SPECTROSCOPIC STUDIES OF MOLECULAR STRUCTURE
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by
G. HERZBERG

Division of Physics, National Research Council of Canada, Ottawa, Canada
K1A OR6

A. INTRODUCTION

The citation for the 1971 Nobel Prize in Chemistry reads “for contribution to the knowledge of electronic structures and geometry of molecules, especially free radicals” and therefore implies that the Prize has been awarded for a long series of studies extending practically over my whole scientific life. I shall try to present in this lecture a few of what I consider to be the more significant results of this work.

It was recognized very early in the development of spectroscopy that the study of molecular spectra is one of the most important tools for the determination of molecular structures. When I began my scientific work it had already been firmly established that band spectra are molecular spectra (in contrast to line spectra which are atomic spectra) and that a band system such as the one shown in Fig. 1 represents all transitions between the vibrational and rotational levels of two electronic states, i.e. corresponds to a single line or a single multiplet in an atomic spectrum.

It was also well-known how, by determining the spacing between appropriate bands in a band system, we obtain the vibrational intervals in the upper and lower electronic states, which are simply and directly related to the vibrational frequencies of the molecule in these states and how, by determining the separations of appropriate lines in a given band, we obtain the rotational intervals in these states which are directly related to the moments of inertia and therefore to the internuclear distances (or in other words the geometrical structures) in the two states.

The fine structure of the rotational lines tells us something about the nature of the electronic states involved, whether they are singlet, doublet, triplet, . . . states while the nature and number of branches tells us, in the case of linear polyatomic molecules and diatomic molecules, whether the electronic states are of the type \( \Sigma^+ \) or \( \Sigma^- \), or \( \Pi \), or \( A \).

In the years 1925-28 the work of James Franck (2) and Birge and Sponer (3) had established how in suitable spectra where a sufficient number of vibrational levels or the limit of an absorption continuum has been observed the dissociation energy can be accurately determined.
Fig. 1.
Band system of the PN molecule after Curry, Herzberg and Herzberg (1).
B. DIATOMIC MOLECULES, RADICALS AND IONS

The spectra of a large number of diatomic molecules, radicals and ions (~600) have been studied by various investigators in the past 50 years. For many of these molecules several band systems have been found and therefore several (sometimes many) electronic states have been established allowing a detailed interpretation in terms of molecular orbital theory [see Mulliken’s Nobel lecture (4)]. If Rydberg series of electronic states are observed (so far only in relatively few cases) it is possible to determine the ionization potential in much the same way as for atoms.

In the following I shall describe the results only with regard to a few of the many diatomic systems studied in our laboratory.

(1) \(H_2, D_2, HD, H_2^+\)

The simplest molecular systems, \(H_2\) and \(H_2^+\) are even now the only ones for which \textit{ab initio} calculations of very high precision can be and have been carried out [Kolos and Roothaan (5), Kolos and Wolniewicz (6), Hunter and Pritchard (7)]. It therefore appeared of considerable interest to improve as much as possible the experimental accuracy of some of the molecular constants. We have attempted to do this for the dissociation energy, the vibrational and rotational intervals in the ground state and for the ionization potential.

![Potential functions of the ground state \((X^1\Sigma_g^+)\) and the second excited state \((B'^1\Sigma_u^+)\) of \(H_2\) showing the relation between the absorption limit and the dissociation energy in the ground state: \(D_0^0 = \nu_{\text{limit}} - E(H, n=2)\).](image-url)
The dissociation energy has been determined from the long wavelength limit of the continuum that joins onto the discrete absorption bands, which correspond to transitions from the ground state to an excited electronic state designated $B$ in Fig. 2. This state dissociates into one normal and one excited ($n = 2$) H atom; therefore the limit of the continuous absorption spectrum occurs at an energy equal to the sum of the dissociation energy of the ground state and the excitation energy of the H atom. As an example Fig. 3 shows a small section of the far ultraviolet absorption spectrum of D$_2$ near 840 Å under very high resolution taken at liquid nitrogen temperature where the absorption limit is clearly visible (for H, the corresponding limit is overlapped by an absorption line which makes it difficult to obtain as precise a limit as for D$_2$). Subtracting the excitation energy of D (or H) from the wave number of the limit and including a very small correction for the rotational barrier at $J = 1$ in the upper state we have obtained the dissociation energies given in the last column of Table 1 which should be compared with the theoretical values in the second column. The agreement between theory and experiment is clearly very good.
Table 1. Calculated and observed dissociation energies of hydrogen

<table>
<thead>
<tr>
<th></th>
<th>Theor. (a)</th>
<th>Obs. (b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$D_0^0(\text{H}_2)$</td>
<td>36 117.9</td>
<td>$&lt;36 118.3$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$&gt;36 116.3$</td>
</tr>
<tr>
<td>$D_0^0(\text{HD})$</td>
<td>36 405.5</td>
<td>${36 406.6$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$36 405.8$</td>
</tr>
<tr>
<td>$D_0^0(\text{D}_2)$</td>
<td>36 748.2</td>
<td>36 748.9 $\pm$ 0.4 cm$^{-1}$</td>
</tr>
</tbody>
</table>

(a) From Kolos and Wolniewicz (6) but including small non-adiabatic corrections according to Bunker (9).

Precise values of the vibrational intervals $\Delta G$ and the rotational constants $B_v$ have been obtained for low $v$ values from the Raman spectrum [Stoicheff (10)] and the infrared quadrupole spectrum [Herzberg (11) Rank and collaborators (12)], and for higher $v$ values from the vacuum ultraviolet emission spectrum [Herzberg and Howe (13), Bredohl and Herzberg (14)]. In Table 2 the observed $\Delta G$ values of $\text{H}_2$ are compared with the theoretical values obtained from the Kolos-Roothaan-Wolniewicz potential (6). The slight systematic differences are in all probability due to the neglect of non-adiabatic corrections [see Poll and Karl (15) and Bunker (9)]. In Fig. 4 the deviations are plotted for $\text{H}_2$, HD and $\text{D}_2$. Similar very small systematic differences are also found for the rotational constants $B_v$ as shown in Fig. 5. There is little doubt that these small discrepancies, of the order of 100 p.p.m., will be fully accounted for once the non-adiabatic corrections have been accurately evaluated.

