7 contains the incremented focal point energies for the linear structures in kcal mol⁻¹.

Vibrational frequencies

Fundamental vibrational frequencies computed at the CCSD(T)/cc-pwCVTZ-PP level of theory for the neutral Ge_2H_2 minimum species are presented in Table 8 and for the cationic species in Table 9. Harmonic vibrational frequencies and

Table 8 CCSD(T)/cc-pwCVTZ-PP fundamental vibrational frequencies in cm⁻¹ and IR intensities in km mol⁻¹ are reported for the neutral minimum structures

Butterfly	Monobridged	Germylidene	<i>trans</i>
C_{2v}	C_s	C_{2v}	C_{2h}
284 (<1, a ₁)	185 (34, a'')	208 (19, b ₂)	220 (0, b _u)
790 (31, a ₁)	307 ^a (5, a')	292 (5, b ₁)	243 (0, a _g)
899 (0, a ₂)	479 (3, a')	305 (6, a ₁)	393 (<1, a _u)
991 (352, b ₂)	916 (112, a')	822 (63, a ₁)	576 (0, a _g)
1362 (32, b ₁)	1494 ^b (75, a')	2036 (74, a ₁)	1975 (0, a _g)
1427 (8, a ₁)	1986 (120, a')	2060 (82, b ₂)	1988 (176, b _u)

^a Member of Fermi dyad: other transition at 383 cm⁻¹. ^b Member of Fermi dyad: other transition at 1370 cm⁻¹.

Table 9 CCSD(T)/cc-pwCVTZ-PP fundamental vibrational frequencies in cm⁻¹ and IR intensities in km mol⁻¹ are reported for the cationic minimum structures

Butterfly	Monobridged	Germylidene	<i>trans</i>
C_{2v}	C_s	C_{2v}	C_{2h}
189 (<1, a ₁)	166 (5, a'')	253 (2, a ₁)	78 (12, a _u)
759 (18, a ₁)	256 (8, a')	260 (7, b ₂)	125 (39, b _u)
922 (0, a ₂)	474 (4, a')	376 (<1, b ₁)	252 (0, a _g)
1039 (457, b ₂)	834 ^a (126, a')	835 (38, a ₁)	588 (0, a _g)
1340 (31, b ₁)	1429 (38, a')	2074 (2, a ₁)	1989 (0, a _g)
1434 (2, a ₁)	2011 (3, a')	2111 (9, b ₂)	2000 (11, b _u)

^a Member of Fermi dyad: other transition at 949 cm⁻¹.

vibration–rotation constants are located in the ESI.[†] As a second order transition state with a doubly degenerate π_g bend, anharmonic treatment of the linear structure was not performed. Containing the largest IR intensity vibrational mode, the butterfly structure has been previously observed *via* matrix isolation in neon and argon by Wang, Andrews, and Kushto in 2002.¹⁷ Wang, Andrews, and Kushto identified the b_2 absorption band of the butterfly structure, having a frequency of 972.3 cm⁻¹ in neon and 964.3 cm⁻¹ in argon, by comparing their results to the previous theoretical study by Pálágyi and Schaefer in 1993.²³ Herein, we present the most accurate theoretical prediction to date, calculating the b_2 absorption band to have a frequency of 991 cm⁻¹. Having by far the largest IR intensity in the neutral and cationic species at 352 and 457 km mol⁻¹, respectively, this band corresponds to the conjoined Ge–H stretch of the hydrogen atoms parallel to the Ge–Ge bond. As the nuclei are displaced along this vibrational mode, their positions begin to resemble that of the germylidene structure. Given that the dipole moments of the butterfly and germylidene structures are perpendicular to one another, the b_2 vibrational mode of the butterfly structure constitutes a significant change in the dipole moment, resulting in a large IR intensity. Also identified in the 2002 paper by Wang, Andrews, and Kushto was the b_1 band, at a frequency of 1352.3 cm⁻¹.¹⁷ Our computations predict this band to be at 1362 cm⁻¹ and it corresponds to the non-conjoined Ge–H stretch perpendicular to the Ge–Ge bond.

All of the remaining unidentified Ge_2H_2 and Ge_2H_2^+ structures contain detectable vibrational modes in the IR region.



