Photoelectron spectroscopy of phosphorus hydride anions

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Negative-ion photoelectron spectroscopy is applied to the PH−, PH2−, P3H−, P2H2−, and P3H3− molecular anions. Franck–Condon simulations of the photoelectron spectra are used to analyze the spectra and to identify various P2Hn− species. The simulations employ density-functional-theory calculations of molecular geometries and vibrational frequencies and normal modes, and coupled-cluster theory calculations of electron affinities. The following electron affinities are obtained: EAn(PH)=1.027±0.006 eV, EAn(PH2)=1.263±0.006 eV, and EAn(P3H)=1.514±0.01 eV. A band is identified as a mixture of trans-HPPH− and cis-HPPH−. Although the trans and cis bands cannot be definitively assigned from experimental information, using theory as a guide we obtain EAn(trans-HPPH)=1.00±0.01 eV and EAn(cis-HPPH)=1.03±0.01 eV. A weak feature tentatively assigned to P2H3− has a vertical detachment energy of 1.74 eV. The derived gas-phase acidity of phosphine is ΔacidG298(PH3)≈1509.7±2.1 kJ mol−1.

I. INTRODUCTION

Phosphorus hydride compounds and radicals are of interest in planetary atmospheres and other astrophysical environments.1,2 They are also important because of the role of organophosphorus compounds in catalysis of radical recombination reactions involved in combustion3,4 and because of the use of phosphine in chemical-vapor deposition of phosphide semiconductors.5–7 Fundamentally, accurate descriptions of multiply bonded compounds of third-row elements by theoretical methods4,8–15 is a challenging problem. Whereas neutral phosphines have been studied extensively,16 their radical and anionic fragments are less well characterized. In this work, we apply negative-ion photoelectron spectroscopy to a series of phosphorus hydride anions, PH−, PH2−, P3H−, P2H2−, and P3H3−. We identify the transitions between electronic and vibrational levels of the anions and the corresponding neutrals and we report electron affinities and other molecular constants. The photoelectron spectra of PH− and PH2− have been reported previously17–19 but are obtained here with higher resolution allowing rotational contours to be examined and more precise electron affinities to be obtained. The photoelectron spectra of the diphosphorus hydride anions, P2Hn− (n=1−3), are reported for the first time.

II. METHODS

Negative-ion photoelectron spectroscopy is performed by crossing a mass-selected ion beam with a fixed-frequency laser. The kinetic energies of photodetached electrons are measured with an electrostatic energy analyzer to obtain the photoelectron spectrum of transitions between initial states of the anion and final states of the corresponding neutral species. Electron binding energies (eBEs) of observed transitions are given by the photon energy minus the measured electron kinetic energy (eKE). The experimental techniques and data analysis procedures have been described in detail in previous reports.20–23

Phosphorus hydride anions are produced by injecting phosphine (PH3) into a flowing afterglow ion source downstream of a microwave discharge, using helium buffer gas at a total pressure of 0.3–0.5 Torr. Phosphine is a highly toxic gas that requires special handling procedures. The PH3 lecture bottle, regulator, and gas handling system were enclosed in a vented cabinet. The major anion formed is PH2− (typically 60 pA), with smaller amounts of PH− (5 pA) and P− (8 pA). Diphosphorus hydride species (P2H2−, 10–20 pA) are presumably produced by secondary-ion–molecule reactions of PH− species with PH3. Ions of a particular mass are selected by passage through a Wien velocity filter. The mass resolution is sufficient to separate PHn− ions completely, but partial mass overlap occurs among various P2Hn− species.

This work employs the 351.1-nm (3.531 eV) or 363.8-nm (3.408 eV) lines of an argon-ion laser, amplified to 30–40 W in a power build-up cavity, to induce photodetachment.22 The resolution of the hemispherical electrostatic energy analyzer is enhanced by imaging the output photoelectrons onto a position-sensitive anode detector.20 The resolution of the kinetic-energy analyzer is 5–8 meV over the entire energy range. The electron kinetic-energy scale is calibrated from the photoelectron spectrum of atomic phosphorus anions, described in detail below. The experimental uncertainty in the absolute electron kinetic energy of
TABLE I. Electron affinities (eV).

<table>
<thead>
<tr>
<th>Species</th>
<th>Theorya</th>
<th>Expt.</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>P</td>
<td>0.613 (0.734)</td>
<td>0.7465±0.0003</td>
<td>33 and 34</td>
</tr>
<tr>
<td>PH</td>
<td>0.922 (1.026)</td>
<td>1.027±0.006</td>
<td>This work</td>
</tr>
<tr>
<td>PH₂</td>
<td>1.201 (1.271)</td>
<td>1.263±0.006</td>
<td>This work</td>
</tr>
<tr>
<td>P₂</td>
<td>0.562</td>
<td>0.589±0.025</td>
<td>56</td>
</tr>
<tr>
<td>P₂H</td>
<td>1.462</td>
<td>1.514±0.010</td>
<td>This work</td>
</tr>
<tr>
<td>trans-HPHHb</td>
<td>1.008</td>
<td>1.00±0.01</td>
<td>This work</td>
</tr>
<tr>
<td>cis-HPHHb</td>
<td>1.043</td>
<td>1.03±0.01</td>
<td>This work</td>
</tr>
<tr>
<td>H₂PP (pyramidal)</td>
<td>1.426</td>
<td>1.028±0.010</td>
<td>Ref. 14</td>
</tr>
<tr>
<td>P₂H₃</td>
<td>1.557</td>
<td>1.74 eV (VDE)c</td>
<td>This work</td>
</tr>
</tbody>
</table>

aCCSD(T)/FC/aug-cc-pVTZ/B3LYP/aug-cc-pVTZ with zero-point energy corrections. Values in parenthesis are higher-level calculations from Ricca and Bauschlicher (Ref. 14). 
bTentative assignment of isomers, as discussed in text.  
cVertical detachment energy.