If one disregards adiabatic and non-adiabatic corrections one obtains from the observed $B_v$ values an internuclear distance for the equilibrium position of

Table 2. Observed and calculated vibrational quanta in the ground state of $\text{H}_2$

| $v$ | $\Delta G(v+1|\ell_i)$ |
|-----|------------------------|
| obs. | theor. |
| 0   | 4161.14                | 4162.06     |
| 1   | 3925.98                | 3926.64     |
| 2   | 3695.24                | 3696.14     |
| 3   | 3468.01                | 3468.68     |
| 4   | 3241.56                | 3242.24     |
| 5   | 3013.73                | 3014.49     |
| 6   | 2782.18                | 2782.82     |
| 7   | 2543.14                | 2543.89     |
| 8   | 2292.96                | 2293.65     |
| 9   | 2026.26                | 2026.81     |
| 10  | 1736.66                | 1737.13     |
| 11  | 1414.98                | 1415.54     |
| 12  | 1049.18                | 1048.98     |
| 13  | 621.96                 | 620.16      |
Deviations of the observed vibrational quanta of H₂, HD, and D₂ from those obtained from theory.

0.74139 and 0.74156 Å for H₂ and D₂, while the theoretical value (the same for both) is 0.74140 Å.

Finally the detailed study of the Rydberg series of H₂ near 800 Å, which is made somewhat complicated by interesting perturbations, by preionization effects and small pressure shifts, has led to the following experimental value for the ionization potential of H₂ [Herzberg and Jungen (16)]

\[ \text{I.P. (H}_2) = 124417.2 \pm 0.4 \text{ cm}^{-1} \]
Deviations of the observed rotational constants $B_v$ of H$_2$, HD, and D$_2$ from those obtained from theory.

and a less detailed study by Takezawa (17) of D$_2$, after a pressure shift correction, to the value

$$\text{I.P. (D}_2\text{)} = 124\,745.6 \pm 0.6 \text{ cm}^{-1}$$

while the theoretical values, including relativistic and Lamb shift corrections, are [Hunter and Pritchard (7), Jezierski and Kołos (18), Bunker (9)]

$$\text{I.P.}\text{theor}(\text{H}_2) = 124\,417.3 \text{ cm}^{-1}$$

$$\text{I.P.}\text{theor}(\text{D}_2) = 124\,745.2 \text{ cm}^{-1}$$

The agreement is well within the error of measurement, i.e. within about 3 p.p.m.

From the ionization potentials and the dissociation energies of H$_2$ and D$_2$ we obtain, according to the general relation

$$D_0(\text{X}_2^+) = \text{I.P. (X)} + D_0(\text{X}_2) - \text{I.P. (X)}$$

the following values for the dissociation energies of the ions
\[ D_0^{\text{obs.}}(\text{H}_2^+) \leq 21 \, 379.9 \pm 0.4 \, \text{cm}^{-1} \]
\[ D_0^{\text{obs.}}(\text{D}_2^+) = 21 \, 711.9 \pm 0.6 \, \text{cm}^{-1} \]

These values may be compared with the theoretical values
\[ D_0^{\text{theor.}}(\text{H}_2^+) = 21 \, 379.3 \, \text{cm}^{-1} \]
\[ D_0^{\text{theor.}}(\text{D}_2^+) = 21 \, 711.6 \, \text{cm}^{-1} \]

which are considered to be accurate to \( \pm 0.2 \, \text{cm}^{-1} \). That the agreement is again very satisfactory is not surprising since \( D_0(\text{H}_2^+, \text{D}_2^+) \) are determined by \( D_0(\text{H}_2, \text{D}_2) \) and I.P.(\text{H}_2, \text{D}_2) and for both of the latter quantities very good agreement between theory and experiment was found.

(2) \( \text{O}_2 \)

The study of forbidden electronic transitions, that is, transitions forbidden by the normal selection rules, has greatly aided in the understanding of the electronic structure of diatomic (and polyatomic) molecules. The \( \text{O}_2 \) molecule is particularly rich in such forbidden transitions. The lowest electron configuration \( \ldots \pi_u \, \pi_g \) gives rise to three states \( ^3 \Sigma_u^-, ^1 \Lambda_g \) and \( ^1 \Sigma_g^* \) of which the first forms the ground state of the molecule. As was first recognized by Van Vleck (19) the forbidden transitions from the ground state to the \( ^1 \Delta_g \) and \( ^1 \Sigma^+ \) states can only occur as magnetic dipole (or much more weakly as quadrupole) radiation. Because of the long absorption path in the atmosphere, these forbidden transitions are very prominent in the solar spectrum observed from the ground. In emission they occur strongly in the red and infrared spectrum of the night sky. Even the transition \( ^1 \Sigma_g^* \rightarrow ^1 \Delta_g \) which can occur only as electric quadrupole radiation has been observed [Noxon (20)].

