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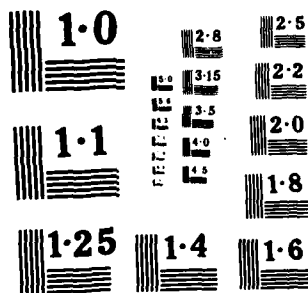
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The inner-valence (20-30eV) photoelectron spectrum of N_2 is interpreted by comparing with various spectra (such as absorption, I_p yield, fluorescence-yield, and core-level photoelectron spectra) and with previously published theoretical calculations. This empirical interpretation allows for unique assignments, and enables us to distinguish between the four previously published theoretical calculations which lead to inconsistent assignments.			

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12

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An Interpretation of the N_2 photoelectron Spectrum**

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Abstract

The inner-valence (20-36eV) photoelectron spectrum of N_2 is interpreted by comparing with various spectra (such as absorption, N^+ -yield, fluorescence-yield, and core-level photoelectron spectra) and with theoretical calculations. The bands at 25.3, 29.0, 32.6, 33.3 (sharp and weak), and 35eV (weak) are attributed to the ${}^2E_u^+(3\sigma_g^{-1}1\pi_u^{-1}1\pi_g)$, ${}^2E_g^+(2\sigma_u^{-1}1\pi_u^{-1}1\pi_g)$, ${}^2E_u^+(3\sigma_g^{-1}1\pi_u^{-1}1\pi_g)$, ${}^2A_u(3\sigma_g^{-1}2\sigma_u^{-1}1\pi_g)$, and ${}^2E_g^+(3\sigma_g^{-1}1\pi_u^{-1}1\pi_g^2)$ states, respectively.



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1. Introduction

The photoelectron spectrum (PES) by Nyholm *et al.* [1], the lower spectrum in Figure 1, clearly shows four bands around 25eV(D), 29eV(F), 33eV(E), and 38eV(H), and probably a weak band around 35eV(G). The recent measurement by Krummacher *et al.* [2], the upper spectrum in Figure 1, confirms the presence of these bands. H band, the strongest and broadest band among the five bands, is known to consist of several $^2\Sigma_g^+$ states that borrow their intensities from $2\sigma_g^{-1}$ configuration. In this letter, we assign symmetries and dominant electronic configurations to the D, F, E, and G bands.

First, we obtain symmetries and dominant configurations from experiments by comparing the PES with other spectra, such as the absorption spectrum, the N₂ vuv spectrum, the core-level PES, and a valence-level PES of a different photon energy. Then, we compare the obtained symmetries and configurations with published theoretical calculations.

2. Observed Rydberg states and the inversion symmetry of its ion-core state.

There is a close relationship between an observed (that is, dipole-allowed) Rydberg state and the inversion symmetry of its ion-core state. Rydberg states that are excited from the N₂ ground state, $^1\Sigma_g^+$, by dipole transitions should have "u" symmetry because the dipole operator has "u" symmetry.

If a Rydberg state I -ns or I -nd is observed, its ion-core state I should have "u" symmetry because Rydberg orbital ns or nd has "g" symmetry. If a Rydberg state I -np is observed, its ion-core state I should have "g" symmetry because Rydberg orbital np has "u" symmetry.

Figure 2 shows how we identify the C-3d, F-3p, F-4p, and E-3s Rydberg states in the absorption (ABS) and N⁺-yield spectra. The solid bars in the middle of the figure denote the Rydberg-electron binding energies that are estimated from the Rydberg states converging to the one-hole (1h) states. Note that the band widths of the C and C-3d states are similar and that those of the F and F-3p states are also similar. Paired vibrational progressions are identified for the C-3d state. Details of this identification will be published elsewhere.

The observation of the C-3d, F-3p (or F-4p), and E-3s Rydberg states implies that the C, F, and E ionized states have "u", "g", and "u" symmetries, respectively.

