# Infrared Spectra of Thallium Hydrides in Solid Neon, Hydrogen, and Argon

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Laser-ablated Tl atoms react with dihydrogen in excess neon, pure hydrogen, and excess argon to form primarily the TlH diatomic molecule. Ultraviolet irradiation increases weak bands that are identified as TlH<sub>2</sub> and TlH<sub>3</sub> on the basis of D<sub>2</sub> and HD substitution and density functional theory (DFT) isotopic frequency calculations. Sample annealing fosters the dimerization of TlH to give Tl<sub>2</sub>H<sub>2</sub> with a rhombic ring structure and the TlTlH<sub>2</sub> isomer. The markedly lower yield of TlH<sub>3</sub> in these experiments compared to AlH<sub>3</sub> in earlier investigations is due to the expected decrease in stability for the Tl(III) oxidation state. The Tl<sup>+</sup>(H<sub>2</sub>)<sub>n</sub> cation complex is also observed, and trends within the group 13 metals are summarized. Excitation at 193 nm gives the  ${}^{2}S_{1/2} \rightarrow$  ${}^{2}P_{1/2,3/2}$  doublet for unreacted Tl atoms, which is blue-shifted 660 cm<sup>-1</sup> from the gas-phase value in solid D<sub>2</sub> but only 210 cm<sup>-1</sup> in solid H<sub>2</sub>, owing to the smaller, more repulsive D<sub>2</sub> matrix cage.

### Introduction

Small group 13 metal hydrides have been investigated extensively (M = Al, Ga, or In), including by gas-phase spectroscopy for the monohydride diatomics<sup>1-9</sup> and matrix infrared studies for the mono-, di-, and trihydrides, which reveal increasing frequencies in the MH<sub>1,2,3</sub> series for each metal.<sup>10-18</sup> Electron spin resonance has been employed to investigate the AlH<sub>2</sub> radical in solid neon.<sup>19</sup> The M<sub>2</sub>H<sub>2</sub> and M<sub>2</sub>H<sub>6</sub> species have been characterized by increasing stability of the former and decreasing stability of the latter with the heavier metals. Thallium hydride, TlH, has only been investigated in the gas phase,<sup>1,20</sup> and by theoretical calculations,<sup>21-24</sup> and there is no experimental evidence to date for TlH<sub>2</sub> and TlH<sub>3</sub>, although the latter have also been subjected to theoretical calculations as heavy metal compounds, owing to interest in relativistic effects.<sup>25,26</sup> The trend of decreasing stability for the heavier metal trihydride is manifest in the lower yield of InH<sub>3</sub> relative to AlH<sub>3</sub>,<sup>18</sup> and this points to difficulty for the preparation of TlH<sub>3</sub>.

### **Experimental and Theoretical Methods**

The laser-ablation matrix-isolation infrared spectroscopy experiment has been described previously.<sup>27,28</sup> Briefly, the Nd: YAG laser fundamental (1064 nm, 10 Hz repetition rate, 10 ns pulse width) was focused on rotating a high purity thallium (Spex Industries, 99.999%) target using 5-20 mJ/pulse, and thallium atoms were co-deposited with pure H<sub>2</sub> or D<sub>2</sub> (Matheson), pure HD (Cambridge Isotope Laboratories), and neon or argon diluted samples onto a CsI window cooled to 3.5 K by a Sumitomo Heavy Industries RDK-205D cryocooler. Infrared spectra were recorded in a Nicolet 750 Fourier transform instrument at 0.5 cm<sup>-1</sup> resolution using a liquid-nitrogen cooled MCTB detector after deposition, annealing, and filtered mercury arc or 193 nm laser (Lambda Physik, Optex) irradiation. Emission spectra were recorded using an Ocean Optics optical fiber spectrometer.



**Figure 1.** Infrared spectra in the 1780–1260 cm<sup>-1</sup> region for normal hydrogen co-deposited at 3.5 K with laser-ablated thallium. Spectra of H<sub>2</sub> + Tl: (a) before irradiation, (b) after 15 min of  $\lambda > 240$  nm irradiation, (c) after 15 min more of  $\lambda > 240$  nm irradiation, (d) after annealing to 6.0 K, and (e) after annealing to 6.5 K.

Density functional theory (DFT) frequency calculations were helpful in assigning lead and bismuth hydride spectra,<sup>29,30</sup> so similar B3LYP/6-311++G\*\*/Stuttgart scalar relativistic pseudo-potential for Tl together with the corresponding basis set (SDD) calculations were performed for thallium hydrides using the Gaussian 98 program system.<sup>31–33</sup> Comparable results were obtained using the BPW91 functional.<sup>34</sup> Although these calculations are only approximate, they provide a useful guide for assigning vibrational spectra.

## Results

Reaction products from laser-ablated Tl and dihydrogen trapped in excess hydrogen, neon, and argon and supporting DFT calculations will be presented.

**Hydrogen.** Laser-ablated Tl co-deposited with hydrogen (deuterium) produces weak bands at 1311 (940) cm<sup>-1</sup> that increase upon ultraviolet irradiation to strong bands at 1311.3 (939.9) cm<sup>-1</sup>, as shown in Figure 1 (Figure 2). This treatment

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**Figure 2.** Infrared spectra in the 1300–900 cm<sup>-1</sup> region for normal deuterium co-deposited at 3.5 K with laser-ablated thallium. Spectra of  $D_2 + Tl$ : (a) before irradiation, (b) after 15 min of  $\lambda > 240$  nm irradiation, (c) after 15 min more of  $\lambda > 240$  nm irradiation, (d) after annealing to 8.0 K, and (e) after annealing to 9.0 K.

also yields new absorptions at 1519.9 (1098.8) and 1390.2 (1007.6)  $\text{cm}^{-1}$  and weak bands at 1748.4 (1254.4)  $\text{cm}^{-1}$ . Annealing increases the strong bands and new features at 1424.3 (1020.5)  $\text{cm}^{-1}$ . Weak bands also appear upon annealing at 908.8 (653.1)  $\text{cm}^{-1}$  that are produced in greater yield using neon host matrixes. Table 1 lists the observed frequencies.