Another group of forbidden transitions in the near ultraviolet corresponds to transitions to states of the configuration \( \ldots \pi_u \, \pi_g \) which gives rise to the six states \( ^1 \Sigma_u^+, ^1 \Sigma_u^-, ^1 \Lambda_u, ^3 \Sigma_u^+, ^3 \Sigma_u^-, ^3 \Delta_u \). Only one of these states can combine with the ground state in an allowed transition, namely, \( ^3 \Sigma_u^- \rightarrow ^3 \Sigma_g^- \); this transition, the well-known Schuman-Runge bands, limits the transparency of air in the ultraviolet. Forbidden transitions to three of the other states, \( ^3 \Sigma_u^-, ^3 \Sigma_u^+ \) and \( ^3 \Delta_u \) have been observed [Herzberg (21)], of which \( ^3 \Sigma_u^- \rightarrow ^3 \Sigma_g^- \) is the best known and is quite prominent in the light of the night sky. Fig. 6 shows a potential diagram of the lower electronic states of \( \text{O}_2 \). The fact that all transitions to the ground state from non-repulsive states arising from normal atoms \((P + P)\) are forbidden accounts for the observation that in the upper atmosphere the radiative recombination of \( \text{O} \) atoms is very slow indeed. On the other hand the weak continuous absorption joining onto the \( ^3 \Sigma_u^- \rightarrow ^3 \Sigma_g^- \) absorption system \((\lambda < 2440 \, \text{Å})\) gives rise to the production of free \( \text{O} \) atoms even at fairly low altitudes and thus accounts for the formation of the ozone layer. Thus the chemistry of the upper atmosphere is greatly affected by the forbidden nature of these transitions i.e. by the electronic structure of the \( \text{O}_2 \) molecule. In Table 3 some of the molecular constants of \( \text{O}_2 \) are summarized and compared with those of \( \text{S}_2 \). In \( \text{S}_2 \), the analogues of the forbidden transitions of \( \text{O}_2 \) have not yet been observed.
Fig. 6.
Potential functions of the ground state and the lower excited states of the O₂ molecule.
The state C \(^3\Delta_u\) should have been designated \(A' \ 3\Delta_u\) to be in accord with Table 3.

### Table 3. Molecular constants of the lower states of the O₂ and S₂ molecules

<table>
<thead>
<tr>
<th></th>
<th>(T_0(\text{cm}^{-1}))</th>
<th>(\omega_0(\text{cm}^{-1}))</th>
<th>(D_0(\text{eV}))</th>
<th>(r_e(\text{Å}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>O₂</td>
<td>(B \ 3\Sigma_u^-)</td>
<td>49 363.1</td>
<td>700.36</td>
<td>0.9627</td>
</tr>
<tr>
<td>A</td>
<td>(a \ 3\Sigma_u^+)</td>
<td>35 007.2</td>
<td>799.07</td>
<td>0.7752</td>
</tr>
<tr>
<td>A'</td>
<td>(3\Delta_u)</td>
<td>34 320</td>
<td>(611.2)</td>
<td>0.8604</td>
</tr>
<tr>
<td>c</td>
<td>(1\Sigma_u^-)</td>
<td>32 664.1</td>
<td>794.29</td>
<td>1.0657</td>
</tr>
<tr>
<td>b</td>
<td>(1\Sigma_g^+)</td>
<td>13 120.9</td>
<td>1432.69</td>
<td>3.4887</td>
</tr>
<tr>
<td>a</td>
<td>(1\Delta_g)</td>
<td>7 882.4</td>
<td>1599.3</td>
<td>4.1382</td>
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<tr>
<td>X</td>
<td>(1\Sigma_g^-)</td>
<td>0</td>
<td>1580.36</td>
<td>5.1155</td>
</tr>
<tr>
<td></td>
<td>(S_2)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>B</td>
<td>(3\Sigma_u^-)</td>
<td>31 689</td>
<td>434.0</td>
<td>1.5828</td>
</tr>
<tr>
<td>A</td>
<td>(2\Sigma_u^+)</td>
<td>(22 682)</td>
<td>481.4</td>
<td>(1.55)</td>
</tr>
<tr>
<td>A'</td>
<td>(2\Delta_u)</td>
<td>(21 855)</td>
<td>488.4</td>
<td>(1.66)</td>
</tr>
<tr>
<td>c</td>
<td>(1\Sigma_u^-)</td>
<td></td>
<td></td>
<td>2.15</td>
</tr>
<tr>
<td>b</td>
<td>(1\Sigma_g^+)</td>
<td>(699.7)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a</td>
<td>(1\Delta_g)</td>
<td>(5,100)</td>
<td>701.94</td>
<td>(3.73)</td>
</tr>
<tr>
<td>X</td>
<td>(1\Sigma_g^-)</td>
<td>0</td>
<td>725.65</td>
<td>4.3662</td>
</tr>
</tbody>
</table>
Section of vacuum ultraviolet absorption spectrum of CH showing the O-O bands of the $E^2\Pi - X^2\Pi$ and $F^2\Delta - X^2\Pi$ transitions after Herzberg and Johns (28).
Many studies of forbidden transitions in other molecules have been made but will not be discussed here [see Herzberg (22), (23)].

(3) CH and CH'  
The visible and near ultraviolet emission spectrum of the CH radical has been known ever since spectra of an ordinary Bunsen burner have been taken. That the well-known bands at 4314 and 3889 Å really belong to CH was established of course only after the theory of diatomic spectra was sufficiently developed [Heurlinger (24), (25), Hultén (26)]. The band at 3 145 Å later discovered by Hori (27) in electric discharges has the same lower state as the other two, the 'II ground state of the molecule. While the ground state has the electron configuration \( \sigma^2 \sigma^2 \sigma^2 \pi \) the three excited states have the configuration \( \ldots \sigma \pi^2 \). This configuration, in addition to \( 2\Sigma^- \), \( 2\Delta \) and \( 2\Sigma'^+ \) also gives rise to a \( 4\Sigma^- \) state which however has not yet been observed even though it is expected to be a stable and fairly low-lying state.

The CH radical is a very reactive radical which under most conditions has a very short life time. It is for this reason that its absorption spectrum has been observed and studied in detail only fairly recently [Herzberg and Johns (28)]. This study has revealed a number of new transitions in the vacuum ultraviolet of which Fig. 7 gives an example. Indeed a whole Rydberg series was found which allowed an accurate determination of the ionization potential \( = 10.64 \text{ eV} \). In addition the value for the dissociation energy was slightly refined in this work and much evidence of predissociation in all excited states except the \( 2\Delta \) state was obtained.