well-resolved peaks is ±0.005 eV (two combined standard uncertainties). Atomic transition intensities and Franck–Condon factors for fitting molecular spectra are calculated using the PESCAL program described previously.

Molecular structure and energy calculations are performed using GAUSSIAN 03. Geometries and vibrational frequencies of the anions and neutral were calculated using density-functional theory (DFT) with Becke’s three-parameter hybrid functional using the Lee–Yang–Parr correlation (B3LYP) and with augmented correlation-consistent polarized valence triple-zeta (aug-cc-pVTZ) or quadruple-zeta (aug-cc-pVQZ) basis sets. Electron affinities are calculated using coupled-cluster theory at the coupled-cluster with single and double excitations and perturbative inclusion of triple excitations (CCSD(T)) level in single- and double-excitation perturbation theory of the anions and neutral were calculated using coupled-cluster theory (CCSD) at the coupled-inclusion of triple excitations cluster with single and double excitations and perturbative density-functional theory parameters.

III. RESULTS AND DISCUSSION

A. Atomic phosphorus, P

The 363.8-nm photoelectron spectrum of P⁻ is displayed in Fig. 1. The electron affinity of phosphorus atom is precisely known from previous high-resolution photodetachment threshold measurements, as are the energy levels of the P levels, namely, P(3P) (0 cm⁻¹), P(3P₁) (181±2 cm⁻¹), and P(3P₀) (263±2 cm⁻¹). The photoelectron spectrum exhibits transitions from these anion levels to the P(3S₂) (0 cm⁻¹) and P(3P₁) (181±2 cm⁻¹) and P(3P₀) (263±2 cm⁻¹). The largest peaks in the two multiplets are due to the P(3P₁) origin transition and the P(3P₀)→P(3P₁) transition, respectively. The P(3P₁) levels of P⁻ are clearly resolved in the spectrum, but the P(3P₀) levels of P⁻ are split by only 15.6 cm⁻¹ (1.9 meV) and are not resolved.

Table II lists the calculated energies and relative intensities of the fine-structure transitions. The relative intensities are calculated in the L-S coupling approximation for one-electron detachment of a 3p electron using statistical factors and Clebsch–Gordan coefficients as described by Engleking and Lineberger. The spectrum is fit by nonlinear least-

FIG. 1. Photoelectron spectrum of P⁻ at a laser wavelength of 363.8 nm (hv=3.408 eV). Photodetachment intensities (solid circles) are plotted as a function of the electron binding energy and electron kinetic energy (eBE=hf−eKE). The bars and curves show fits using calculated relative intensities with anion temperature of 390 K, as described in the text and listed in Table II. The inset shows the instrumental resolution function.
TABLE II. Fine-structure transitions for \( P^- + h\nu \rightarrow P + e^- \).

<table>
<thead>
<tr>
<th>Transition (P( \rightarrow )P( ^- ))</th>
<th>Energy (cm(^{-1}))</th>
<th>eBE (eV)</th>
<th>Calc. (( T=\infty ))</th>
<th>Calc. (( T=390 ) K)</th>
<th>Expt.</th>
</tr>
</thead>
<tbody>
<tr>
<td>( ^4S_{3/2} \rightarrow ^3P_0 )</td>
<td>( -263 \pm 2^{b} )</td>
<td>0.7139</td>
<td>0.111</td>
<td>0.055</td>
<td>0.06</td>
</tr>
<tr>
<td>( ^4S_{3/2} \rightarrow ^3P_1 )</td>
<td>( -181 \pm 2^{b} )</td>
<td>0.7241</td>
<td>0.333</td>
<td>0.222</td>
<td>0.22</td>
</tr>
<tr>
<td>( ^4S_{3/2} \rightarrow ^3P_2 )</td>
<td>0</td>
<td>0.7465(^b)</td>
<td>0.556</td>
<td>0.723</td>
<td>0.72</td>
</tr>
<tr>
<td>( ^2D_{3/2} \rightarrow ^3P_0 )</td>
<td>( 11098 \pm 2^{c} )</td>
<td>2.1225</td>
<td>0.111</td>
<td>0.055</td>
<td>0.05</td>
</tr>
<tr>
<td>( ^2D_{3/2} \rightarrow ^3P_1 )</td>
<td>( 11114 \pm 2^{c} )</td>
<td>2.1245</td>
<td>0.000</td>
<td>0.000</td>
<td>⋯</td>
</tr>
<tr>
<td>( ^2D_{3/2} \rightarrow ^3P_2 )</td>
<td>( 11180 \pm 2^{c} )</td>
<td>2.1326</td>
<td>0.233</td>
<td>0.156</td>
<td>0.14</td>
</tr>
<tr>
<td>( ^2D_{5/2} \rightarrow ^3P_1 )</td>
<td>( 11196 \pm 2^{c} )</td>
<td>2.1346</td>
<td>0.100</td>
<td>0.067</td>
<td>0.08</td>
</tr>
<tr>
<td>( ^2D_{5/2} \rightarrow ^3P_2 )</td>
<td>( 11361 \pm 2^{c} )</td>
<td>2.1551</td>
<td>0.056</td>
<td>0.072</td>
<td>0.12</td>
</tr>
<tr>
<td>( ^2D_{5/2} \rightarrow ^3P_0 )</td>
<td>( 11377 \pm 2^{c} )</td>
<td>2.1571</td>
<td>0.500</td>
<td>0.650</td>
<td>0.61</td>
</tr>
</tbody>
</table>