The HD experiment provides important diagnostic information on the thallium hydrides in Figure 3. The strong initial bands are observed slightly shifted in the pure HD host at 1320.1 and 945.9 cm<sup>-1</sup>. The sharp 1519.9 (1098.8) and 1390.2 (1007.6) cm<sup>-1</sup> bands are replaced by a new set at 1461.8 and 1046.2 cm<sup>-1</sup>. The weaker 1748.4 (1254.4) cm<sup>-1</sup> absorptions shift to 1747.3 and 1258.1 cm<sup>-1</sup>. Finally, a weak 839.0 cm<sup>-1</sup> band is observed instead of the 908.8 (653.1) cm<sup>-1</sup> pair.

The higher frequency region for hydrogen (deuterium) reveals sharp new 4113.6 (2962.8) cm<sup>-1</sup> absorptions and other metal independent absorptions observed previously at 4143.4 (2982.4)



**Figure 3.** Infrared spectra in the 1800–800 cm<sup>-1</sup> region for pure HD co-deposited at 3.5 K with laser-ablated thallium. Spectra of HD + TI: (a) before irradiation, (b) after 30 min of  $\lambda > 240$  nm irradiation, (c) after annealing to 7.3 K, (d) after 15 min of  $\lambda > 240$  nm irradiation, and (e) after annealing to 8.0 K.

and 3972 (2870) cm<sup>-1.16-18</sup> The spectra in Figure 4 show that  $\lambda > 240$  nm irradiation effectively destroys all product bands in this region. However, in the more robust deuterium lattice, the 2870 cm<sup>-1</sup> absorption for (D<sup>-</sup>)(D<sub>2</sub>)<sub>n</sub> is regenerated in part by 193 nm irradiation: this destruction–regeneration cycle is repeated. Likewise, the higher frequency region with HD shows analogous sharp bands at 3621.7 and 3596.3 cm<sup>-1</sup> and a broader feature at 3478 cm<sup>-1</sup>. The sharp 3621.7 cm<sup>-1</sup> and broad 3478 cm<sup>-1</sup> bands are common to other metal experiments with HD.

**Neon.** Weak bands are observed at 1327 (950) cm<sup>-1</sup> for H<sub>2</sub> (D<sub>2</sub>) in excess neon co-deposited with Tl atoms. Spectra in Figure 5 compare deposited samples after deposition and 193 nm irradiation (5 min at 4 mJ/pulse). Stronger bands at 919.8 (667.3) cm<sup>-1</sup> acquire a 851.1 cm<sup>-1</sup> HD counterpart. Irradiation also produces weaker 1525.9 (1103.5) cm<sup>-1</sup> bands with a 1468 cm<sup>-1</sup> HD counterpart.

<b>IABLE I:</b> Infrared Absorptions (cm <sup>-1</sup> ) Observed from Reaction of Inallium and Dinvdrog	en in Neor	i. Hvarogen	and Argon.
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	neon			hydrogen			argon		
H <sub>2</sub>	HD	D <sub>2</sub>	H <sub>2</sub>	HD	D <sub>2</sub>	H <sub>2</sub>	HD	D <sub>2</sub>	identity
			4113.6	3596.3	2962.8				$Tl^+(H_2)_n$
			1748.4	1747.3					TlH <sub>3</sub>
				1259.9	1254.4				$TlD_3$
			1700.2						anneal pdt
					1224.3				anneal pdt
			1522.4						TlH <sub>2</sub> (site)
1525.9			1519.9			1503.3			$TlH_2$
	1468			1461.8, 1046.2					TIHD
		1112.3			1101.4				TlD <sub>2</sub> (site)
		1103.5			1098.8			1082.0	$TlD_2$
			1394.7						TlH <sub>2</sub> (site)
			1390.2						$TlH_2$
					1010.3				TlD <sub>2</sub> (site)
					1007.6				$TlD_2$
			1430.3	1430.8					TlTlH <sub>2</sub> (site)
1431.7			1424.3	1424.8			1427.6		$TITIH_2$
				1424.2, 1423.0					TITIHD
				1023.9, 1022.6					TITIHD
				1023.5	1021.6				TlTlD <sub>2</sub> (site)
				1027.7	1020.5				TITID <sub>2</sub>
			1328.0	1339.6					TlH (site)
1327	1327		1311.3	1320.1		1309.4	1309.4		TlH
				959.4	953.4		938.0		TlD (site)
	950	950		945.9	939.9			938.0	TlD
919.8			908.8			888.2	888.0		$Tl_2H_2$
	851.1			839.0			825.3		$Tl_2HD$
		667.3			653.1		642.8	642.6	$Tl_2D_2$



**Figure 4.** Infrared spectra in the H–H and D–D stretching regions for hydrogen and deuterium co-deposited at 3.5 K with thallium. Spectra of H<sub>2</sub> + TI: (a) before irradiation, (b) after 15 min of  $\lambda > 240$  nm irradiation, (c) after 15 min more of  $\lambda > 240$  nm irradiation, (d) after annealing to 6.0 K, and (e) after annealing to 6.5 K. Spectra of D<sub>2</sub> + TI: (f) before irradiation, (g) after 15 min of  $\lambda > 240$  nm irradiation, (h) after 3 min of 193 nm irradiation (5 mJ/pulse at 10 Hz), (i) after  $\lambda > 240$  nm irradiation.



**Figure 5.** Infrared spectra in the  $600-1550 \text{ cm}^{-1}$  region for hydrogen co-deposited at 3.5 K with thallium in neon. Spectra of 4% H<sub>2</sub> in neon + Tl: (a) before irradiation and (b) after 5 min of 193 nm (5 mJ/pulse at 10 Hz) irradiation. Spectra of 4% HD in neon + Tl: (c) before irradiation and (d) after 193 nm irradiation. Spectra of 2% H<sub>2</sub> + 2% D<sub>2</sub> in neon + Tl: (e) before irradiation and (f) after 193 nm irradiation. Spectra of 4% D<sub>2</sub> in neon + Tl: (g) before irradiation and (h) after 193 nm irradiation.