Fig. 8. shows a diagram of the observed electronic states of CH and Table 4 gives some of the principal molecular constants.

The CH radical was the first molecule recognized in the interstellar medium [Swings and Rosenfeld (29), McKellar (30)]. The life time of CH under the action of the interstellar radiation field is relatively short (about 30 years) because of the predissociation that has been established to occur in all absorption systems of CH except the longest wavelength one \( (2\Delta—2\Pi) \). It is therefore somewhat surprising that CH is present in sufficient concentration to show its absorption spectrum.

The ion CH' has also been observed, first in interstellar absorption [Adams (31)] and then in emission in the laboratory [Douglas and Herzberg (32)]. The observed transition is the expected \( \sigma \pi \Pi—\sigma \pi 1 \Sigma \) transition. The molecular constants are included in Table 4. Fig. 9 shows a laboratory spectrum. The interstellar lines \( \text{[R (0) lines coming from the lowest rotational level]} \) are marked. In spite of much effort we have not yet been able to observe this spectrum \textit{in absorption} in the laboratory.

(4) C_2  
A few years ago, in an attempt to observe absorption spectra of CH, C_2, CH', and CH', using flash discharges through CH, we observed a new very simple spectrum shown in Fig. 10 [Herzberg and Lagerqvist (33)]. The analysis was very easy and showed immediately and conclusively that the carrier of this
Fig. 8.
Energy level diagram of the electronic states of the CH radical showing the observed transitions. The Rydberg transitions are not marked.
Table 4. Molecular constants of the known electronic states of CH and CH+.

<table>
<thead>
<tr>
<th>State</th>
<th>$T_0$ (cm$^{-1}$)</th>
<th>$\Delta G(1/2)$ (cm$^{-1}$)</th>
<th>$B_0$ (cm$^{-1}$)</th>
<th>$r_0$(Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH+</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$^1\Pi$</td>
<td>109446</td>
<td>1642.16</td>
<td>11.428</td>
<td>1.2596</td>
</tr>
<tr>
<td>$^3\Sigma^+$</td>
<td>85850</td>
<td>3251.5</td>
<td>13.932</td>
<td>1.1409</td>
</tr>
<tr>
<td>CH</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$^3\Delta$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$^3\Pi$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$^3\Sigma^+$</td>
<td>3d</td>
<td>72960</td>
<td>12.6</td>
<td>1.20</td>
</tr>
<tr>
<td>$^3\Pi_T$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$^3\Sigma^+$</td>
<td>3p</td>
<td>64532</td>
<td>12.17</td>
<td>1.221</td>
</tr>
<tr>
<td>$^3\Pi_L$</td>
<td>58981</td>
<td>2613</td>
<td>13.7</td>
<td>1.12</td>
</tr>
<tr>
<td>$^3\Sigma^+$</td>
<td>C</td>
<td>31778</td>
<td>14.25</td>
<td>1.128</td>
</tr>
<tr>
<td>$^3\Sigma^-$</td>
<td>B</td>
<td>23698</td>
<td>12.64</td>
<td>1.198</td>
</tr>
<tr>
<td>$^3\Delta$</td>
<td>A</td>
<td>23217</td>
<td>14.58</td>
<td>1.115</td>
</tr>
<tr>
<td>$^3\Pi$</td>
<td>X</td>
<td>0</td>
<td>14.19</td>
<td>1.130</td>
</tr>
</tbody>
</table>

Fig. 9. Two emission bands of CH' after Douglas and Herzberg (32). The interstellar lines are the R(0) lines.
Fig. 10.
Absorption spectrum of the Ca⁺ ion after Herzberg and Lagerqvist (33).
spectrum must be the C\(_2\) radical or one of its ions. It seemed impossible to fit this transition into the known system of energy levels of C\(_2\). Therefore and because of the similarity with N\(_2^+\) we suggested that the spectrum belongs to the C\(_2^-\) ion (i.e. represents the first discrete spectrum of a negative molecular ion) even though the required doublet structure of the spectrum was not very clearly recognizable.

In Fig. 11 the observed electronic energy levels of N\(_2^+\), CN and CO\(^+\) are compared with those of the new transition. All these molecules or ions have 13 electrons. Comparisons such as this historically formed the starting point of molecular orbital theory. Here the comparison serves to strengthen the suggestion that the new transition belongs to a 13 electron system, that is, C\(_2^-\). This suggestion was strikingly confirmed first by the work of Milligan and Jacox (34) on the analogous spectrum observed in a matrix since it is considerably strengthened by the addition of an electron donor like Cs; and recently even more conclusively by the work of Lineberger and Patterson (35) who photo-ionized a C\(_2^-\) beam by a two-photon absorption of a tunable laser beam and found, as shown in Fig. 12, a photoionization exactly at the wave lengths of the new absorption bands.

Meinel (36) in our laboratory has recently observed the spectrum of C\(_2^-\). Other diatomic molecules, radicals and ions studied in our laboratory are listed
in Table 5. For all these systems, information about their geometrical structure (internuclear distances), their vibrational frequencies in various electronic states, and about their electronic structure has been obtained. Much work on these molecules and many other molecules which we have not studied has proceeded in many laboratories throughout the world notably here in Stockholm in Professor Lagerqvist’s laboratory.