3. Photon-energy variation of PES

Ionization into two-hole, one-electron (2h-1e) states or three-hole, two-electron (3h-2e) states is forbidden, because it involves a two- or three-electron-jump transition. 2h-1e or 3h-2e states gain their intensity through configuration-interaction (CI) mixing with 1h states. In a configuration interaction, only the states

that have the same symmetry can mix with each other. Therefore, the bands that are clearly observed in PES (such as the C, F, E, and G bands) should have either $^2\Sigma_g^+$ (the symmetry of the $2\sigma_g^{-1}$ and $3\sigma_g^{-1}$ states), $^2\Sigma_u^+$ (the symmetry of the $2\sigma_u^{-1}$ state), or $^2\Pi_u$ (the symmetry of the $1\pi_u^{-1}$ state).

Figure 1 compares PES's of different photon energies. When the photon energy is decreased from 1487eV to 50.3eV, the $3\sigma_g^{-1}$ and $1\pi_u^{-1}$ intensities increase relative to the $2\sigma_u^{-1}$ intensity and also the F intensity increases relative to the C, E, G, and H intensities. These relative increases are depicted by shaded areas in the figure. The simultaneous increases of the $3\sigma_g^{-1}$, $1\pi_u^{-1}$, and F intensities suggest that the F state mixes with either the $3\sigma_g^{-1}$ or $1\pi_u^{-1}$ state and also that the C, E, and G states do not mix with these states (that is, the C, E, and G states mix with either the $2\sigma_g^{-1}$ or $2\sigma_u^{-1}$ state).

In the preceding section, we have concluded that the F state should have "g" symmetry. Combining this and the above conclusion that the F state must mix with either the $3\sigma_g^{-1}$ or $1\pi_u^{-1}$ state, we conclude that the F state mixes with the $3\sigma_g^{-1}$ and $2\sigma_g^{-1}$ states. Since the C and E states should have "u" symmetry and must mix with either the $2\sigma_g^{-1}$ or $2\sigma_u^{-1}$ states, we conclude that the C and E states mix with the $2\sigma_u^{-1}$ state. In summary, the F state has the $^2\Sigma_g^+$

symmetry; the C and E states have the 2E_u symmetry; and the G state has the 2E_g symmetry.

4. The ${}^2E_u(n\sigma^{-1}\pi_u^{-1}\pi_g)$ shake-up states

The $n\sigma^{-1}\pi_u^{-1}\pi_g$ configuration gives two 2E_u states. We expect that their energies relative to the ${}^2E_u(n\sigma^{-1})$ state (that is, the ${}^2E_u(n\sigma^{-1}\pi_u^{-1}\pi_g) - {}^2E_u(n\sigma^{-1})$ energies) are approximately independent of the $n\sigma^{-1}$ hole. Here, the $n\sigma^{-1}$ hole may be $1\sigma^{-1}$, $2\sigma_u^{-1}$, or $3\sigma_g^{-1}$. The following analysis is based on this assumption.

The ${}^2E_u(1\sigma^{-1}\pi_u^{-1}\pi_g)$ states have been identified as the lowest two shake-up peaks in the core-level PES (the middle and bottom spectra of Figure 3) [13]. The second-lowest, shake-up peak is much stronger than the lowest shake-up peak, as seen in Figure 3, indicating that the higher ${}^2E_u(1\sigma^{-1}\pi_u^{-1}\pi_g)$ state mixes more strongly with the ${}^2E_u(1\sigma^{-1})$ state.

When the $1\sigma^{-1}$ peak of the core-level PES (the middle) is aligned with the $3\sigma_g^{-1}$ peak of the valence-level PES (the top), the ${}^2E_u(1\sigma^{-1}\pi_u^{-1}\pi_g)$ shake-up peaks nearly align with the C and E bands, as shown in the figure. This agreement suggests that the C and E bands are the ${}^2E_u(3\sigma_g^{-1}\pi_u^{-1}\pi_g)$ shake-up states from the $3\sigma_g^{-1}$ hole. In the preceding section, we have concluded that the C and E bands should have the 2E_u symmetry. This conclusion confirms the above

statement. In summary, the C and E bands should have the ${}^2E_u^+$ symmetry and the $3\sigma_g^{-1}1\pi_u^{-1}1\pi_g$ configuration.