**Argon.** The weak argon matrix  $H_2$  ( $D_2$ ) reaction products at 1309.4 (938.0) cm<sup>-1</sup> increase markedly upon annealing to allow diffusion and reaction of trapped atomic species, as shown in Figure 6a,b and i,j. Additional bands appear at 888.2 (642.6) cm<sup>-1</sup> upon annealing. Using the HD reagent, the latter two bands plus a new 825.3 cm<sup>-1</sup> absorption are produced: Increased laser energy favors the yield of TIH and TID, as illustrated in Figure 6c-f. The Ar<sub>n</sub>H<sup>+</sup> and Ar<sub>n</sub>D<sup>+</sup> species were also trapped in solid argon.<sup>35,36</sup>

**Emission Spectra.** Emission spectra were recorded from the 1064 nm laser-generated plume above the target surface during ablation–deposition. The strong Tl atomic spin–orbit doublets<sup>37</sup> were observed at 377.7 and 535.2 nm and at 276.9 and 353.0 nm, as shown in Figure 7. In addition, weaker Tl<sup>+</sup> lines were found at 329.1, 507.7, 515.4, and 595.0 nm.<sup>38</sup> During 193 nm



**Figure 6.** Infrared spectra in the  $600-1550 \text{ cm}^{-1}$  region for hydrogen co-deposited at 3.5 K with thallium in argon. Spectra of 4% H<sub>2</sub> in argon + TI: (a) before annealing and (b) after annealing to 30 K. Spectra of 3% HD in argon + TI (10 mJ/pulse): (c) before annealing and (d) after annealing to 30 K. Spectra of 6% HD in argon + TI (30 mJ/pulse): (e) before annealing and (f) after annealing to 30 K. Spectra of 1.5% H<sub>2</sub> + 1.5% D<sub>2</sub> in argon + TI: (g) before annealing and (h) after annealing to 30 K. Spectra of 4% D<sub>2</sub> in argon + TI: (i) before annealing and (j) after annealing to 30 K.



Figure 7. Emission spectra from hydrogen-thallium experiments: (a) emission from the 1064 nm laser-ablation plume, (b) emission from solid  $D_2 + Tl$  using 193 nm excitation at 1 mJ/pulse, (c) emission from solid  $H_2 + Tl$  using 193 nm excitation at 1 mJ/pulse, (d) emission from neon + Tl using 193 nm excitation, and (e) emission from argon + Tl using 193 nm excitation.

irradiation of the  $H_2$  and  $D_2$  samples, strong Tl atomic doublets were observed at 374.7 and 526.2 nm and 368.5 and 513.8 nm, respectively. Similar 193 nm irradiation of neon and argon matrix samples containing Tl and  $H_2$  produced broader features and sharper bands for atomic Tl at 548 nm in the former and 393 and 549 nm in the latter.

### Discussion

New thallium hydride absorptions will be assigned on the basis of isotopic substitution, matrix shifts, and quantum chemical calculations.

**TIH.** Experiments with laser-ablated Tl and hydrogen (deuterium) gave weak bands at 1311.3 (939.9) cm<sup>-1</sup>, which increased markedly upon UV irradiation (Figures 1 and 2) and are slightly lower than the gas-phase TlH (TlD) fundamentals<sup>1,20</sup> of 1345.3 (963.7) cm<sup>-1</sup> and are due to the diatomic hydride molecules in solid H<sub>2</sub> (D<sub>2</sub>). These TlH (TlD) bands are bracketed by the argon matrix (1309.4 (938.0) cm<sup>-1</sup>) and neon matrix (1327 (950) cm<sup>-1</sup>) values. Two sharp bands were observed with pure HD, but both were blue-shifted 0.6% from the H<sub>2</sub> (D<sub>2</sub>) matrix values. Our simple B3LYP calculation predicts 1306 (926) cm<sup>-1</sup> for these heavy metal hydride frequencies, which is excellent, considering the approximations involved. The BPW91 functional is slightly lower and not in as good agreement.

**TIH<sub>2</sub>.** Weak sharp absorptions are produced at 1519.9 (1098.8) and 1390.2 (1007.6) cm<sup>-1</sup> upon 193 nm irradiation of solid hydrogen (deuterium) samples with H/D ratios of 1.383 and 1.380. The solid HD sample revealed sharp 1461.8 and 1046.2 cm<sup>-1</sup> bands, which are 7 cm<sup>-1</sup> above and below the medians, respectively, for the two hydrogen and deuterium modes. This substantiates assignment of the two bands to antisymmetric and symmetric stretching modes of TlH<sub>2</sub> (TlD<sub>2</sub>), where interaction of the Tl–H and Tl–D stretching modes in TlHD forces the two modes apart from the median TlH<sub>2</sub> (TlD<sub>2</sub>) values. The stronger antisymmetric mode is observed at 1525.9 cm<sup>-1</sup> in solid neon and at 1503.3 cm<sup>-1</sup> in solid argon, which bracket the solid H<sub>2</sub> value.

Our B3LYP calculations predict the two TlH<sub>2</sub> modes at 1530 and 1415 cm<sup>-1</sup>, which are 0.6 and 1.8% higher than the hydrogen matrix values. On the other hand, our BPW91 computations find values of 1483 and 1354 cm<sup>-1</sup>, which are 2.4 and 2.6% lower than the observed values. These are the comparisons expected for DFT frequencies.

**TIH<sub>3</sub>.** Weak, sharp bands appear at 1748.4 (1254.4) cm<sup>-1</sup> upon irradiation of solid H<sub>2</sub> (D<sub>2</sub>) samples. With solid HD, these bands shift to 1747.3 and 1258.1 cm<sup>-1</sup>, which are the effects expected for TIH<sub>2</sub>D and TIHD<sub>2</sub>. Solid molecular hydrogens are required for the product, as no counterparts were observed in solid neon or argon.

B3LYP calculations found a trigonal planar TlH<sub>3</sub> species with a 1719 cm<sup>-1</sup> antisymmetric mode and a 1.730 Å bond length. Furthermore, our B3LYP calculations also find TlH<sub>2</sub>D to be 2.1 cm<sup>-1</sup> lower than TlH<sub>3</sub> and TlHD<sub>2</sub> and 3.3 cm<sup>-1</sup> higher than TlD<sub>3</sub>, which is in excellent agreement with the observed -1.1 and +5.4 cm<sup>-1</sup> observed isotopic shifts for these mixed isotopic trihydrides. Relativistic MP2 calculations earlier predicted a 1.728 Å distance and a 1741 cm<sup>-1</sup> frequency,<sup>26</sup> and these higher level calculations are closer to our experimental observation for TlH<sub>3</sub>. On the basis of the neon matrix shift for the antisymmetric mode for TlH<sub>2</sub>, we expect the gas-phase TlH<sub>3</sub> fundamental to be at 1760  $\pm$  10 cm<sup>-1</sup>.