This work.
\(^b\)Slater and Lineberger (Ref. 33), EA\( (P) = 0.7465 \pm 0.0003 \) eV.
\(^c\)Martin et al. (Ref. 35).

The photoelectron spectrum of \( P^- \) is shown in Fig. 2. We observe transitions from the \( X^2 \Pi_j \) ground state of \( P^- \) to the \( X^3 \Sigma^- \), \( a^1 \Delta \), and \( b^1 \Sigma^+ \) states of \( P \). Each of these transitions is completely vertical—the \( 0 \rightarrow 0 \) origins are observed, but no \( 1 \rightarrow 0 \) or higher vibrational transitions or \( 0 \rightarrow 1 \) hot bands. For all three electronic transitions, the electron is detached from one of the two nonbonding \( p \)-type orbitals centered on phosphorus. The three neutral electronic states represent the different ways of coupling the two electrons in these two orbitals. The absence of the \( 1 \rightarrow 0 \) fundamental transitions places an upper limit of 0.02 for the Franck-Condon factors for each transition, limited by the base line noise level. Limits on the bond length of \( P^- \) may be estimated from this experimental upper limit for the Franck-Condon factors. For this purpose we calculate Franck-Condon factors using the Morse oscillator approximation.\(^{25,37}\) The spectroscopic bond lengths and vibrational constants for the neutral electronic states are listed in Table III.\(^{38-46}\) For the anion vibrational constants, we use the theoretical values of Rosmus and Meyer\(^{45}\) in Table III. These values agree with the less-precise experimental fundamental frequency of Zittel and Lineberger,\(^{18}\) who were able to observe the \( 0 \rightarrow 1 \) hot band in the photoelectron spectrum because of a higher anion temperature than is available with the present ion source. The Morse oscillator calculations show that a maximum Franck-Condon intensity of 0.02 corresponds to a maximum change in the bond length of 0.024 Å for the anion relative to any of the three observed electronic states of \( P \). Zittel and Lineberger\(^{18}\) assumed that the bond length of \( P^- \) is shorter than that of \( P \) on the basis of comparisons with isoelectronic molecules, but \( ab \) initio calculations indicate that the anion is slightly longer. Rosmus and Meyer\(^{45}\) and Meyer and Rosmus\(^{47}\) find from coupled electron pair approach (CEPA) calculations that the ground-
state bond length is 0.015 Å longer in the anion, while our density-functional theory calculations at the B3LYP/aug-cc-pVQZ level give 0.011 Å. These values are completely consistent with our experimental upper limit of 0.024 Å. Taking the shortest of the neutral states,\(^\text{40}\) \(r_e = 1.418\) Å for \(\text{PH}^-\ 1\Sigma^+\), our maximum bond-length change yields an upper limit of \(r_e < 1.442\) Å for \(\text{PH}^-\ X\ 2\Pi\)). The minimum change of 0.008 Å found by Zittel and Lineberger\(^\text{39}\) from the intensity of the \(0 \rightarrow 1\) hot band of the \(\text{PH}^+\ X\ 3\Sigma^+ \rightarrow \text{PH}^-\ X\ 2\Pi\) transition gives a lower limit of \(r_e > 1.430\) Å. Thus, the bond length of the anion is fairly tightly established as \(1.430 \leq r_e \leq 1.442\) Å, in excellent agreement with the CEPA theoretical value\(^\text{45}\) of 1.441 Å and our calculation of 1.439 Å at the B3LYP/aug-cc-pVQZ level.

Figure 3 presents each electronic transition in the photoelectron spectrum of \(\text{PH}^-\) on an expanded scale, showing that these transitions are structured. The structure arises from the spin–orbit splitting of the \(^1\Pi_3/2\) and \(^3\Pi_1/2\) states of \(\text{PH}^-\) and from the contours of partially resolved rotational branches, which were not resolved in the previous photoelectron spectra.\(^\text{18}\) To simulate the contours, line positions and line strengths are calculated for \([\text{PH}(1\Sigma^+) + e^-] \rightarrow \text{PH}^- (^3\Pi_1)\) and \([\text{PH}(1\Delta) + e^-] \rightarrow \text{PH}^- (^3\Pi_1)\) absorption transitions using the DIATOMIC program\(^\text{48}\) with the rotational and spin–orbit coupling constants in Table III, and then convolved over the instrumental resolution function obtained from the atomic P\(^+\) spectrum. The spin–orbit constant for the anion and the anion temperature are obtained by manually adjusting the values in DIATOMIC until a reasonable match to the observed contours is obtained. We obtain good fits with \(T = 300\) K and \(A(\tilde{\Pi}_1) = -185 \pm 10\) cm\(^{-1}\). As shown in Fig. 3, very good but not perfect agreement with the observed contours is obtained. The small deviations may arise from the approximation of treating the photodetachment process as an absorption transition (i.e., neglecting the detailed interaction of the outgoing electron with the states of the neutral) or the approximation that the rotational and spin–orbit temperatures of the anion are identical. Modeling with values of the spin–orbit splitting outside this \(-185\pm 10\) cm\(^{-1}\) range results in visually degraded fits to the main branches. To our knowledge, this result is the first experimental measurement of the spin–orbit splitting in \(\text{PH}^-\). It is in excellent agreement with the theoretical value of \(-185.7\) cm\(^{-1}\) calculated by relativistic self-consistent-field methods.\(^\text{39}\) The rotationless origins for each electronic transition are marked by vertical bars in Fig. 3. The energies of these origin transitions give the adiabatic