When the $1\sigma^{-1}$ peak of the core-level PES (the bottom) is aligned with the $2\sigma_u^{-1}$ peak of the valence-level PES (the top), the ${}^2E^+(1\sigma^{-1}1\pi_u^{-1}1\pi_g)$ shake-up peaks nearly align with the F and G bands (see Figure 3). This agreement would suggest that the F and G bands are the ${}^2E_g^+(2\sigma_u^{-1}1\pi_u^{-1}1\pi_g)$ states of the $2\sigma_u^{-1}$ hole. However, the core-level PES also suggests that the G band should be stronger than the F band, contrary to Figure 3. We attribute the H' band, which is indicated in Figure 1, to the higher ${}^2E_g^+(2\sigma_u^{-1}1\pi_u^{-1}1\pi_g)$ state. This assignment will be discussed elsewhere. In the preceding section, we concluded that the F band should have the ${}^2E_g^+$ symmetry. This conclusion confirms the ${}^2E_g^+(2\sigma_u^{-1}1\pi_u^{-1}1\pi_g)$ assignment of the F band.

G. Band widths of PES

Based on their bonding ($3\sigma_g$ and $1\pi_u$) and antibonding ($2\sigma_u$ and $1\pi_g$) characters, we expect that the FWHM's of the two ${}^2E_u^+(3\sigma_g^{-1}1\pi_u^{-1}1\pi_g)$ states are similar to each other and are larger than the FWHM's of the ${}^2E_g^+(2\sigma_u^{-1}1\pi_u^{-1}1\pi_g)$ states. Schirmer et al. [4] calculated the FWHM's for these states and confirmed the above expectation: 2.06eV and 2.12eV for the ${}^2E_u^+(3\sigma_g^{-1}1\pi_u^{-1}1\pi_g)$ states and 1.24eV for the lower ${}^2E_g^+(2\sigma_u^{-1}1\pi_u^{-1}1\pi_g)$ state.

The observed FWHM's (≈ 2.2 eV) of the C and E bands are similar to each other and are larger than the observed FWHM (≈ 1.6 eV) of the F band. These observed band widths support the C ${}^2\Sigma_u^+(3\sigma_g^{-1}1\pi_u^{-1}1\pi_g)$, E ${}^2\Sigma_u^+(3\sigma_g^{-1}1\pi_u^{-1}1\pi_g)$, and F ${}^2\Sigma_g^+(2\sigma_u^{-1}1\pi_u^{-1}1\pi_g)$ assignments. The C ${}^2\Sigma_u^+(3\sigma_g^{-1}1\pi_u^{-1}1\pi_g)$ assignment has been well established through studies on the C $\rightarrow$ X transition [5,6].

6. Theoretical calculations

Figure 4 compares the experimental results (the first column) with published theoretical calculations (the remaining columns). The shaded bars depict the band-peak energies from HFS. The symmetries and the dominant configurations in the first column are those determined in the previous sections. The (2/3) indicates the presence of both $2\sigma_g^{-1}$ and $3\sigma_g^{-1}$ mixing, and the (1/3) indicates the absence of the $3\sigma_g^{-1}$ mixing.

Theoretical calculations shown are the lowest six states that have either ${}^2\Sigma_g^+$, ${}^2\Pi_u$, or ${}^2\Sigma_u^+$ symmetry, excluding the X ${}^2\Sigma_g^+(3\sigma_g^{-1})$, A ${}^2\Pi_u(1\pi_u^{-1})$ and B ${}^2\Sigma_u^+(2\sigma_u^{-1})$ states. Solid bars show the calculated vertical ionization potentials, and the numbers above or below the bars give the one-hole mixing intensities. The ${}^2\Sigma_g^+$ state mixes with two one-hole states, the $2\sigma_g^{-1}$ and $3\sigma_g^{-1}$ states; the upper number denotes the $2\sigma_g^{-1}$ mixing intensity. The symmetries and the dominant configurations in the last column are determined from calculations.

The computational methods used were the partitioning CI (Kosugi et al. [7]), the polarization CI (Langhoff et al. [8]), the multiconfigurational Green's function (Nichols et al. [9]), and the equation-of-motion Green's function (Herman et al. [10]). Kosugi's, Langhoff's and Nichols' calculations include the $1h$, $2h-1e$, and $3h-2e$ configurations, but Herman's calculation includes only the $1h$ and $2h-1e$ configurations. Calculations that are not shown in Figure 4, such as those by Schirmer et al. [4,11], and the single-excitation CI and 2ph-TDA Green's function calculations in Ref. [8], also include only the $1h$ and $2h-1e$ configurations and give results similar to Herman's calculation. It has been demonstrated [7,8] that the inclusion of the $3h-2e$ configurations is essential to provide accuracy for the higher states above 32eV.