Our TlH<sub>3</sub> band absorbance is much less than what we observed for InH<sub>3</sub>, which in turn is much less than what we observed for GaH<sub>3</sub> and AlH<sub>3</sub>, and is the expected trend.<sup>26</sup> Evaporation of the H<sub>2</sub> (D<sub>2</sub>) matrix gave no evidence of broad absorption bands in the 1500–700 cm<sup>-1</sup> region that might be due to solid thallium hydride. The low yield of TlH<sub>3</sub> and our failure to observe (TlH<sub>3</sub>)<sub>n</sub> at low temperature probably is due to the inherent instability of Tl(III) hydride, which casts doubt on the claimed synthesis of (TlH<sub>3</sub>)<sub>n</sub>.<sup>39,40</sup>

**Tl<sub>2</sub>H<sub>2</sub> and TlTlH<sub>2</sub>.** The large yield of TlH in these experiments invites the consideration of dimers. On the basis of observations for  $Ga_2H_2$  and  $In_2H_2$  and calculations from several groups,<sup>41–45</sup> the rhombic dimer is expected to be the minimum energy structure and several higher energy isomers, including TlTlH<sub>2</sub>, must be considered.

The evidence for  $Tl_2H_2$  is summarized in Figure 5 for neon experiments where the yield is high. The weak band at 919.8 cm<sup>-1</sup> with H<sub>2</sub> increases markedly upon 193 nm irradiation and shifts to 677.3 cm<sup>-1</sup> with D<sub>2</sub> (H/D ratio: 1.378). Experiments with a H<sub>2</sub> + D<sub>2</sub> mixture give primarily TlH, TlD, Tl<sub>2</sub>H<sub>2</sub>, and Tl<sub>2</sub>D<sub>2</sub> without evidence of the Tl<sub>2</sub>HD species. However, an investigation with HD gives a strong intermediate band at 851.1 cm<sup>-1</sup> and very weak Tl<sub>2</sub>H<sub>2</sub> (Tl<sub>2</sub>D<sub>2</sub>) bands at 918.8 (653.1) cm<sup>-1</sup>.

In solid argon, the yield of TlH is high after annealing and a sharp doublet is observed for  $Tl_2H_2$  at 897.0 and 888.2 cm<sup>-1</sup>, which shifts to 648.9 and 642.6 cm<sup>-1</sup> with D<sub>2</sub> (H/D ratios: 1.382 and 1.382). Using higher laser energy with HD (Figure 6e,f) gives a large yield of TlH and TlD and a statistical yield of  $Tl_2H_2$ ,  $Tl_2HD$ , and  $Tl_2D_2$ . However, using lower laser energy with HD (Figure 6d,e) gives a small yield of TlH and TlD and primarily  $Tl_2HD$  at 833.0 and 825.3 cm<sup>-1</sup>.

In solid molecular hydrogen, the weak 908.8 (653.1) cm<sup>-1</sup>  $H_2$  (D<sub>2</sub>) bands are between solid neon and argon values, and pure HD gives an intermediate 839.0 cm<sup>-1</sup> band. Such is also the case for  $In_2H_2$ .<sup>18,41</sup>

Our B3LYP calculations find a very intense  $b_{1u}$  mode for rhombic Tl<sub>2</sub>H<sub>2</sub> at 915 cm<sup>-1</sup>, which is in excellent agreement with the 918.8 cm<sup>-1</sup> neon matrix observation. Stronger interactions with the H<sub>2</sub> and Ar matrixes shift this absorption to lower wavenumbers. The analogous In<sub>2</sub>H<sub>2</sub> species absorbs higher at 982.6 cm<sup>-1</sup> in solid neon. Our B3LYP calculations predict Tl<sub>2</sub>D<sub>2</sub> at 648.8 cm<sup>-1</sup> in the harmonic approximation (H/D ratio: 1.410), and the lower 1.378 neon matrix isotopic frequency ratio is due to anharmonicity in the vibration. Upon symmetry lowering to Tl<sub>2</sub>HD, the normal mode changes and our B3LYP calculation predicts the strongest Tl<sub>2</sub>HD mode at 832.0 cm<sup>-1</sup>, 83/266 of the way from the Tl<sub>2</sub>H<sub>2</sub> to Tl<sub>2</sub>D<sub>2</sub> frequencies. Our observed Tl<sub>2</sub>HD band is 70/256 of the way from the neon matrix Tl<sub>2</sub>H<sub>2</sub> to Tl<sub>2</sub>D<sub>2</sub> values.

Annealing increases sharp new bands at 1430.8 and 1424.3 cm<sup>-1</sup> in H<sub>2</sub> with a weaker 1020.5 cm<sup>-1</sup> D<sub>2</sub> counterpart: the H/D ratio 1.396 is slightly higher than the values for the nearby TlH<sub>2</sub> absorptions. The growth upon annealing invites consideration of another Tl<sub>2</sub>H<sub>2</sub> isomer. Calculations for the Ga and In analogues<sup>17,18,41</sup> show that the  $C_{2\nu}$  MMH<sub>2</sub> isomer is next in energy, but our DFT calculations find the  $C_s$  isomer next with the  $C_{2\nu}$  and  $C_{2h}$  isomer energies within 6 kcal/mol. However, the observed frequencies match best for the  $C_{2\nu}$  TITlH<sub>2</sub> isomer (Table 2). Annealing increases split bands at 1424.2 and 1423.0 cm<sup>-1</sup> and at 1023.5 and 1022.6 cm<sup>-1</sup> in pure HD that show the small sretch–stretch interaction calculated for the TITlH<sub>2</sub> isomer. Our B3LYP calculation predicts close b<sub>2</sub> and a<sub>1</sub> modes, and the 1430.8 and 1424.3 cm<sup>-1</sup> bands are assigned accordingly.