<table>
<thead>
<tr>
<th>Value</th>
<th>(\text{PH}^-\ X^2\Pi)</th>
<th>(\text{PH}^+\ X^2\Sigma^+)</th>
<th>(\text{PH}^+\ 1\Delta)</th>
<th>(\text{PH}^+\ 1\Sigma^+)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(T_0)</td>
<td>(-1.027\pm 0.006) eV</td>
<td>0</td>
<td>(7558\pm 15) cm(^{-1})</td>
<td>(14310\pm 30) cm(^{-1})</td>
</tr>
<tr>
<td>(r_e) (Å)</td>
<td>(1.430 &lt; r_e &lt; 1.442)</td>
<td>(1.422)</td>
<td>(1.420)</td>
<td>(1.418)</td>
</tr>
<tr>
<td>(r_0) (Å)</td>
<td>(1.430)</td>
<td>(1.430)</td>
<td>(1.428)</td>
<td>(1.428)</td>
</tr>
<tr>
<td>(\omega_e) (cm(^{-1}))</td>
<td>(224)</td>
<td>(2363.8)</td>
<td>(2415)</td>
<td>(2403.0)</td>
</tr>
<tr>
<td>(\omega_r) (cm(^{-1}))</td>
<td>(50)</td>
<td>(43.9)</td>
<td>(48)</td>
<td>(42.0)</td>
</tr>
<tr>
<td>(A) (cm(^{-1}))</td>
<td>(-185\pm 10)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\(^\text{a}\)This work.
\(^\text{b}\)Reference 38.
\(^\text{c}\)Reference 39.
\(^\text{d}\)Estimated from experimental \(r_e\) by interpolation between other states.
\(^\text{e}\)Theoretical values used for Franck–Condon calculations (Ref. 45).
\(^\text{f}\)Theoretical values used for Franck–Condon calculations (Ref. 46).
electron binding energy for each electronic state. The resulting value for the adiabatic electron affinity is $E_A(\text{PH}) = 1.027 \pm 0.006 \text{ eV}$, which is in complete agreement with the previous experimental value of 1.028 \pm 0.010 \text{ eV}.^{18}$ The measured electronic term energies are $T_0(\text{PH} a^1 \Delta) = 7555.1 \text{ cm}^{-1}$ and $T_0(\text{PH} b^1 \Sigma^+ ) = 14345 \text{ cm}^{-1}$. These energies agree with literature values of $T_0(\text{PH} a^1 \Delta) = 14345 \text{ cm}^{-1}$ and $T_0(\text{PH} b^1 \Sigma^+ ) = 7555.1 \text{ cm}^{-1}$ from electronic spectroscopy (Table III).^{38,40}

### C. Phosphino radical, PH$_2^-$

Figure 4(a) shows the vibrational transitions of the PH$_2(X^2 B_1) \rightarrow PH_2(X^1 A_1)$ band in the photoelectron spectrum of PH$_2^-$- No other transitions are observed in the energy range of the experiment. The spectrum is nearly vertical, dominated by the 0–0 origin transition, consistent with detachment of the electron from a nonbonding phosphorus p-type orbital. A weak 1–0 transition of the $v_9$ symmetric stretch vibration is clearly observed and a slight rise above the background is observed at the expected position of the 1–0 transition of the $v_9$ bending vibration. The asymmetric stretch mode is not active by symmetry selection rules, and the relatively high anion frequencies and small Franck-Condon factors suppress the hot bands. The origin transition and symmetric stretch fundamental transition exhibit partially resolved rotational contours with a central Q branch peak and adjacent P and R branches, as shown in the expanded plot of the origin transition in Fig. 4(b).

To model the rotational contour, the photodetachment transition is treated as a two-step process,^{50}$\text{PH}_2(X^2 A_1) \rightarrow \text{PH}_2^-(1B_1) \rightarrow \text{PH}_2(X^2 B_1) + e^-(k$s). The virtual intermediate anion state represents the PH$_2^-$ anion with the extra electron in a continuum $s$ orbital. As described in a previous work on the isovalent anion NH$_2^-$,^{50}$ the rotational energy levels and transition intensities are calculated for an asymmetric rotor using type-C selection rules. Rotational constants are known for the neutral and are presented in Table IV.$^{1,51-53}$ For the anion we used calculated geometries for initial fits, but the contours shown in Fig. 4 use the rotational constants in Table IV obtained from our Franck-Condon analysis described below. A temperature of 300 K was used for Boltzmann rotational energy-level populations and the calculated lines are broadened by the instrumental resolution function obtained from the atomic phosphorus spectrum. This simulated contour agrees well with the experiment, as shown in Fig. 4(b). The binding energy of the rotationless origin marked in Fig. 4(b) represents the measured adiabatic electron affinity, $E_A(\text{PH}_2) = 1.263 \pm 0.006 \text{ eV}$, in agreement with the previous photoelectron spectroscopy value^{18} of 1.271 \pm 0.010 eV.