For the C and F bands, the theoretical calculations almost reproduce the experimental results for the band energies, the band intensities, the symmetries, and the dominant electronic configurations. The theoretical calculations except Nichols' calculation agree on the amount of $3d_{5/2}$ mixing in the F state. The calculated intensity for the ${}^2P_{1/2}$ state around 28eV is too weak to be observed.

For the E band, Kosugi's and Langhoff's calculations agree with the experimental results for the band energies, the band intensities, the symmetries, and the dominant electronic configuration. Note that the calculated

intensity for the $^2P_{1/2}$ state around 33eV is too weak to assign to the E band.

For the G band, the theoretical calculations agree with the experimental results for the band energy, band intensity, the symmetry, and the near absence of $3d_{5/2}^{-1}$ mixing. The theoretical dominant configuration $3d_{5/2}^{-1}1\pi_{u/2}^{-1}2\pi_{g/2}^2$ is also consistent with the experimental evidence that $2d_{5/2}^{-1}1\pi_{u/2}^{-1}1\pi_{g/2}^2$ is not a dominant configuration. Based on this agreement, we assign the $^2P_{1/2}(3d_{5/2}^{-1}1\pi_{u/2}^{-1}2\pi_{g/2}^2)$ state to the G band.

The above comparisons suggest that Kosugi's and Langhoff's calculations are reliable. These two calculations predict a low-intensity $^2P_{1/2}(3d_{5/2}^{-1}2d_{5/2}^{-1}1\pi_{g/2})$ band around 33eV. This band is expected to be sharper than the F band (broadening from its configuration) and to show the photon-energy variation similar to that of the F band (see Fig. 1). In Figure 2 of Reference [2], we can recognize a sharp low-intensity band at 33.3eV, which seems to show the expected photon-energy variation. Furthermore, a 31.5eV peak in the fluorescence spectrum [12] can be attributed to the 3d or 4s Rydberg state converging to this ionized state at 33.3eV. Based on these evidence, we identify the sharp low-intensity band at 33.3eV in the PES as the $^2P_{1/2}(3d_{5/2}^{-1}2d_{5/2}^{-1}1\pi_{g/2})$ state.

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Figure Captions

- Fig. 1. Comparison of photoelectron spectra taken at two different incident photon energies. The shaded areas indicate the relative increase in the band intensities at the 50.3eV photon energy in comparison with those at the 1487eV photon energy.
- Fig. 2. Identification of the Rydberg states converging to the L, F, and E ionized states.
- Fig. 3. The $2\Sigma^+(n\sigma^{-1}\pi_u^{-1}\pi_g)$ shake-up states observed in the valence-level and core-level PES.
- Fig. 4. Comparison of the experimental results with published theoretical calculations.