 $Tl^+(H_2)_n$ . The sharp bands at 4113.6 (2962.8) cm<sup>-1</sup> (Figure 4) decrease upon full arc irradiation, never to return. An intermediate 3596.3 cm<sup>-1</sup> counterpart is observed with pure HD. The former absorptions follow a series of like bands with Al (4108.7 and 2959.4 cm<sup>-1</sup>), Ga (4108.9 and 2960.4 cm<sup>-1</sup>), and In (4113.1 and 2961.9 cm<sup>-1</sup>) for the metal cation trapped in the solid molecular hydrogen lattice and the resulting perturbation on the solvating H<sub>2</sub> (D<sub>2</sub>) ligand vibration. The small metal dependence on this species arises from the weak interaction. Our B3LYP calculation shows that the side-bound  $Tl^+(H_2)$ 

		structure:		
species	state	bond length,	rel energy, kcal/mol	frequencies $cm^{-1}$ (sym intensities km/mol)
species	state	A, of aligic, deg	Keal/1101	
	150	<b>THE 2024</b>		B3LYP/6-311++G(d,p)/LanL2DZ
TIH $(C_{\infty v})$	1 <u>&gt;</u> 2 A	TIH: 2.024		THE 1256.4 (1206). The $8/6.7$ (606)
$\Pi H_2(C_{2v})$	$^{-}A_{1}$	11H: 1.929 UTIL: 120.2		$\Pi_{2}$ : 1555.0 (02, 042), 1214.0 ( $a_1$ , 251), 500.4 ( $a_1$ , 95) $\Pi_{2}$ : 062.4 (224), 860.2 (117), 250.5 (48)
$TH_{1}(D_{11})$	14.1	HIIH: 120.2 TH: 1830		$IID_2: 902.4 (324), 800.2 (117), 359.5 (48)$ $TIH_2: 1657.7 (9.70) 1645.6 (9.7498 \times 2), 564.8 (9.77200), 531.8 (9.729 \times 2)$
$\Pi\Pi_3(D_{3h})$	$A_1$	1111. 1.650		11113. 1057.7 ( $a_1$ , 0), 1045.0 ( $e_1$ , 498 × 2), 504.8 ( $a_2$ , 200), 551.8 ( $e_1$ , 229 × 2)
	100			B3LYP/6-311++G(d,p)/SDD
TlH $(C_{\infty v})$	$1\Sigma$	TlH: 1.905		TIH: 1306.0 (1159). TID: 926.1 (583)
$\operatorname{TlH}_2(C_{2\nu})$	$^{2}A_{1}$	TIH: 1.787	0.0	$TH_2$ : 1530.8 (b <sub>2</sub> , 660), 1415.3 (a <sub>1</sub> , 178), 616.3 (a <sub>1</sub> , 133)
$TIII = \langle G \rangle$	1.	HTIH: 121.9	22.2	$IID_2: I086.9 (333), I002.1 (90), 437.7 (67)$
$\Pi H_2 (C_{2v})$	$^{1}A_{1}$	11H: 1.988	-32.3	$IIH_2$ : 1029.0 (a <sub>1</sub> , 1880), 1019.4 (b <sub>2</sub> , 2010), 622.5 (a <sub>1</sub> , 78) TID =, 720.2 (042), 722.0 (1000), 441.2 (27)
$T^{+}(U)$	1.4	HIIH: 91.8	102	$IID_2 : 730.2 (942), 722.9 (1009), 441.3 (57)$
$\Pi^{*}(\Pi_{2})(C_{2v})$	$A_1$	ПП: 5.120 ЦЦ: 0.747	105	$(\Pi_2)$ : 4309.3 ( $a_1$ , 32), 193.4 ( $b_2$ , 17), 193.8 ( $a_1$ , 13) $(\Pi_2)$ : 2000.0 (26), 128.2 (9), 127.7 (7)
$T_{1}H_{2}^{+}(D_{1})$	$1\Sigma +$	TIH: 1.660	150	TH ( $D_2$ ). 5050.5 (20), 150.5 (6), 157.7 (7) TH ( $+$ : 2054.2 ( $\sigma$ 15) 1961.7 ( $\sigma$ 0) 713.7 ( $\pi$ 20 × 2)
$IIII_2 (D_{\infty h})$	∠g	HTIH: 180.0	150	TID $^+$ : 1460 1 (8) 1387 7 (0) 507 3 (16 $\times$ 2)
$T_{1}H_{2}(D_{2k})$	$^{1}A_{1}'$	TIH: 1,730		$TH_{2}^{-1}$ 17661 (6), 15071 (6), 50715 (16 × 2) TH <sub>2</sub> <sup>-1</sup> 1753 5 (a', 0) 1718 8 (e', 445 × 2) 666 6 (a''', 181) 614 1 (e', 210 × 2)
$11113(D_{3n})$	111	HTIH: 120.0		TiD <sub>3</sub> : 1240.4 (0) 1219.5 (227 $\times$ 2) 475.0 (92) 436.2 (106 $\times$ 2)
		111111 12010		$TH_{2}D$ : 1742.2 (142), 1716.7 (463), 1229.6 (136), 612.6 (214), 609.5 (151), 503.0 (139)
				$TIHD_2$ : 1728.7 (306), 1233.0 (84), 1222.8 (206), 559.8 (178), 546.4 (122), 436.5 (105)
$\mathrm{TlH}_4^-(T_d)$	$^{1}A_{1}$	TlH: 1.807		$TIH_4^-$ : 1536.0 (a <sub>1</sub> , 0), 1415.5 (t <sub>2</sub> , 956 × 3), 642.1 (e, 0 × 2), 608.0 (t <sub>2</sub> , 583 × 3)
1 ( 4)				$TID_4^-$ : 1086.6 (0), 1003.4 (483 × 3), 454.2 (0 × 2), 433.3 (288 × 3)
$Tl_2H_2(D_{2h})$	${}^{1}A_{g}$	TlH: 2.157	0.0	Tl <sub>2</sub> H <sub>2</sub> : 982.0 (a <sub>g</sub> , 0), 914.9 (b <sub>1u</sub> , 2435), 707.9 (b <sub>2u</sub> , 491), 664.1 (b <sub>3g</sub> , 0), 303.3 (b <sub>3u</sub> , 11), 95.0 (a <sub>g</sub> , 0)
	8	HT1H: 109.9		Tl <sub>2</sub> HD: 891.7 (114), 832.0 (1714), 552.1 (255), 517.5 (116), 262.5 (8), 95 (0)
				Tl <sub>2</sub> D <sub>2</sub> : 694.8 (0), 648.8 (1224), 502.0 (247), 470.3 (0), 215.1 (60), 95.0 (0)
$Tl_2H_2(C_s)$	$^{1}A'$	TlH: 1.845	12.4	Tl <sub>2</sub> H <sub>2</sub> : 1381.6 (a', 1293), 1003.7 (a', 506), 592.6 (a', 674), 318.6 (a', 5), 196.1 (a'', 5), 50.7 (a', 2)
		TlH': 2.340		
		TITI': 3.285		
		Tl'H': 2.041		
		HT1H': 99.2		
$\mathrm{Tl}_{2}\mathrm{H}_{2}\left(C_{2h}\right)$	$^{1}A_{g}$	TlH: 1.867	17.3	$Tl_2H_2$ : 1358.0 (b <sub>u</sub> , 2193), 1350.7 (a <sub>g</sub> , 0), 333.6 (a <sub>g</sub> , 0), 130.9 (a <sub>u</sub> , 48), 40.7 (a <sub>g</sub> , 0), 35.4 (b <sub>u</sub> , 14)