Table 5. Diatomic molecules, radicals and ions studied at the National Research Council of Canada

H₂, He₂, B₂, C₂, N₂, O₂, F₂, Mg₂, Si₂, P₂, S₂, Cl₂, I₂
LiH, BH, CH, NH, OH, SiH, PH, SH, HCl, HBr, CrH, CuH
BN, CN, PN
CO, NO, PO, SO, IO
BS, CS, SnS
BF, NF, NaF, SiF, PF, KF, RbF
BCl, CCl, SiI, NCl, AlCl
NBr, AlBr, CP, AIC
CH⁺, NH⁺, SiH⁺, PH⁺
C⁻, C₂⁻, N⁺, N₂⁺, P⁺, CO⁺, CN⁺, NO⁺
C. Polyatomic Molecules, Radicals and Ions

(1) General remarks

In polyatomic molecules and radicals there are in general several geometrical parameters which are required to describe their structures (rather than one, \( r_e \), for diatomic molecules). For a large number of stable molecules these parameters have been determined in their ground states by the techniques of infrared, Raman and microwave spectroscopy. In excited electronic states these parameters can only be obtained by a detailed study of electronic band systems in absorption or emission. In many cases it has been found that the shape (point group) of the molecule in an excited state is different from its shape in the ground state. For example the molecules \( \text{C}_2\text{H}_2 \) and HCN, well-known to be linear in their ground states are found to be strongly bent in their first (singlet) excited states as shown in Fig. 13. It is clear that this behaviour throws an interesting light on the way in which the electronic structure determines molecular shape.

For polyatomic free radicals and ions one is dependent both for the ground states and the excited states on the study of electronic spectra to obtain the shapes and the geometrical parameters. Only for a few radicals have microwave and infrared spectra been obtained in the gaseous state. Not infrequently, just as for stable molecules, electronic spectra of radicals are diffuse or even continuous in which case no information about geometrical structure can be obtained. The \( \text{BH}_3 \) radical may be such a case. An additional difficulty in the interpretation of free radical spectra is the problem of identification, that is the problem to which particular radical an observed spectrum belongs. As an example of these difficulties I should like to discuss the history of the discovery of the spectra of \( \text{CH}_3 \) and \( \text{C}_3 \).
In 1941 the Belgian astronomer Swings wrote to me about a problem that had arisen in the interpretation of the spectra of comets. Fig. 14 shows the spectrum of a comet. In this spectrum the emission bands of CN, C₂, CH, NH, OH can be clearly seen, but in addition there is a group of bands near 4050 Å whose origin nobody had been able to identify at that time. On the basis of the structure of this spectrum, I thought I could eliminate the possibility that it was due to a diatomic free radical. Rather, the 4050 group appeared to me like a perpendicular band of a nearly symmetric top molecule, and because of the wide spacing of the subbands I concluded (38) that it must be due to a nonlinear molecule XH₂ with a bond angle of the order of 140°. The most likely identification appeared to be CH₂, particularly since at that time Mulliken (39) had just predicted a spectrum of CH₂ to occur in the region 4000-4500 Å. Since CH was known to be present in comets, the identification of the 4050 group as due to CH₂ seemed eminently reasonable.

On the basis of these considerations I proceeded to do some laboratory experiments. I tried the obvious, passing a discharge through methane (CH₄) in the hope of obtaining in this discharge a spectrum of CH₂. While the continuous discharge through methane showed only well-known features such as CH and H₂, I noticed that the colour of the discharge in the first instant after it was turned on was slightly different from the later colour. I therefore took a spectrum with the discharge turned on and off repeatedly. On this spectrum, in addition to the bands of CH₂, a new feature appeared precisely at 4050 Å which agreed in almost every detail with the 4050 group as observed in comets. This agreement is shown in Fig. 15. Thus, for the first time [Herzberg (40)], the 4050 group of comets had been observed in the laboratory, and this had been done by choosing conditions suggested by the assumption that the spectrum was due to CH₂. Therefore, it was perhaps excusable that I felt confirmed in my belief that this spectrum was due to CH₂.

However, in 1949 Monfils and Rosen (41) at Liege repeated our experiment but replaced the hydrogen by deuterium. The spectrum that they observed was identical in every detail with the spectrum that I had observed,
whereas of course small isotope shifts would have been expected had this spectrum been due to \( \text{CH}_2 \). Dr. Douglas and I at Ottawa immediately repeated this experiment using much higher resolution and confirmed the result of the Belgian physicists, thus establishing without doubt that neither the cometary spectrum nor the laboratory spectrum was due to \( \text{CH}_2 \). Douglas (42) then proceeded to find the true carrier of this spectrum by using methane with \( ^{13}\text{C} \) in it [supplied by the late K. Clusius]. He observed that the main emission band at 4050 Å in a 50-50 mixture of \( ^{12}\text{CH}_4 \) and \( ^{13}\text{CH}_4 \) was replaced by six bands, showing immediately that the molecule responsible for this spectrum must have three carbon atoms in it. Further consideration of the fine structure of this band left no doubt that the spectrum must be due to the free \( \text{C}_3 \) radical. At the time when Douglas established this conclusion, the \( \text{C}_3 \) radical had not even been postulated in any chemical reaction, but since that time it has been found to be one of the dominant constituents of carbon vapour as obtained by the evaporation of graphite.

The question now arose, if the 4050 group is not due to \( \text{CH}_2 \), where is the true spectrum of \( \text{CH}_2 \)? or does such a spectrum not exist? It was almost ten years after the identification of the carrier of the 4050 group before a spectrum of \( \text{CH}_2 \) was found. From photochemical evidence it was well known that there are two molecules which on photolysis give \( \text{CH}_2 \) - namely, ketene (CH\(_2\)CO) and diazomethane (CH\(_2\)N\(_2\)). Since the latter compound is rather explosive, we began by studying the continuous photolysis of ketene and, when that failed, by turning to flash photolysis, which had in the meantime been developed [Norrish and Porter (43)]. Even though we extended our search into the vacuum ultraviolet, we did not find a spectrum of \( \text{CH}_2 \). As a last resort we decided that we should try diazomethane in spite of its hazardous properties. Almost the first absorption spectrum of flash-photolyzed diazomethane showed

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Fig. 15.
The \( \lambda 4050 \) group in the laboratory and in a comet after Herzberg (40).
a new transient feature (of a lifetime of about 10 µsec), which turned out to be the spectrum of CH₂. We did not observe this particular feature in the flash photolysis of ketene because ketene itself absorbs strongly at this particular wavelength.