The position of the 1–0 transition of the symmetric stretch gives a vibrational frequency for neutral PH$_2$ of 2925 \pm 15 cm$^{-1}$, in reasonable agreement with 2310 \pm 2 cm$^{-1}$ from a high-temperature Raman spectrum^{54} (obtained by

### Table IV. Molecular constants for PH$_2$ and PH$_2^-$

<table>
<thead>
<tr>
<th>Constant</th>
<th>PH$_2$</th>
<th>PH$_2^-$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A$ (cm$^{-1}$)</td>
<td>8.96</td>
<td>9.113$^b$</td>
</tr>
<tr>
<td>$B$ (cm$^{-1}$)</td>
<td>7.76</td>
<td>8.083$^b$</td>
</tr>
<tr>
<td>$C$ (cm$^{-1}$)</td>
<td>4.16</td>
<td>4.214$^d$</td>
</tr>
<tr>
<td>$r$ (Å)</td>
<td>$r' + (0.016 \pm 0.004) = 1.439$</td>
<td>1.423$^b$</td>
</tr>
<tr>
<td>$\theta$ (°)</td>
<td>$\theta' + (0.6 \pm 0.2) = 92.3$</td>
<td>91.7$^b$</td>
</tr>
<tr>
<td>$v_1$ (cm$^{-1}$)</td>
<td>1060$^f$</td>
<td>1101.4$^c$</td>
</tr>
<tr>
<td>$v_2$ (cm$^{-1}$)</td>
<td>2221$^d$</td>
<td>2295 \pm 15$^a$</td>
</tr>
<tr>
<td>$v_3$ (cm$^{-1}$)</td>
<td>2225$^d$</td>
<td>2310 \pm 2$^e$</td>
</tr>
<tr>
<td>$E_A$ (eV)</td>
<td>1.263 \pm 0.006$^d$</td>
<td></td>
</tr>
</tbody>
</table>

$^a$This work except as noted. Rotational constants are calculated from the estimated geometry neglecting the inertial defect.

$^b$Reference 51.

$^c$References 53 and 55.

$^d$Frequencies calculated at the B3LYP/aug-cc-pVTZ level with a scaling factor of 0.98.

$^e$References 54 and 55.
subtracting the PH$_3$ signal from the spectrum of PH$_2$ and PH$_3$). The bending vibration is too weak for a precise determination of its frequency; we use the spectroscopic value\textsuperscript{53,55} of 1101.4 cm$^{-1}$. By comparison with these experimental frequencies, an empirical scaling factor of 0.98 is determined for frequencies calculated by the B3LYP/aug-cc-pVTZ method. These scaled frequencies for the anion and the inactive symmetric stretch are listed in Table IV. To model the intensities, Franck–Condon factors are calculated in the harmonic-oscillator approximation using methods described previously.\textsuperscript{26} Duschinsky rotation is included using normal coordinate vectors from the B3LYP/aug-cc-pVTZ frequency calculations, but the Duschinsky rotation effect is minor for this system. The geometry displacements between the known values for the neutral (Table IV)\textsuperscript{51} and the anion are found by fitting the Franck–Condon intensities. The Franck–Condon fit can establish the magnitude but not the sign of the changes in the bond length and angle. Calculations show that both increase from neutral to the anion. Using the calculated geometries to assign the direction of the change, we find that the bond length in the anion is 0.016±0.004 Å longer than in the neutral and that the bond angle is 0.6°±0.2° wider. Using geometry values for the neutral from Chen \textit{et al.},\textsuperscript{51} we can estimate geometries and rotational constants for PH$_2^-$, as listed in Table IV.

D. Identification of P$_2$H$_n^-$ species

The P$_2$H$_n^-$ species are incompletely resolved in the mass spectrum. Survey photoelectron spectra, taken by adjusting the setting of the Wien filter to the low- or high-mass side of the P$_2$H$_n^-$ peak, are compared in Figs. 5(a) and 5(b). These spectra show that at least two different molecular anions are present. The major species in Fig. 5(a) has an origin peak at an electron binding energy of about 1.52 eV and is on the low-mass side of the peak. Three additional transitions arise when the mass setting is moved to the higher-mass side of the peak, as shown in Fig. 5(b). To identify the major transitions in Figs. 5(a) and 5(b), we compare them with simulated spectra for P$_2^-$, P$_2$H$^-$, P$_2$H$_2^-$, and P$_2$H$_3^-$ in Figs. 5(c)–5(h). The simulated spectra use B3LYP/aug-cc-pVTZ geometries, frequencies, and normal-mode vectors, with electron affinities calculated at the CCSD(T)/aug-cc-pVTZ//B3LYP/aug-cc-pVTZ level. The Franck–Condon profiles are calculated in the harmonic-oscillator approximation with full treatment of Duschinsky rotation.\textsuperscript{25,26}