		TITI: 3.352		
		TITIH:		
$\mathrm{Tl}_{2}\mathrm{H}_{2}\left(C_{2v}\right)$	$^{1}A_{1}$	TlH: 1.793	18.2	$Tl_2H_2$ : 1508.4 (b <sub>2</sub> , 710), 1505.8 (a <sub>1</sub> , 1069), 642.7 (a <sub>1</sub> , 498), 315.1 (b <sub>1</sub> , 81), 160.8 (b <sub>2</sub> , 35), 87.6 (a <sub>1</sub> , 4)
		TITI': 3.129		
	1.	HTIH: 107.9		
$\mathrm{Tl}_{2}\mathrm{H}_{6}\left(D_{2h}\right)$	$^{1}A_{g}$	TIH: 2.012		$T1_2H_6$ : 1833.3 (b <sub>2u</sub> , 653), 1828.6 (a <sub>g</sub> ,0), 1824.9 (b <sub>3u</sub> , 164), 1824.0 (b <sub>1g</sub> , 0),
		TIH: 1.700		1186.5 ( $a_g$ , 0), 10/8.2 ( $b_{3u}$ , 1333), 1001.3 ( $b_{2g}$ , 0), 914.3 ( $b_{1u}$ , 460),
		11H11: 100./		$680.1 (b_{2u}, 196), 648.9 (b_{3g}, 0), 611.1 (a_g, 0), 606.5 (b_{1u}, 109), 503.8 (b_{3u}, 972),$
		H IIH : 137.1		$401.5 (D_{1g}, 0), 508.8 (a_u, 0), 190.5 (D_{2u}, 0), 134.5 (D_{2g}, 0), 107.8 (a_g, 0)$
				BPW91/6-311++G(d,p)/SDD
TlH $(C_{\infty v})$	$1\Sigma$	TIH: 1.923	0.0	TIH: 1278.3 (1039). TID: 906.4 (522)
$\operatorname{TlH}_2(C_{2v})$	$^{2}A_{1}$	TIH: 1.864	0.0	$TH_2$ : 1483.4 (b <sub>2</sub> , 605), 1353.9 (a <sub>1</sub> , 173), 591.3 (a <sub>1</sub> , 110)
	1.	HTIH:		TID <sub>2</sub> : 1053.3 (305), 958.6 (87), 419.9 (55)
$\operatorname{TIH}_2^-(C_{2\nu})$	$^{1}A_{1}$	TIH: 2.006	-31.1	$T1H_2^-$ : 1011.8 (a <sub>1</sub> , 1738), 1008.3 (b <sub>2</sub> , 1839), 594.1 (a <sub>1</sub> , 58)
$T^{+}(\mathbf{I})$	1.4	HTIH: 91.4	102	$IID_2 : /I/.6 (8/0), /I5.0 (920), 421.2 (2/)$
$\Pi^{+}(\mathbf{H}_{2})(\mathbf{C}_{2v})$	$^{1}A_{1}$	IIH: 5.004	103	$(H_2)$ : 4204.8 (a <sub>1</sub> , 85), 232.7 (a <sub>1</sub> , 20), 225.4 (b <sub>2</sub> , 18) T <sup>+</sup> (D <sub>2</sub> ): 2016.8 (41), 165.4 (11), 150.5 (0)
$TIH_{+}^{+}(D_{-})$	15	$\Pi\Pi: 0.731$ $\Pi\Pi: 1.674$	151	TH $(D_2)$ : 5010.8 (41), 105.4 (11), 159.5 (9) TH $(+, 1004.7 (a, 0), 1982.8 (a, 0), 708.5 (a, 18 \times 2))$
$\Pi\Pi_2  (D_{\infty h})$	∠g	HTH: 1.074	151	$1111_2$ . 1994.7 (eu, 9), 1885.8 (eg, 0), 708.5 ( $h_u$ , 18 × 2)
$T_{1}H_{a}(D_{ar})$	14.1	TIH: 1 744		TH.: $1685 1 (a, 0) 1668 A (a' A23 \times 2) 647 1 (a'' 143) 583 6 (a' 181)$
$1111_3(D_{3h})$	$A_1$	$HTH \cdot 120.0$		1113. 1065.1 ( $a_1$ , 0), 1006.4 ( $c$ , 425 × 2), 047.1 ( $a_2$ , 145), 565.0 ( $c$ , 161)
$T H_{4}^{-} (T_{4})$	<sup>1</sup> <b>A</b> <sub>1</sub>	TIH: 1 818		$T_{1}H_{4}^{-1}$ : 1479 6 (a) 1382 4 (to 927 x 3) 617 8 (e 0 x 2) 578 8 (to 504 x 3)
$Tl_2H_2(D_{2k})$	1A.	TIH: 2 162	0.0	The H <sub>2</sub> : $957.1 (a_1, 0)$ , $922.3 (b_{12}, 2003)$ , $739.3 (b_{22}, 0)$ , $715.4 (b_{22}, 424)$ , $291.3 (b_{22}, 11)$ , $94.4 (a_2, 0)$
$112112(D_{2n})$	r ig	TIHTI: 109.5	0.0	$T_{2}T_{2}$ : $5577.2(0), 654.0(1007), 523.6(213), 206.5(5), 94.4(0)$
$Tl_2H_2(C_s)$	$^{1}A'$	TIH: 1.853	12.1	$T_{2}D_{2}^{(i)}$ (0), 00 10 (1007), 000 (010), 2000 (0), 9 (110) $T_{2}D_{2}^{(i)}$ (1358.3 (a', 1103), 952.7 (a', 409), 638.3 (a', 531), 336.9 (a', 0), 205.6 (a'', 5), 66.6 (a', 2)
(-3)		TIH': 2.284		<u>2</u> - <u>2</u>
		TITI': 3.179		
		HTIH': 101.0		
$Tl_2H_2(C_{2h})$	$^{1}A_{g}$	TlH: 1.870	18.3	Tl <sub>2</sub> H <sub>2</sub> : 1333.4 (b <sub>u</sub> , 1861), 1323.3 (a <sub>g</sub> , 0), 363.0 (a <sub>g</sub> , 0), 146.1 (a <sub>u</sub> , 45), 89.7 (b <sub>u</sub> , 18), 56.1 (a <sub>g</sub> , 0)
		TITI: 3.199		
		HTlH: 121.8		
$\mathrm{Tl}_{2}\mathrm{H}_{2}\left(C_{2v}\right)$	$^{1}A_{1}$	TlH: 1.810	18.7	$Tl_{2}H_{2}:\ 1459.1\ (b_{2},\ 650),\ 1448.7\ (a_{1},\ 939),\ 612.2\ (a_{1},\ 396),\ 302.3\ (b_{1},\ 66),\ 150.7\ (b_{2},\ 30),\ 87.8\ (a_{1},\ 3)$
		TITI': 3.122		
		HTIH: 107.2		
$\mathrm{Tl}_{2}\mathrm{H}_{6}\left(D_{2h}\right)$	$^{1}A_{g}$	TlH: 2.026		Tl <sub>2</sub> H <sub>6</sub> : 1787.6 (b <sub>2u</sub> , 598), 1777.1 (b <sub>1g</sub> , 0), 1765.4 (a <sub>g</sub> , 0), 1763.1 (b <sub>3u</sub> , 162), 1143.7 (a <sub>g</sub> , 0),
		TlH': 1.711		1040.8 ( $b_{3u}$ , 1191), 985.3 ( $b_{2g}$ , 0), 909.8 ( $b_{1u}$ , 380), 635.7 ( $b_{2u}$ , 151), 617.9 ( $b_{3g}$ , 0),
		T1HT1: 100.1		586.1 ( $a_g$ , 0), 585.6 ( $b_{1u}$ , 82), 482.7 ( $b_{3u}$ , 809), 391.2 ( $b_{1g}$ , 0), 355.6 ( $a_u$ , 0), 210.0 ( $b_{2g}$ , 0),
		H'TIH': 138.3		$186.2 (b_{2u},0), 105.2 (a_g,0)$