We were fortunate in being able to obtain immediately, with the help of Dr. Leitch at Ottawa, a quantity of deuterated diazomethane, and in this way were able to verify, as shown by Fig. 16, that the new feature, at 1415 Å, actually does shift when hydrogen is substituted by deuterium. Thus, at least we were sure that the molecule or radical responsible for this feature contained hydrogen, but of course this observation did not yet prove that the radical was CH₂.

Spectra taken at high resolution in partially or fully deuterated diazomethane showed three different positions for the principal band depending on the deuterium concentration. These spectra, reproduced in Fig. 17, show for two of the isotopes a clear and simple fine structure. For CD₂ there is in addition a characteristic intensity alternation indicating a symmetric position of the two D atoms. The simple structure of the bands suggests that the molecule is linear. According to molecular orbital theory the ground state of linear CH₂ must be

\[ \ldots \sigma_{g}^\pi \tau_{u}^{\pi} \Sigma_{g}^{-} \]

and in agreement with that the odd lines are strong in the CD₂ band. We concluded therefore that the ground state of CH₂ is a triplet state and that the molecule is linear even though the triplet splitting was not resolved.

During the last year several theoretical calculations (45), (46) as well as electron-spin resonance studies (47), (48) of CH₂ and CD₂ in inert matrices
have strongly suggested that CH$_2$ in its lowest triplet state is not linear. If that is so, the molecule is a symmetric top and in addition to the $K = 0$ subband there should be several other subbands with $K \neq 0$. In 1960 the non-observation of these subbands was considered as strong evidence that the molecule is linear. However in view of the new ESR evidence one must consider the possibility that the $K \neq 0$ subbands are so strongly predissociated that they escape observation [Herzberg and Johns (49)]. A decepta calculation of the bond angle from the observed $B_0$ values of CD$_2$ and CHD leads to 136° and a bond distance of 1.078 Å. We believe now that this is the correct structure for the ground state of CH$_2$.

In addition to the VUV spectrum a series of bands with complicated fine structure were found [(44), (50)] in the red and photographic infrared of which Fig. 18 gives an example. This spectrum was analyzed as that of an asymmetric top and corresponds to a second modification of CH$_2$ with zero spin (singlet CH$_2$). Its life time is shorter than that of the first (triplet)
Fig. 18.
Two subbands (1-0 and 1-2) of the 0 13 O-O 0 0 band of the red absorption system of CH₂.
modification suggesting that the latter represents the ground state. In Fig. 19 an energy level diagram of the observed states is shown. The energy difference of the lowest singlet and triplet state is uncertain. It should be noted that the observed low-lying states are precisely those predicted from the electron configuration \( ^3\Sigma_u^-, ^1\Delta_g, ^1\Sigma_g^- \) for the linear conformation.

In Fig. 20 the geometrical structure of \( \text{CH}_2 \) in the lowest states is illustrated graphically.

(3) Other dihydrides
Several years before the spectrum of \( \text{CH}_2 \) was observed Dressler and Ramsay (51) at Ottawa observed and analyzed the spectrum of the \( \text{NH}_2 \) radical obtained by the flash photolysis of \( \text{NH}_3 \). The spectrum is quite complicated since the molecule is an asymmetric top and its analysis by Dressler and Ramsay represented a very considerable accomplishment. The experience gained by them was of great help in the analysis of the singlet spectrum of \( \text{CH}_2 \).
to earlier. Fig. 21 shows the structure of NH₂ derived from the spectrum.

More recently Dr. Johns and I (52) obtained the spectrum of BH₂ by the flash photolysis of BH₃CO. This spectrum lies in the same region as that of NH₂. Fig. 22 shows a section of the spectrum showing the B¹⁰—B¹¹ isotope shift and Fig. 23 shows the resulting structure. Table 6 summarizes the information of observed bond angles in these and other dihydrides.

Several years before these structures were spectroscopically established Walsh (53) gave some rules for the prediction of such structures on the basis of certain semi-empirical assumptions about the molecular orbitals in these systems. In the Walsh diagram for XH₂ molecules shown in Fig. 24 the predicted energies of the lowest orbitals are plotted as a function of the bond angle (going from 90° to 180°). The orbitals designated 2a₁ and 1b₂ (or 2σg and 1a₁) favour slightly the linear conformation while the orbital 3a₁ which arises as one component of 1πu strongly favours the bent conformation. The other orbital 1b₁ arising from 1σu favours neither conformation. In the ground state of BH₂ the electron configuration is

$$(2a_1)^2 (1b_2)^2 3a_1$$

Fig. 21.
Geometrical structure of NH₂ in the two known electronic states.
Fig. 22.
Subbands of the BH$_2$ band near 8520 Å after Herzberg and Johns (52).
Lines of $^9$B H$_2$ are marked above, those of $^{11}$B H$_2$ below the spectra.
and therefore in this state the molecule is bent (see Table 6); but in the first excited state

\[(2a_1)^2 (b_2)^2 1b_1\]

there is no tendency for bending and in agreement with observation the molecule is predicted to be linear. In a similar way the other molecules of Table 6 can be treated: Two electrons in the 3a1 orbital always lead to strong bending with an angle of about 105° as in the ground state of H₂O.

(4) Triatomic monohydrides
A number of triatomic radicals with one H atom have been studied. The first was HCO of which Fig. 25 shows one of the absorption bands [Herzberg and Ramsay (54)]. Here again there is a striking change of shape in the electronic transition as shown in Fig. 26. Moreover this is another case (the first to be recognized) in which only one \(K\) value, here \(K'' = 1\), appears since in the upper state (where the molecule is linear) all levels with \(I \neq 0\) are strongly predissociated.