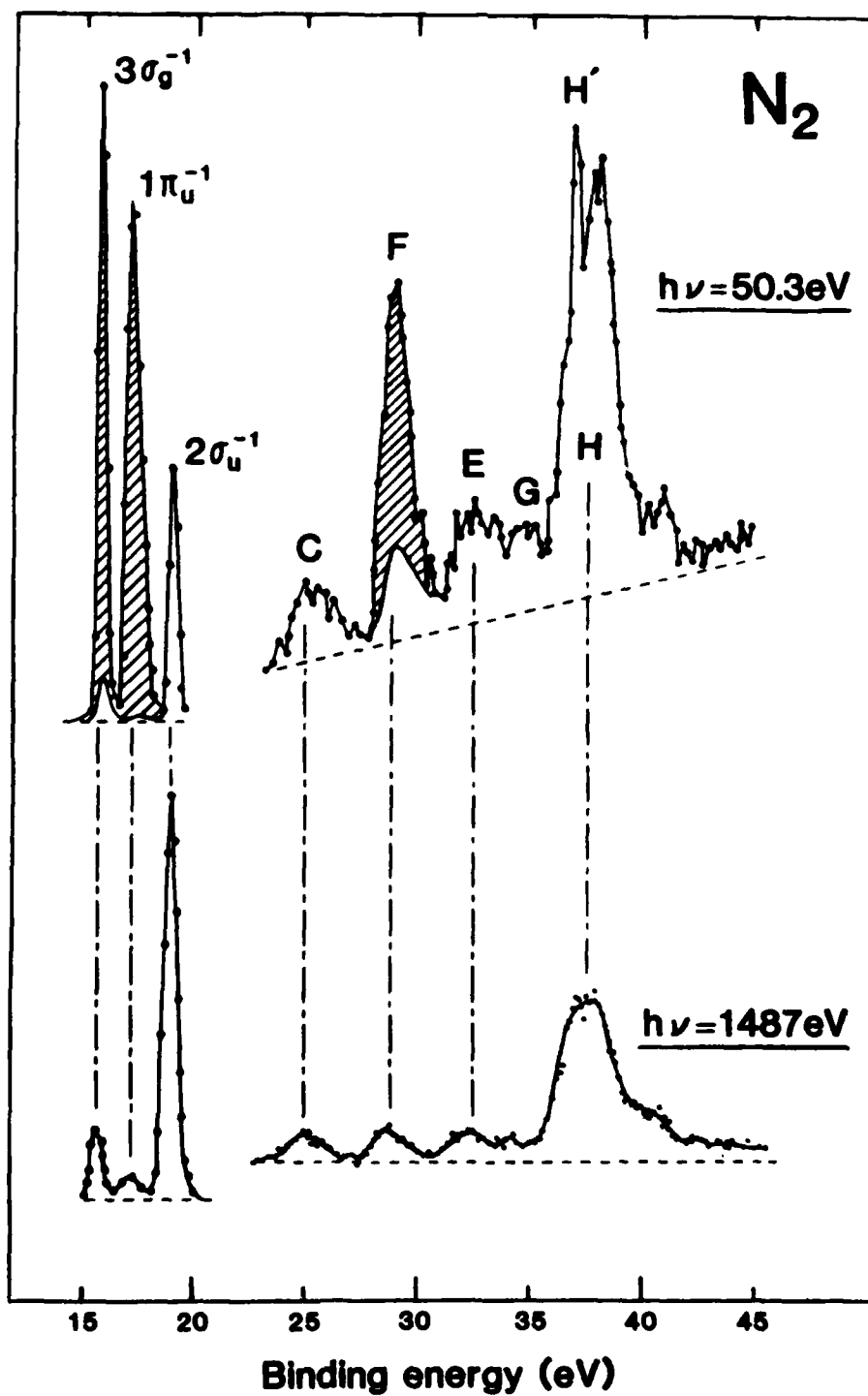


Fig. 1

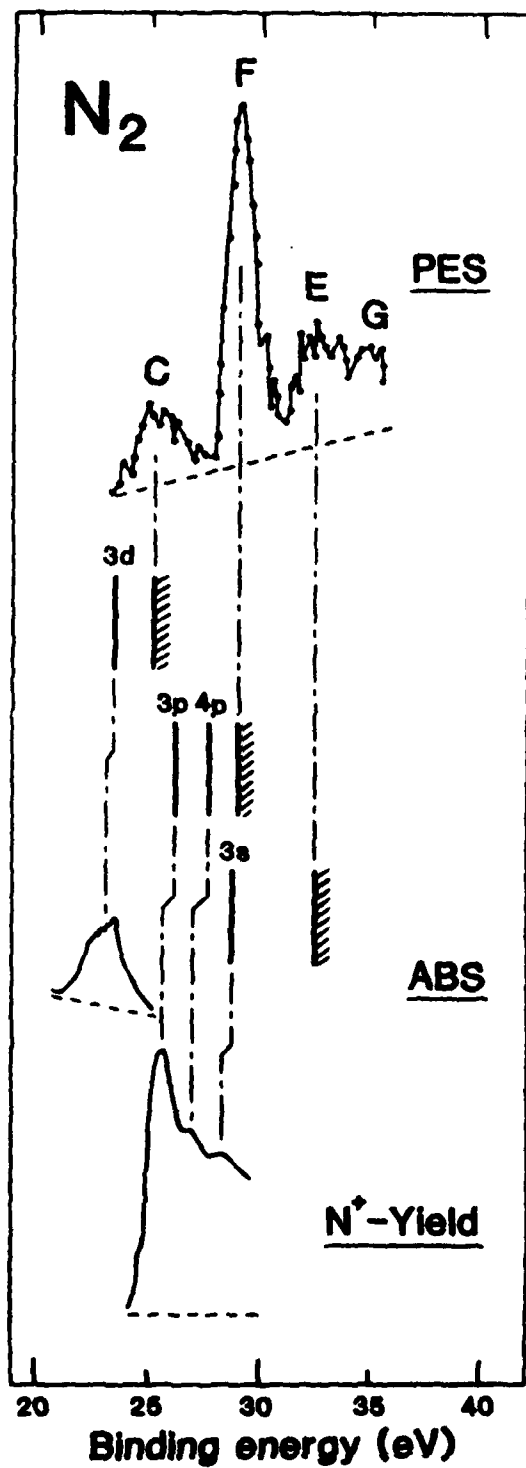