TABLE 3: Group 13 Metal Hydride Stretching Frequencies Observed in Solid Hydrogen<sup>a-c</sup>

	$M^+(H_2)_n$	(H <sub>2</sub> )MH <sub>3</sub>	$M_2H_6$	$MH_3$	$M_2H_5$	$M_2H_4$	$MH_2$	$\rm MMH_2$	$\mathrm{MH_4}^-$	MH	$\mathrm{MH_2}^-$	$M_2H_2 \\$	$(MH_3)_n$
Al	4108.7	4061.6	1932 1915	1884	1918 1845	1838 1826	1822 1788		1638	1599	(1430)	1156	1720
Ga	4108.9	4087.3	1995 1976	1929	1967	1875 1863	1815 1746	1783	1774	1517	1356	1035	1900 1500
In	4113.1	4098.5	1820 1803	1761		1707 1704	1629 1563	1530	1608	1393	(1225)	980	1460
Tl	4113.6			1748			1520 1390	1424		1311		909	

<sup>*a*</sup> References 14–18. <sup>*b*</sup>  $M^+(H_2)_n$  and  $(H_2)MH_3$  are sharp bands for H–H ligand stretching frequencies. The AlH<sub>2</sub><sup>-</sup> value is calculated from the AlD<sub>2</sub><sup>-</sup> value in D<sub>2</sub> times the AlH<sub>2</sub>/AlD<sub>2</sub> frequency ratio. (MH<sub>3</sub>)<sub>*n*</sub> are solid film spectra. <sup>*c*</sup> The values for B<sup>+</sup>(H<sub>2</sub>)<sub>*n*</sub>, 4089.3 cm<sup>-1</sup>, and B<sup>+</sup>(D<sub>2</sub>)<sub>*n*</sub>, 2940.4 cm<sup>-1</sup>, are unpublished results from this laboratory.

structure is more stable than the linear (HTIH)<sup>+</sup> structure, but here we have a more highly coordinated Tl<sup>+</sup> species that is displaced 39.2 cm<sup>-1</sup> below the 4152.8 cm<sup>-1</sup> frequency for the solid n-H<sub>2</sub> lattice.<sup>46</sup> Finally, charge balance is achieved in these experiments by trapping H<sup>-</sup> (D<sup>-</sup>) in solid H<sub>2</sub> (D<sub>2</sub>), as evidenced by strong 3972 (2870) cm<sup>-1</sup> absorptions.<sup>47</sup>

**Emission Spectra.** Two properties of the 193 nm induced Tl emission spectra in solid H<sub>2</sub> and D<sub>2</sub> are interesting. In H<sub>2</sub>, the two  ${}^{2}S_{1/2} \rightarrow {}^{2}P_{1/2,3/2}$  components are blue-shifted 210 and  $320 \pm 20 \text{ cm}^{-1}$ , and in solid D<sub>2</sub>, these bands are blue-shifted 660 and 780  $\pm 20 \text{ cm}^{-1}$ . Not only has D<sub>2</sub> blue-shifted the emissions ~450 cm<sup>-1</sup> more than H<sub>2</sub>, but also the spin—orbit splitting (7793 cm<sup>-1</sup> in the gas phase) has decreased by  $120 \pm 10 \text{ cm}^{-1}$  in solid D<sub>2</sub> and by  $110 \pm 10 \text{ cm}^{-1}$  in solid H<sub>2</sub>. The smaller lattice parameter (3.600 Å) for solid n-D<sub>2</sub> suggests a more repulsive cage interaction than that for solid n-H<sub>2</sub> (3.769 Å).<sup>48,49</sup> Spin—orbit splitting is often diminished in the matrix,<sup>50</sup> and these small decreases (1.4–1.9%) are reasonable. Finally, the emission yield for Tl was much higher than that found for In and Ga in solid hydrogen, which attests the lower reactivity of Tl with H<sub>2</sub>.