The spectra of HNO [Dalby (55)] and HCF [Merer and Travis (56)] shown in Figs. 27 and 28 are examples for cases in which the radicals are bent in both upper and lower state and where no predissociation occurs, i.e. several

![Fig. 23. Geometrical structure of BH₂ in the two known electronic states.](image)

Table 6. Bond angles in the ground states of triatomic dihydrides

<table>
<thead>
<tr>
<th>BH₂</th>
<th>CH₃</th>
<th>NH₃</th>
<th>H₂O</th>
</tr>
</thead>
<tbody>
<tr>
<td>131°</td>
<td>136° (triplet)</td>
<td>103.3°</td>
<td>105.2°</td>
</tr>
<tr>
<td></td>
<td>104° (singlet)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>AlH₃</th>
<th>SiH₂</th>
<th>PH₃</th>
<th>H₂S</th>
</tr>
</thead>
<tbody>
<tr>
<td>119°</td>
<td>92.1° (singlet)</td>
<td>91.5°</td>
<td>92.2°</td>
</tr>
</tbody>
</table>
The variation of the orbital energies in going from a bond angle of 90° to 180° is shown.

subbands of different K are observed. The geometrical structures resulting from the analysis of these spectra are shown in Figs. 29 and 30.

The most recent radicals studied in this group are HNF [Woodman (57)] and HSil [Billingsley (58)] for which Figs. 31 and 32 show the resulting structures. HNF is one of the first examples of an asymmetric top in which the spin doubling is fully resolved and analyzed.

(5) Triatomic non-hydrides

Among triatomic non-hydrides the C₃ radical has already been discussed. It is linear in both the upper and the lower state of the only known electronic transition which is of type \(^1Π_u^-1Σ_u^+\). In the excited state \(^1Π_u^-\) there is a strong interaction between electronic and vibrational angular momentum (Renner-Teller effect) leading to considerable anomalies in the spacing of the vibrational levels. In the ground state \(^1Σ_u^+\) the bending frequency is surprisingly small, only 63 cm\(^{-1}\). The corresponding force constant is 1/100 of that in CO, i.e. the molecule is very floppy. The electron configurations of the two states are \((1π_u)^4\) (30,) \(1π_π\) and \(\ldots \) \(1π_u\) \(4\) \(3σ_u\) \(2\).
Fig. 25.
The 0110—000 band of the A—X system of HCO at 5624 Å after Herzberg and Ramsay (54).

Only the 0—1 subband is observed. Each line is an unresolved doublet.
Fig. 26.
Geometrical structure of HCO in the two known electronic states.

Radicals with one more electron which has to go into a π_g orbital are CCN and CNC observed by Merer and Travis (59) at Ottawa. The NCN molecule with two electrons in the π_g orbital and therefore with a \(3\Sigma^-\) ground state, was studied by Travis and myself (60), the similar molecule CCO by Devillers and Ramsay (61). We have looked for the corresponding spectrum of the ion N_3^+ which has the same number of electrons. Ledbetter (62) found a complicated absorption band in flash discharges through nitrogen which is not otherwise identifiable, but no proof for its belonging to N_3^+ has yet been obtained. Several radicals and ions with three electrons in the π_g orbital have been studied by various investigators.

Table 7 summarizes the molecular constants for the ground states of this group of molecules. Note the strong increase of the bending frequency \(v_b\) as a function of the number of π_g electrons.

(6) CH_3 and CH_3^+

The spectrum of the methyl radical was easier to obtain than that of methylene (44). First by the flash photolysis of Hg(CH_3)_2, and later that of many other methyl containing compounds a spectrum consisting of two diffuse peaks near 2160 Å was obtained as shown in Fig. 33 a. For CD_3, a simple (even though still diffuse) fine structure is observed (Fig. 33 b) which yields the conclusion that CD_3 is planar (or very nearly planar) in its ground state and that the CD distance is 1.069 Å. In addition several Rydberg series have been observed in the region 1500-1280 Å which yield accurate values for the ionization potential \([\text{I.P.} (\text{CH}_3) = 9.843\text{eV}]\).

The electron configurations of the two lowest states of CH_3 assuming D_3h symmetry are
Fig. 27.
Two subbands of the 000—000 band of the $\tilde{A}-\tilde{A}$ system of the HNO radical after Dalby (55).
Subbands of the 000-000 band of the $\tilde{A}$$-\tilde{X}$ system of the HCF radical after Merer and Travis (56).

$$\ldots (1e')^4 (1a_2'')^2 A_2''$$
$$\ldots (1\epsilon')^3 (1a_2'')^2 E'$$

The transition between these two states should lie in the visible region but for planar CH$_3$ it is a forbidden transition and so far has not been observed. The upper states of all observed transitions are Rydberg states of the type

$$\ldots (1\epsilon')^4 nsa_1'; nda_1'; nd\epsilon''$$

The spectrum of SiH$_3$ is of course expected to be very similar to that of CH$_3$ but in spite of considerable effort we have not yet been able to observe

\[ \text{Fig. 28.} \]
Subbands of the 000-000 band of the $\tilde{A}$$-\tilde{X}$ system of the HCF radical after Merer and Travis (56).

\[ \text{Fig. 29.} \]
Geometrical structure of the HNO radical in the two known electronic states.
Fig. 30.
Geometrical structure of the HCF radical in the two known electronic states.

Fig. 31.
Geometrical structure of the HNF radical in the two known electronic states.