Fig. 2

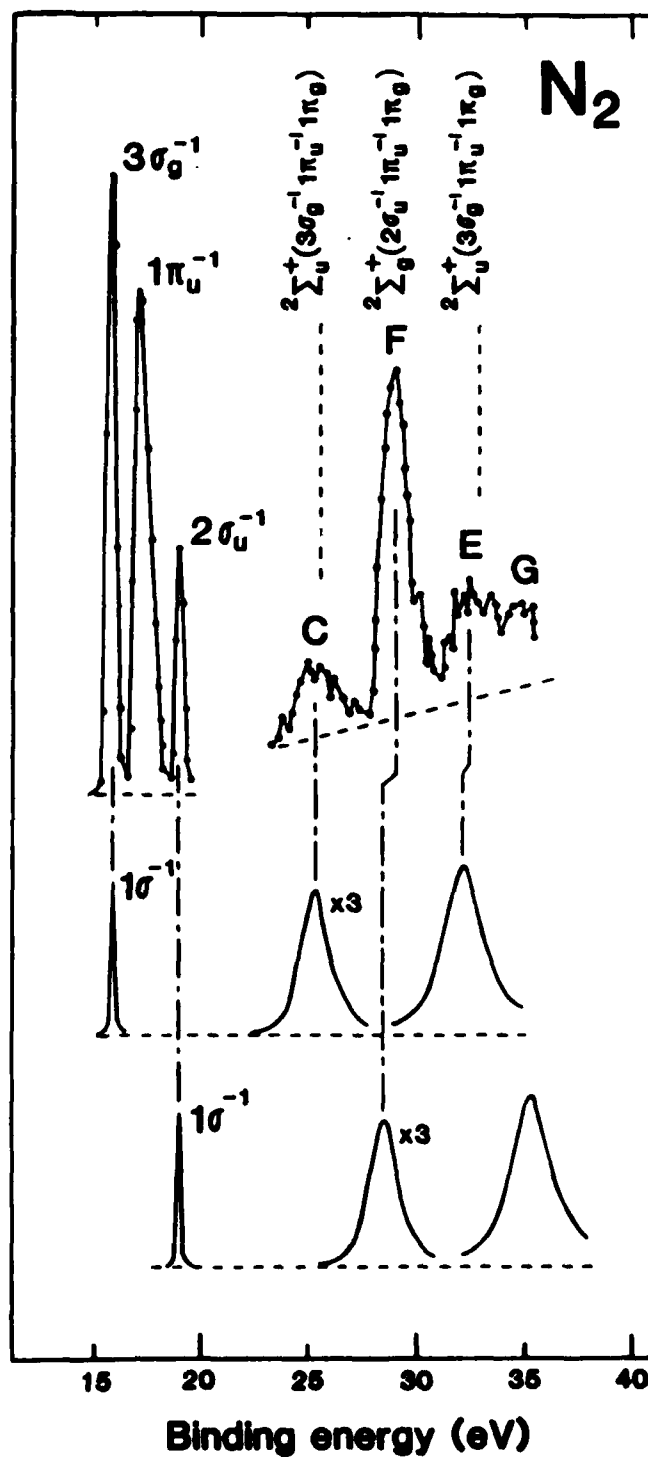


Fig. 3