Several low-lying electronic states and isomers of P$_2$H$_2^-$ and P$_2$H$_3^-$ exist, as previously studied by Schaefer and co-workers.\textsuperscript{9–11} Our theoretical calculations generally agree with those results; here we use the CCSD(T)/aug-cc-pVTZ//B3LYP/aug-cc-pVTZ relative energies as shown in Fig. 6 for consistent comparisons. The Franck–Condon simulations for trans-HPPH$^-$ and cis-HPPH$^-$ are shown in Figs. 5(e) and 5(f), respectively. The trans isomer is the ground doublet state of the anion and the neutral singlet ground state, but the cis isomer is only 0.10 eV higher for the anion and 0.14 eV higher for the neutral according to the calculations. The stable geometry of the triplet excited state of neutral HPPH is nonplanar (skewed geometry), with a dihedral angle of

![Figure 5](image-url)
The cis and trans isomers on the neutral triplet surface represent transition state geometries for the rotation around the PP bond of the skewed, nonplanar HPPH. The geometry difference between the anion (either trans or cis) and the nonplanar neutral triplet is thus quite large and the Franck–Condon factors in the harmonic approximation are not meaningful. The simulations in Fig. 5 for cis-HPPH− and trans-HPPH− instead show the vertical transitions to the cis-HPPH and trans-HPPH triplet transition states, respectively, obtained by simply ignoring the mode with imaginary frequency in the upper state. That is an oversimplification; these simulations are intended merely to show the regions where the transition to the neutral triplet surface might occur. The transition states marked with a superscript dagger, which have one imaginary frequency, are thus saddle points on the neutral doublet surface. The triplet anion has the same asymmetric geometry as the neutral, but is unbound with respect to electron detachment. Because the anion has poor Franck–Condon overlap with the neutral, the harmonic approximation for Franck–Condon factors is not useful. In Fig. 5(h), we instead plot the simulated spectrum for vertical transitions from the C₆ anions to the C₆ saddle points on the neutral surface, with two anion conformations weighted for a 300-K population. The actual spectrum to the asymmetric neutral would be expected to be in the same energy region but with greater broadening because of the neglected displacement in the H₂P–PH internal rotation coordinate.

We now compare the simulated spectra in Figs. 5(c)–5(h) to the experimental spectra in Figs. 5(a) and 5(b) to aid species assignments. It is clear that the major observed transition near eBE=1.5 eV is P₂H⁺, as simulated in Fig. 5(d). This band is shown in Fig. 7 under conditions that minimize interferences and is considered in detail in Sec. III E. No transitions matching P₂H⁺ are observed, but the simulated spectrum in Fig. 5(e) is similar to the experimental spectra reported by Snodgrass et al. and by Jones et al. with EA(P₂)=0.589±0.025 eV. The observed band near eBE=1.0 eV appears to be due to HPPH− or cis-HPPH−. Figure 8 shows the HPPH− transition under conditions where P₂H⁺ is suppressed; this spectrum is considered further below. Judging by the intensities as the mass setting is varied, the band near eBE=2.9 eV likely arises from anions of the overlap with the ground-state anion. Barriers for interconversion of the neutral P₂H₂ isomers are expected to be high. The geometries of the calculated low-energy isomers of P₂H₁⁻ and P₂H₂⁻ are also shown in Fig. 6. Compared with the lowest-energy singlet H₂PP⁺ anion geometry with Cᵥ symmetry, the neutral molecule is asymmetric with the PPH plane containing the lone hydrogen rotated out of the anionic Cᵥ plane by about 72°. The singlet anion has two nearly degenerate conformations in Cᵥ symmetry, which correspond to saddle points on the neutral doublet surface. The triplet anion has the same asymmetric geometry as the neutral, but is unbound with respect to electron detachment. Because the anion has poor Franck–Condon overlap with the ground-state neutral, the harmonic approximation for Franck–Condon factors is not useful. In Fig. 5(h), we instead plot the simulated spectrum for vertical transitions from the C₆ anions to the C₆ saddle points on the neutral surface, with two anion conformations weighted for a 300-K population. The actual spectrum to the asymmetric neutral would be expected to be in the same energy region but with greater broadening because of the neglected displacement in the H₂P–PH internal rotation coordinate.

We now compare the simulated spectra in Figs. 5(c)–5(h) to the experimental spectra in Figs. 5(a) and 5(b) to aid species assignments. It is clear that the major observed transition near eBE=1.5 eV is P₂H⁺, as simulated in Fig. 5(d). This band is shown in Fig. 7 under conditions that minimize interferences and is considered in detail in Sec. III E. No transitions matching P₂H⁺ are observed, but the simulated spectrum in Fig. 5(e) is similar to the experimental spectra reported by Snodgrass et al. and by Jones et al. with EA(P₂)=0.589±0.025 eV. The observed band near eBE=1.0 eV appears to be due to trans-HPPH− or cis-HPPH−. Figure 8 shows the HPPH− transition under conditions where P₂H⁺ is suppressed; this spectrum is considered further below. Judging by the intensities as the mass setting is varied, the band near eBE=2.9 eV likely arises from anions of the
above would be a transition to the triplet surface in the re-
tatively assign this band to triplet HPPH, which as discussed
ization between the two possible nonplanar HPPH geom-

e beam. No origin transition can be discerned, which is con-
have the highly excited H 2 PP − isomer present in the ion
or origins of the PP-stretch mode progressions, tentatively assigned to
trans-HPPH as discussed in the text.