The respective 916 and 840 cm^{-1} a' modes of the neutral and cationic monobridged structure have intensities of 112 and 126 km mol^{-1} , which correspond to the Ge–H stretch of the bridged hydrogen between the two germanium atoms. Although the cationic butterfly structure also contains a mode at 916 cm^{-1} , this non-conjoined Ge–H stretching mode has no effect on the dipole and is therefore undetectable in the IR region. The 2036 cm^{-1} symmetric and 2060 cm^{-1} asymmetric Ge–H stretches have the greatest IR intensities in the neutral germylidene structure at respective values of 74 and 82 km mol^{-1} , but the intensities of these two modes decrease dramatically in the cationic structure to 2 and 9 km mol^{-1} . Both the neutral and cationic germylidene structures have measurable a_1 Ge–H in-plane wagging modes, respectively at 822 and 835 cm^{-1} with intensities of 63 and 38 km mol^{-1} . Measuring the neutral *trans* structure in the IR region is more likely than the cationic *trans* structure due to the 1988 cm^{-1} b_u Ge–H stretching mode with an intensity of 176 km mol^{-1} . This mode shifts to 2000 cm^{-1} and decreases to an intensity of 11 km mol^{-1} in the cationic *trans* structure. However, the 125 cm^{-1} b_u in-plane wagging mode has a substantial intensity of 39 km mol^{-1} in the cationic *trans* structure, whereas this vibrational mode has a near-zero IR intensity in the neutral *trans* structure. Table 10 reports, in Debye, vibrationally averaged zero-point dipole moments originating from the center of mass. Fig. 2 depicts the dipole moments of the neutral structures and Fig. 3 the dipole moments of the cationic structures. The dipole moment of the *trans* and linear structures are necessarily zero.

Similar to carbon dioxide, in which the dipole moment points to carbon, the dipole moment of the neutral and cationic germylidene species point toward the less electronegative germanium atoms. However, the combination of a Ge–Ge double bond and a lone pair on $\text{Ge}_{(1)}$ supplements the direction of the dipole moment in the germylidene structure. Likewise, the dipole moment of the neutral butterfly structure points in the

Table 10 Vibrationally averaged zero-point dipole moments originating from the center of mass are given in Debye

Geometry	Cation	Neutral
Butterfly	0.20	0.70
Monobridged	0.43	0.29
Germylidene	0.21	0.51

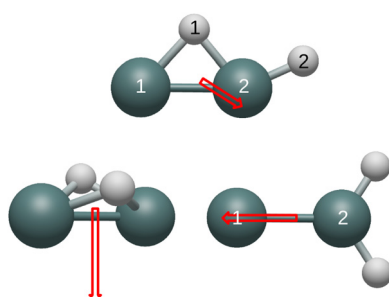


Fig. 2 Dipole moments originating from the center of mass of the neutral structures are displayed.

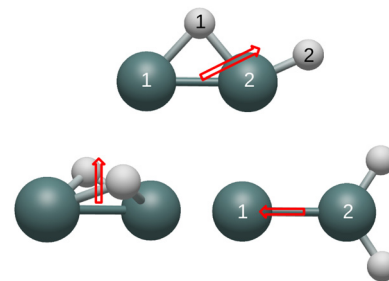


Fig. 3 Dipole moments originating from the center of mass of the cationic structures are displayed.

direction of the Ge–Ge bond. Containing a lone pair on both germanium atoms, the neutral butterfly structure also contains sufficient electron density within and around the Ge–Ge bond for the dipole moment to point to the less electronegative germanium atoms. On the other hand, the cationic butterfly structure has a dipole moment pointing to the hydrogen atoms, opposite of the neutral butterfly structure. The singly-occupied molecular orbital (SOMO) of the butterfly structure is the sole Ge–Ge bond; having one less electron in this molecular orbital is enough to flip the direction of the dipole moment from germanium to hydrogen. With both a Ge–Ge double bond and a lone pair on $\text{Ge}_{(1)}$, it is unclear why the dipole moment of the neutral monobridged structure points in the direction of $\text{Ge}_{(2)}$. As also seen in the butterfly structure, the dipole moment of the cationic monobridged structure changes direction from germanium to hydrogen in its neutral counterpart.

Natural bonding orbital analysis

Natural bond orders are reported in Table 11 and partial charges in Table 12. The electron removed from neutral Ge_2H_2 is primarily a Ge–Ge bonding electron as suggested by the decrease in bond order and the distribution of positive charge. Already containing a partial positive charge in the neutral species, the germanium atoms unsurprisingly gain more positive charge in the cationic species. Also considering the size, ionization potential, and electronegativity of the

Table 11 HF/def2-TZVP natural bond orders computed at CCSD(T)/cc-pwCVQZ-PP optimized geometries