**Reaction Mechanisms.** The reactions of Tl and H<sub>2</sub> observed here are common to those for the other group 13 metals. The initial reaction (eq 1) is endothermic ( $\Delta E = 58 \text{ kcal/mol})^1$  and must be driven by exciting the metal atom. The insertion reaction (eq 2) also requires activation energy. Note that  $\lambda > 240$  nm irradiation (Figure 1b) markedly increases the yield of both TlH and TlH<sub>2</sub> and produces TlH<sub>3</sub> from reactions 3 and 4.

$$TI + H_2 \rightarrow TIH + H \tag{1}$$

$$Tl + H_2 \rightarrow TlH_2$$
 (2)

$$TIH + H_2 \rightarrow TIH_3 \tag{3}$$

$$\Gamma IH_2 + H \to T IH_3 \tag{4}$$

Unreacted Tl clusters form Tl<sub>2</sub>, and this reaction is favored in the soft neon matrix. Then irradiation leads directly to Tl<sub>2</sub>H<sub>2</sub> formation (reaction 5). Such is the mechanism proposed for In<sub>2</sub>H<sub>2</sub>, but the reactions with Al<sub>2</sub> and Ga<sub>2</sub> are spontaneous. Note the experiments with HD in neon gave major Tl<sub>2</sub>HD and those with H<sub>2</sub> + D<sub>2</sub> in neon favored Tl<sub>2</sub>H<sub>2</sub> and Tl<sub>2</sub>D<sub>2</sub> (Figure 5), which confirms the reduction in reaction 5. Very similar results are observed in solid argon (Figure 6). In the more rigid argon matrix at 3.5 K, unreacted Tl is isolated and H atoms from dissociation by laser plume radiation react upon annealing to form TlH, which also can dimerize (reactions 6 and 7). With higher laser energy, the yields of TlH and TlD from HD are large and the  $Tl_2H_2$ ,  $Tl_2HD$ , and  $Tl_2D_2$  yields from reaction 7 are statistical (Figure 6).

T TT 7

$$Tl_2 + H_2 \xrightarrow{OV} Tl_2H_2$$
 (5)

$$\Gamma l + H \rightarrow T l H$$
 (6)

$$2\text{TlH} \rightarrow \text{Tl}_2\text{H}_2 \tag{7}$$

Annealing in solid hydrogen fosters the further reaction of Tl with TlH<sub>2</sub>, as reactions 1-3 require activation energy.

$$TI + TIH_2 \rightarrow TITIH_2 \tag{8}$$

The laser-ablation process produces metal cations as well as neutral atoms,<sup>51</sup> as seen from the emission spectrum (Figure 7), and the  $Tl^+$  cation complex is formed upon trapping of  $Tl^+$  in solid H<sub>2</sub>.

$$\mathrm{Tl}^{+} + n\mathrm{H}_{2} \rightarrow \mathrm{Tl}^{+}(\mathrm{H}_{2})_{n} \tag{9}$$

### Conclusions

Laser-ablated Tl atoms react with  $H_2$  much the same as In, Ga, and Al, but major differences do exist. Thallium is less reactive: The lower product band and higher 193 nm atomic emission yield attest this fact. The dominant thallium product is Tl(I)H, whereas the major aluminum product is Al(III)H<sub>3</sub>. This substantiates the observation of Al<sub>2</sub>H<sub>6</sub> and solid (AlH<sub>3</sub>)<sub>n</sub> and the absence of Tl<sub>2</sub>H<sub>6</sub> and solid (TlH<sub>3</sub>)<sub>n</sub> in the product spectrum. Although the yield of TlH<sub>3</sub> is small, its identification with TlH<sub>2</sub>D, TlHD<sub>2</sub>, and TlD<sub>3</sub> isotopic modifications is definitive. Table 3 summarizes the group 13 metal hydride stretching frequencies: The TlH<sub>3</sub> fundamental is only 13 cm<sup>-1</sup> below the InH<sub>3</sub> value, whereas the TlH<sub>2</sub> and TlH modes are 173–82 cm<sup>-1</sup> below their indium counterparts.

The reaction of metal dimer  $M_2$  with  $H_2$  gives the rhombic  $M_2H_2$  dimer, although the reactions are spontaneous for  $Al_2$  and  $Ga_2$  but they require photochemical activation for  $In_2$  and  $Tl_2$ . A significant yield of the TITIH<sub>2</sub> isomer is also observed. Sharp  ${}^2S_{1/2} \rightarrow {}^2P_{1/2,3/2}$  emission from unreacted atomic Tl induced by 193 nm radiation is blue-shifted 660 cm<sup>-1</sup> in solid  $D_2$  and 210 cm<sup>-1</sup> in solid  $H_2$ .

Although the DFT calculated electron affinity for  $TIH_2$  is 32 kcal/mol, we have no definitive evidence for the  $TIH_2^-$  anion in the spectrum. This may arise from detachment by irradiation in the experiment. On the other hand,  $AIH_4^-$  is the dominant anion observed with Al.

The group 13 metal cations all form weak complexes with  $H_2$  in the solid lattice with a H–H stretching mode 39–44 cm<sup>-1</sup> below the solid  $H_2$  fundamental at 4153 cm<sup>-1</sup>.

Density functional theory using the relativistic SDD pseudopotential is adequate to calculate frequencies for thallium hydrides, but these calculations fit better for TIH > TIH<sub>2</sub> > TIH<sub>3</sub>. However, relativistic MP2 calculations produce more accurate frequencies for TIH<sub>3</sub>. It must be noted that the LANL2DZ pseudopotential and basis is vastly inferior for thallium hydrides.

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#### **References and Notes**

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