Figure 7 shows the spectrum of the P 2 H (2A′) ←P 2 H − (1A′) transition and a Franck–Condon fit with opti-
mized displacements and neutral frequencies. The spectrum
shown was taken on the low-mass side of the P 2 H − mass
peak and is predominantly due to P 2 H − . Two vibrational pro-
gressions can be identified. The calculations for neutral P 2 H
give frequencies of 2267 cm −1 (PH stretch), 666 cm −1 (PPH
bend), and 611 cm −1 (PP stretch). The first two frequencies
have the largest calculated normal-mode displacements be-
tween the singlet anion and the neutral, and can therefore be
assigned as the observed modes. We therefore assign the
2160±30 cm −1 progression to the PH stretch (r P ) and
the 630±20 cm −1 progression to the PPH bend (r H ). For calcu-
ulating the Franck–Condon fits, the frequencies for unob-
erved modes are fixed at the calculated values, scaled by
95% to match the two experimental frequencies. The peak
profiles are simulated by treating P 2 H and P 2 H − as asymmet-
ic rotors and convoluting over the instrumental resolution
function. The overall peak contour has a full width at half
maximum of 20 meV, and the effective rotational correction
of the origin peak position is −2 meV. The electron affinity
from the optimized fits is EA 0 (P 2 H )=1.514±0.010 eV. To
our knowledge, there are no previous experimental reports of
this electron affinity.

Because the PP-stretching mode is not observed, the ex-
act geometry change cannot be derived from the data. We
obtain approximate values by fixing the displacement for that
mode at the (small) calculated value, and using the Franck–
Condon intensities to fit the displacements in the other two
modes. This gives estimates of an increase of 0.023 Å for the
PP distance, an increase of 0.068 Å for the PH distance, and
an increase of 8° in the PPH angle from the neutral to the
anion. The calculated neutral geometry is r(PP)=2.007 Å,
r(PH)=1.435 Å, and α=98°; experimental values are un-
known.

F. Diphosphiene, HPPH

The experimental spectrum of the HPPH − band is shown
on an expanded scale in Fig. 8. The signal is weak, but there
is a clear vibrational progression starting at an origin at
e BE=1.00 eV with a frequency of about 600 cm −1 . As noted
above, the trans-HPPH − and cis-HPPH − isomers have very
similar calculated energies, frequencies, and simulated
Franck–Condon profiles. Therefore, a definitive isomeric as-
signment of the experimental spectrum is not possible. The
major vibrational progression can be fit starting with the
simulated spectra of either isomer and making minor adjust-
ments to the PP-stretch mode frequency (calculated as
616 cm −1 in trans-HPPH and 603 cm −1 in cis-HPPH) and
geometry displacements. The other two active, totally sym-
metric vibrational modes—the PH symmetric stretches with
calculated frequencies of 2335 cm −1 and 2366 cm −1 for trans
and cis, respectively, and the PPH symmetric in plane bends
consistent with a broad, nonvertical transition as expected from
H 3 PPH − to the asymmetric H 3 PPH neutral. This band is too
weak for more detailed analysis.
at 970 and 741 cm\(^{-1}\) — have small calculated Franck–Condon factors and are not observed. The activity of the PP-stretch mode results from the removal of an electron from the highest-occupied molecular orbital in the anion, which is essentially a phosphorus out-of-plane \(p\pi^*\) antibonding orbital. The \textit{trans}-HPPH parameters do provide a slightly better fit, consistent with the 0.1-eV lower energy calculated for \textit{trans}-HPPH\(^-\) compared with \textit{cis}-HPPH\(^-\), but either fit of the major progression is acceptable given the quality of the data. However, there is an additional weaker secondary vibrational progression that is not reproduced by either simulation. It has an apparent origin at \(600\) cm\(^{-1}\). The 240-cm\(^{-1}\) spacing between the origin of the major progression at 1.00 eV and the peak at 1.03 eV is too small to match any of the vibrational modes of either \textit{trans}-HPPH or \textit{cis}-HPPH. Therefore, we provisionally assign this progression to the isomer \textit{other} than the one responsible for the major progression. Using the theoretical energies as a guide, we can tentatively identify the major progression as \textit{trans}-HPPH\(^-\) with \(EA_0(\textit{trans}-\text{HPPH}) = 1.00\pm0.01\) eV and the secondary progression as \textit{cis}-HPPH\(^-\) with \(EA_0(\textit{cis}-\text{HPPH}) = 1.03\pm0.01\) eV. Overlap of the two isomer transitions and the weak signal intensity preclude further analysis.