Geometry	Bond	Cation	Neutral
Butterfly	Ge–Ge	0.79	1.31
	Ge–H	0.43	0.42
Monobridged	Ge–Ge	1.59	2.23
	$\text{Ge}_{(1)}\text{–H}_{(1)}$	0.37	0.36
	$\text{Ge}_{(2)}\text{–H}_{(1)}$	0.49	0.48
	$\text{Ge}_{(2)}\text{–H}_{(2)}$	0.91	0.90
Germylidene	Ge–Ge	1.48	1.99
	Ge–H	0.95	0.96
<i>trans</i>	Ge–Ge	2.01	2.60
	Ge–H	0.76	0.80
Linear	Ge–Ge	2.40	2.97
	Ge–H	0.95	0.95



Table 12 HF/def2-TZVP partial charges computed at CCSD(T)/cc-pwCVQZ-PP optimized geometries

Geometry	Atom	Cation	Neutral
Butterfly	Ge	+0.83	+0.27
	H	-0.33	-0.27
Monobridged	Ge ₍₁₎	+0.79	+0.22
	Ge ₍₂₎	+0.61	+0.11
	H ₍₁₎	-0.11	-0.14
	H ₍₂₎	-0.28	-0.19
Germlylidene	Ge ₍₁₎	+0.70	+0.24
	Ge ₍₂₎	+0.57	+0.07
	H	-0.13	-0.15
trans	Ge	+0.67	+0.18
	H	-0.17	-0.18
Linear	Ge	+0.52	+0.07
	H	-0.02	-0.07

germanium and hydrogen atoms, chemical intuition would also suggest this location.⁴⁴

Ge–H bond orders have small changes between the cationic and neutral structures. If the positive charge of Ge_2H_2^+ is located completely within Ge–Ge bonding orbitals, a decrease of a half of a bond order could be expected for the Ge–Ge bond. Slightly larger decreases than half of a bond order are seen in these five structures when comparing the cationic to the neutral species.

Structures containing the most Ge–H bonds might be assumed to have the weakest Ge–Ge bond order. However, this is not the case when considering the lower Ge–Ge bond order in the germlylidene structure compared to the monobridged structure. Summing the Ge–H bond orders in the germlylidene and monobridged structures, respectively 1.92 and 1.74 in the cations and 1.90 and 1.77 in the neutral structures, illustrates why the additional Ge–H bond in the monobridged structure does not necessitate the Ge–Ge bond to be weaker than that of the germlylidene structure. Due to the asymmetric distribution of Ge–H bonds in the germlylidene and monobridged structures, the positive charge is not equally shared between the two germanium atoms. In both cases, the germanium with fewer bonds to hydrogen has the lower partial positive charge.

Because the hydrogens are symmetrically bound about the germanium atoms of the butterfly, *trans*, and linear structures, the respective germanium atoms of these structures contain equal partial positive charges. Having the most Ge–H bonds (four) of the five structures considered, the butterfly structure also has the lowest Ge–Ge bond order in both the cationic and neutral species. The bond order of the bridged hydrogen atoms in the butterfly structure lies directly in between the $\text{Ge}_{(1)}\text{H}_{(1)}$ and $\text{Ge}_{(2)}\text{H}_{(1)}$ bond orders of the bridged hydrogen in the monobridged structure for both the cationic and neutral species. Similar to the butterfly structure, the *trans* and linear structures equally share the partial positive charge between the two germanium atoms and exhibit respective decreases of 0.59 and 0.57 in Ge–Ge NBO from the neutral to cationic structures.

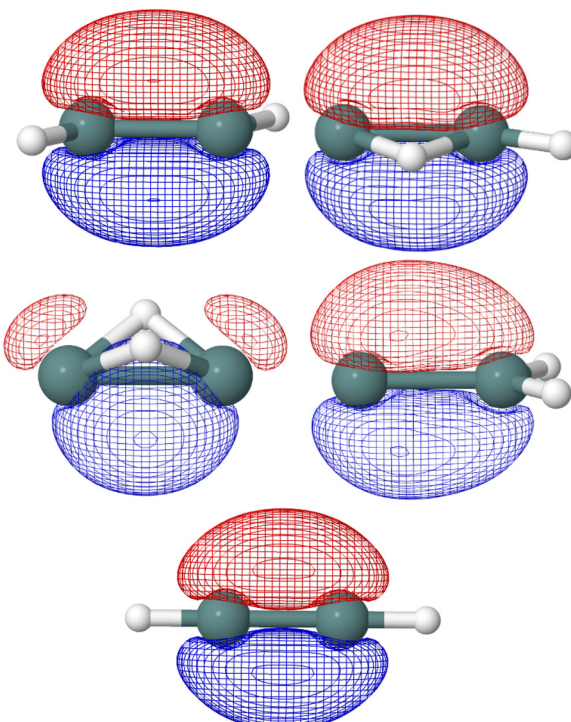


Fig. 4 HF/def2-TZVP singly-occupied molecular orbitals of the Ge_2H_2^+ structures computed at CCSD(T)/cc-pwCVQZ-PP optimized geometries are shown. From left to right and top to bottom: *trans* (C_{2h} ; $2A_1$), monobridged (C_s ; $2A''$), butterfly (C_{2v} ; $2A_1$), germlylidene (C_{2v} ; $2B_2$), and linear ($D_{\infty h}$; $2\Pi_u$).