Table 7. Molecular constants in the ground states of linear triatomic non-hydrides

<table>
<thead>
<tr>
<th>Molecule</th>
<th>State</th>
<th>$\Delta G$ cm$^{-1}$</th>
<th>$B_0$ cm$^{-1}$</th>
<th>$r_0$ Å</th>
<th>$N$</th>
</tr>
</thead>
<tbody>
<tr>
<td>C$_2$</td>
<td>$^1\Sigma_g^+$</td>
<td>63.1</td>
<td>0.4305</td>
<td>1.277</td>
<td>12</td>
</tr>
<tr>
<td>CCN</td>
<td>$^2\Pi_r$</td>
<td>(325)</td>
<td>0.3981</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CNC</td>
<td>$^2\Pi_g$</td>
<td>321</td>
<td>0.4535</td>
<td>1.245</td>
<td>13</td>
</tr>
<tr>
<td>NCN</td>
<td>$^1\Sigma_g^-$</td>
<td>(423)</td>
<td>0.3968</td>
<td>1.232</td>
<td>14</td>
</tr>
<tr>
<td>CCO</td>
<td>$^2\Sigma_g^-$</td>
<td>379.4</td>
<td>0.3851</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NCO</td>
<td>$^2\Pi_t$</td>
<td>(539)</td>
<td>0.3894</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N$_2$</td>
<td>$^2\Pi_g$</td>
<td>0.4312</td>
<td>1.182</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td>BO$_4$</td>
<td>$^2\Pi_g$</td>
<td>464</td>
<td>0.3292</td>
<td>1.265</td>
<td>16</td>
</tr>
<tr>
<td>CO$_4$</td>
<td>$^1\Sigma_g^+$</td>
<td>667.4</td>
<td>0.3902</td>
<td>1.162</td>
<td></td>
</tr>
</tbody>
</table>

$N$ = number of valence electrons
Fig. 32. Geometrical structure of the HSiI radical in the two known electronic states.

Fig. 33. Absorption bands of (a) CH₃ and (b) CD₃ near 2150 Å after Herzberg (44).
it. Thus it is still an open question whether SiH₃ is planar or non-planar in its ground state although the latter appears more likely.

A molecular system with the same number of electrons as CH₄ is the CH₄⁺ ion. If CH₄⁺ were tetrahedral like CH₄ its ground state would have the electron configuration

\[(1a₁)² \ (2a₁)² \ (1f₂)⁶ \ ^2F₂\]

that is, would be a triply degenerate state. However, according to the Jahn-Teller theorem, in such a triply degenerate state the equilibrium conformation cannot be the symmetrical tetrahedral one. Recent theoretical calculations [see for example Dixon (63)] suggest, that CH⁺₄ has a D₂d structure and that the electron configurations of the two lowest electronic states derived from \(^2F₂\) are

\[(1a₁)² \ (2a₁)² \ (1e)⁴ \ (1b₂) \ ^2B₂\]
\[\ldots \ldots \ldots \ldots \ (1e)⁵ \ (1b₂)² \ ^2E\]

If, instead, CH⁺₄ had C₃ᵥ symmetry (i.e. one C-H bond longer than the other three) the states \(^2A₁\) and \(^2E\) with electron configurations even more similar to those of the two low-lying states of CH₃ would result. But now the transition between these two low-lying states is not forbidden. That these two low-lying states of CH⁺₄ exist has been conclusively shown by photo-electron spectroscopy. Fig. 34 shows the photo-electron spectrum as observed by Price (64). The two maxima at the right correspond to photoelectrons with CH⁺₄ left in one or the other of the two states. The allowed transition between these two states would lie in the visible region but has not yet been observed.

Fig. 34. Photo-electron spectrum of CH₄ obtained with the He⁺ line at 304 Å by Price (64).
One reason (in addition to its intrinsic interest) why we have spent considerable effort to observe this spectrum is it possible astrophysical importance. It is possible that the diffuse interstellar lines (which represent the last major identification problem in astronomical spectroscopy) may be due to this transition in CH⁺, the diffuseness being due to predissociation of the upper state (the dissociation energy of CH⁺ is only about 1 eV). Since CH₄ is almost certainly present in the interstellar medium it would not be surprising if a sufficient stationary concentration of CH⁺ arises for the appearance of the 'E→B₁ transition.

(7) HNCN
As a final example I should like to mention a radical which, like many others, was observed in the flash photolysis of diazomethane, namely HNCN [Herzberg and Warsop (65)]. As shown in Fig. 35 a particularly simple spectrum arises, a textbook example of a perpendicular band of a nearly symmetric top, consisting of a series of almost equidistant subbands. The large separation of the subbands indicates that the heavy atoms lie very nearly on a straight line while the H atom lies off that line. Isotope effects with D, C⁰ show that only one hydrogen and one carbon atom is present. Fig. 36 shows the derived structure; the position of the C atom cannot be accurately determined from the present data.

D. CONCLUSION
In the preceding discussion I have not described any of the techniques used in our work since the principal method, the flash-photolysis technique, was described by Norrish and Porter in their Nobel lectures (43). We have more recently developed two methods for the study of spectra of molecular ions: the flash discharge technique which is closely related to the flash photolysis technique and the flash radiolysis (or pulse radiolysis) technique in which a powerful pulse of electrons is sent through the absorption cell and produces many ions. Both methods have been described in more detail in my Faraday lecture (66). The results obtained with the second method have as yet been very sparse but we hope to overcome some of the difficulties and obtain absorption spectra of ions such as C₆H₆⁺, HCN⁺, CH⁺, CH⁺, CH⁺, H₂O⁺H₂O⁺, ... C₆H₆⁺ and many others about which nothing is known at present. The
information so obtained would be of immense help for a deeper understanding of the electronic structure of these ions as well as the corresponding neutral molecules. The new methods of tunable lasers as shown by Patel and his collaborators (67) promise to give infrared spectra of radicals and ions about which up to now very little is known. For example the vibrational frequencies in the ground state of CH$_2$ are unknown at present. These few remarks may suffice to indicate that a great deal remains to be done in the study of the geometric and electronic structure of radicals and molecular ions.

Acknowledgement

The work described in this lecture could not have been accomplished without the effective help and independent work of many collaborators. The list of references gives the names of most of them. I should like to single out particularly the constant support by most valuable criticisms and suggestions which I have had from Dr. A. E. Douglas, now my successor as director of the Physics Division of the National Research Council. I should also like to acknowledge the devoted help I have had for twenty years from Mr. J. Shoo-smith who carried out all the experimental work with great efficiency, energy and skill.

In receiving the Nobel Prize I am deeply aware of the debt I owe to all these collaborators and technical assistants for their continued help and support.
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