A more remote possibility for the assignment of the secondary progression is to the \(1\rightarrow1\) hot sequence band of the out-of-plane bending mode, noting that \(\Delta v=0\) transitions are allowed for nontotally symmetric modes. The out-of-plane bend exhibits a large calculated frequency change from 449 cm\(^{-1}\) (289 cm\(^{-1}\)) in the anion to 775 cm\(^{-1}\) (699 cm\(^{-1}\)) in the neutral for the \textit{trans}(cis) isomer. These transitions are seen in the simulation in Fig. 8(b) for the \textit{trans} isomer as small peaks at \(eBE=1.042, 1.117,\) and 1.192 eV. Fitting the intensities and transition energies of this mode to the experimental peaks would require a shift of frequency of 8\% (21\% for \textit{cis}) from the calculated value for the neutral and a non-Boltzmann vibrational population of the anion, at least in the harmonic-oscillator approximation. Therefore, this assignment is less satisfactory. Observation of these sequence transitions in a higher-resolution experiment could help confirm the identities of the isomers.

### G. Thermochemistry

The electron affinities determined in this work are collected in Table I and compared with literature and calculated values. The gas-phase acidities of PH, PH\(_2\), and PH\(_3\) can be derived from the electron affinities reported here and the bond dissociation energies from Berkowitz et al.\(^{58–62}\) using the negative-ion thermochemical cycle, \(\Delta_{\text{acid}} H(RH) = D(RH) + IE(H) - EA(R)\).

<table>
<thead>
<tr>
<th>Value</th>
<th>PH</th>
<th>PH(_2)</th>
<th>PH(_3)</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>(EA_0(R)) (eV)</td>
<td>0.7465±0.0003</td>
<td>1.027±0.006</td>
<td>1.263±0.006</td>
<td>Expt.(^a)</td>
</tr>
<tr>
<td>(D_{\text{f}}(R-H)) (kJ mol(^{-1}))</td>
<td>293.3±2.1</td>
<td>&gt;312.5±2.1</td>
<td>=345.0±1.9</td>
<td>Expt.(^b)</td>
</tr>
<tr>
<td>(\Delta_{\text{acid}} H_0(RH)) (kJ mol(^{-1}))</td>
<td>1533.3±2.1</td>
<td>&gt;1525.6±2.2</td>
<td>=1535.2±2.0</td>
<td>Expt.(^c)</td>
</tr>
<tr>
<td></td>
<td>1534.6</td>
<td>1530.9</td>
<td>1529.4</td>
<td>Theory(^d)</td>
</tr>
<tr>
<td>(\Delta_{\text{acid}} G_{298}(RH)) (kJ mol(^{-1}))</td>
<td>1537.6±2.1</td>
<td>&gt;1530.9±2.2</td>
<td>=1541.3±2.0</td>
<td>Expt.(^e)</td>
</tr>
<tr>
<td>(\Delta_{\text{acid}} G_{298}(RH)) (kJ mol(^{-1}) K(^{-1}))</td>
<td>80.1±2</td>
<td>96.7±2</td>
<td>105.9±2</td>
<td>Expt.(^f)</td>
</tr>
<tr>
<td>(\Delta_{\text{acid}} G_{298}(RH)) (kJ mol(^{-1}))</td>
<td>1513.7±2.2</td>
<td>&gt;1502.1±2.2</td>
<td>=1509.7±2.1</td>
<td>Expt.(^g)</td>
</tr>
<tr>
<td></td>
<td>1520±8.4</td>
<td></td>
<td></td>
<td>Expt.(^h)</td>
</tr>
</tbody>
</table>

\(^a\)From Table I.
\(^b\)CCSD(T) with complete basis-set extrapolation, zero-point energy, and spin–orbit corrections from Ricca and Bauschlicher (Ref. 14).
\(^c\)Gas-phase phase acidities derived using IE(H)=0.5 hartree.
\(^d\)Berkowitz and co-workers (Refs. 58–62).
\(^e\)\(\Delta_{\text{acid}} H(RH) = D(RH) + IE(H) - EA(R)\).
\(^f\)Thermal enthalpy correlations and entropies calculated using the harmonic-oscillator, rigid-rotor approximations: \(\Delta H_{298} = \Delta H_0 + \int_0^{298} \Delta C_v dT\) and \(\Delta S_{298} = \int_0^{298} (\Delta C_P/T) dT\).
\(^g\)\(\Delta G = \Delta H - T\Delta S\). 
\(^h\)Ion cyclotron resonance equilibrium (Ref. 65).
with theoretical calculations reported by Ricca and Bauschlicher, Jr. at the CCSD(T) level with complete basis-set extrapolation, zero-point energy corrections, and spin–orbit energy corrections. The theoretical values for the three electron affinities and for $D_0(P-H)$ are in superb agreement with experiment, within about 1 kJ/mol. The theoretical values of $D_0(HP-H)$ and $D_0(H_2P-H)$ are higher and lower by about 5 kJ/mol than the experimental lower and upper limits, respectively, which is consistent and implies that the experimental upper limit for the appearance energy of PH$_3^-$ from PH$_3$ could be higher than the true value by that amount. The deviations between theory and experiment for the gas-phase acidities in Table V mirror those of the dissociation energies.

IV. CONCLUSION

We have reported electron affinities and other molecular parameters for small phosphorus hydride species. Simulated Franck–Condon profiles using geometries and normal modes from density-functional theory have proven very useful for identifying the bands in photoelectron spectra. Without high-quality theoretical input, the assignment of the P$_2$H$_5^-$ species would be difficult. Accurate molecular structure calculations and the development of software for routine Franck–Condon simulations of photoelectron spectra using calculated vibrational force fields make the spectroscopic assignments possible.

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