In the five Ge_2H_2 structures considered, the highest occupied molecular orbital (HOMO) in the neutral species is the SOMO of the respective cationic species. As seen in Fig. 4, the SOMOs of the *trans* (C_{2h} ; $2A_1$), monobridged (C_s ; $2A''$), germlylidene (C_{2v} ; $2B_2$), and linear ($D_{\infty h}$; $2\Pi_u$) structures have almost entirely p-character. The butterfly (C_{2v} ; $2A_1$) structure is the only exception, as the SOMO has 90 percent p-character, 8 percent s-character, and 2 percent d-character. Knowing the linear structure has a triple Ge–Ge bond, first and foremost a σ -bonding molecular orbital is expected but also a degenerate π -bonding orbital to the SOMO. NBO analysis confirms this, as the HOMO of the neutral linear structure (SOMO in the cation) has a degenerate, 99 percent p-character π -bonding orbital. NBO analysis also indicates the *trans* structure to have three Ge–Ge bonds, with an NBO of 2.60. One could visualize the hydrogen of the *trans* structure sticking into where one of the degenerate Ge–Ge π -bonding orbitals of the linear structure was located, resulting in a π -like b_u orbital with 77 percent p-character and 22 percent s-character. In addition to the a'' π -character HOMO, the monobridged structure also has a σ -like a' Ge–Ge bonding orbital. $\text{Ge}_{(1)}$ contributes 29 percent of the a' Ge–Ge bond while $\text{Ge}_{(2)}$ contributes 71 percent. The contribution of $\text{Ge}_{(1)}$ is 90 percent p-character, 9 percent s-character while the contribution of $\text{Ge}_{(2)}$ is 59 percent s-character, 41 percent p-character. Similarly, the germlylidene structure has a σ -like Ge–Ge bonding orbital, which is of a_1 symmetry. Because $\text{Ge}_{(2)}$ is bound to both hydrogen atoms,



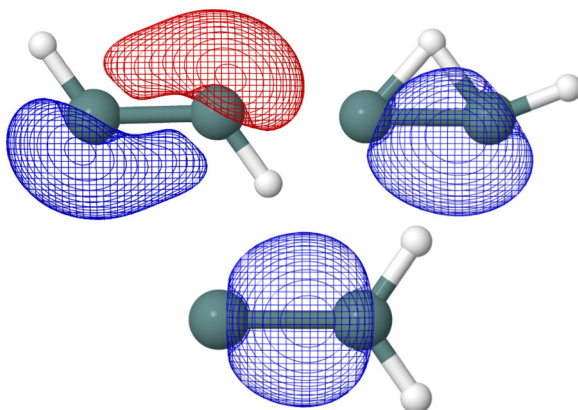


Fig. 5 *Trans* b_u , monobridged a' , and germylidene a_1 Ge–Ge bonding orbitals computed at HF/def2-TZVP are shown.

it may be surprising that $Ge_{(2)}$ contributes 62 percent of the a_1 Ge–Ge bond. However, the lone pair orbital on $Ge_{(1)}$ lies only slightly higher in energy than the a_1 Ge–Ge bonding orbital, suggesting a propensity for electronic localization around $Ge_{(1)}$. The contribution of $Ge_{(1)}$ to the a_1 Ge–Ge bonding orbital of the germylidene structure is 14 percent s-character, 85 percent p-character and the contribution of $Ge_{(2)}$ is 42 percent s-character, 58 percent p-character. Fig. 5 shows the *trans* b_u , monobridged a' , and germylidene a_1 Ge–Ge bonding orbitals.

Conclusions

Using state-of-the-art *ab initio* methods, it is determined that the relative ordering in energy for these five structures of $Ge_2H_2^+$ is butterfly, germylidene, monobridged, *trans*, then linear. The ordering in the neutral species is different, as the monobridged structure has a significantly lower energy than that of the germylidene structure. Analysis of NBOs and partial charges revealed the positive charge is localized between the germanium atoms. As the first characterization of these five $Ge_2H_2^+$ structures, this study should serve to instigate future experimental work.

Conflicts of interest

There are no conflicts to declare.

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