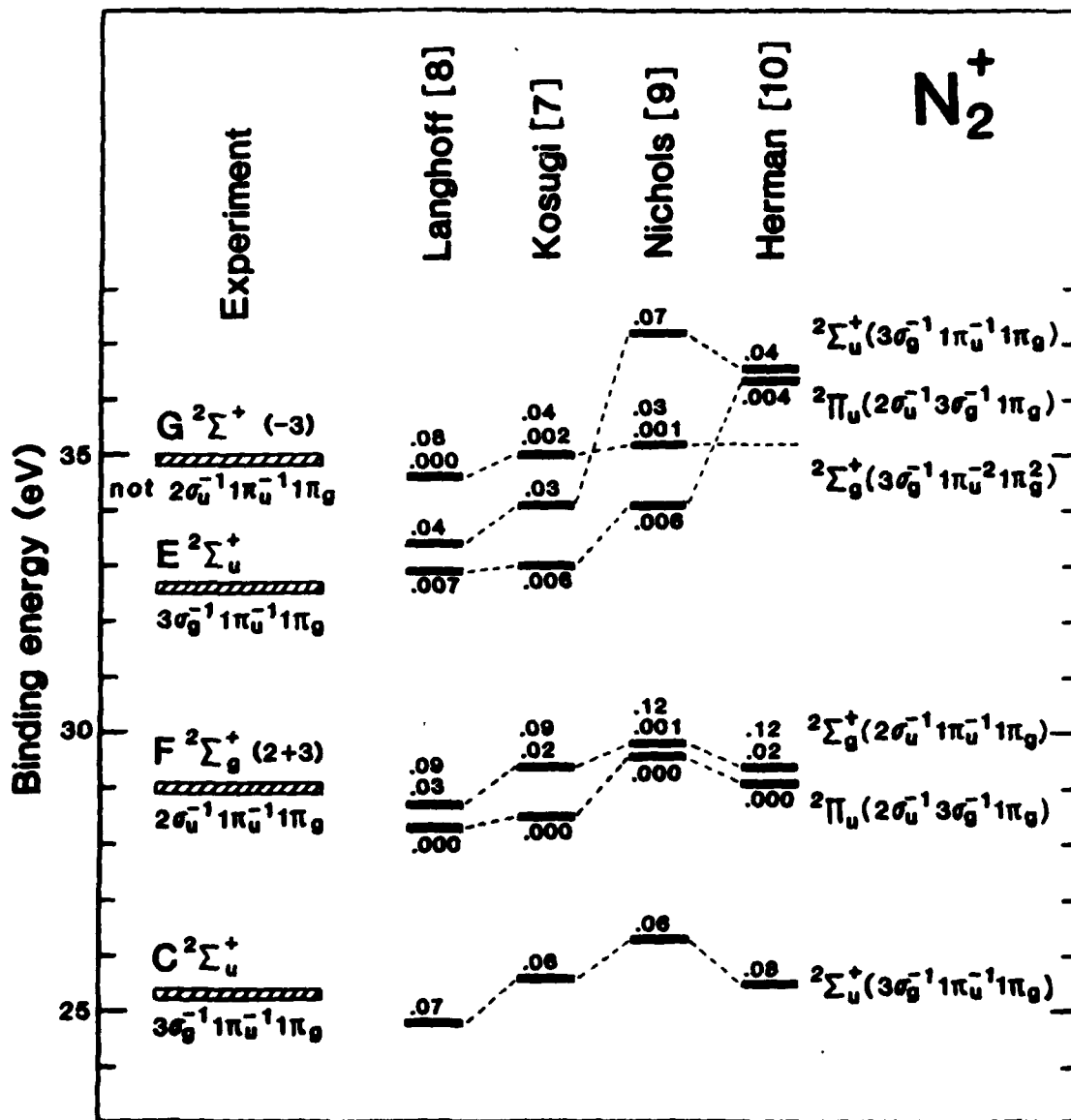


Fig. 4

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