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WILTHS OF ATOMIC 42 AND (NASA-TE-X-72900) 4p VACANCY STATES, 46 LESS THAN OF EQUAL TO Z LESS THAN OF EQUAL TO 50 (NASA) 30 p HC Unclas CSCL 20L \$4.00 01934 G3/76

WIDTHS OF ATOMIC 4s and 4p VACANCY STATES, 46 < 2 < 50

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Abstract

X-ray photoelectron and Auger spectra involving N_1 , N_2 , and N2 vacancy states of Pd, Ag, Cd, In, and Sn were measured and compared with results of free-atom calculations. As previously observed in Cu and Zn Auger spectra that involve 3d-band electrons, we now also find free-atom characteristics, with regard to widths and structure, in the Ag and Cd $M_4 - N_{4,5}N_{4,5}$ and $M_5 - N_{4.5}N_{4.5}$ Auger spectra that arise from transitions of 4d-band electrons. Theoretical N₁ widths computed with calculated free-atom Auger energies agree well with measurements. Theory however predicts wider N₂ than N₃ vacancy states (as observed for Xe), while the measured N_2 and N_3 widths are nearly equal to each other and to the average of the calculated N_2 and N_3 widths. The calculations are made difficult by the exceedingly short lifetime of some 4p vacancies and by the extreme sensitivity of super-Coster-Kronig rates, which dominate the deexcitation, to the transition energy and to the fine details of the atomic potential.

I. INTRODUCTION

The lifetimes of atomic inner-shell vacancies are uniquely related, through the uncertainty principle, to the widths of the corresponding atomic energy levels. In many cases, atomic level widths can be measured quite accurately by x-ray photoelectron spectroscopy,^{1,2} providing an *i*-portant check on calculations of transition probabilities and lending insight into deexcitation processes.³ In particular, considerable difficulties are still encountered in theoretical estimates of Coster-Kronig rates, 4-6 which often determine the dominant partial widths of states that are characterized by vacancies in the lower L, M, and N subshells⁷; more experimental data are needed as a guide for refined calculations. Available information on atomic level widths is quite incomplete as yet, in spite of its relevance to fundamental theory and applications.⁸ In the present paper, we report on measurements of N_1 , N_2 , and N_3 widths in Pd, Ag, Cd, In, and Sn, and compare these with new calculations.

II. EXPERIMENT

A. Measurements

The x-ray photoelectron spectrometer used in these experiments has been described previously.¹ Photoelectrons were produced by (nonmonochromatized) Mg K $\alpha_{1,2}$ x rays and retarded to ~100 eV before they entered a ll-cm radius hemispherical electrostatic analyzer. Samples consisted of spectroscopically pure foils. Sample surfaces were sputter-cleaned with Ar ions until the 1s photoelectron lines of O and C became nondetectable or, at most, barely discernible above background. Spectra were measured at pressures of $\leq x10^{-8}$ Torr. All measurements were repeated with three different samples of each element.

Lines with favorable signal-to-background ratio were measured at a resolution characterized by 1.1-eV full width at half-maximum (fwhm) of the Au N₇ photoelectron peak. Many N-shell photoelectron lines measured in this work are quite wide (>4 eV), however, and rather weak; in these cases, the spectrometer resolution was reduced to 1.5-eV fwhm for the Au N₇ line, thus enhancing the signal-to-background ratio. Even so, instrumental broadening remained negligible in comparison with the intrinsically large experimental uncertainties associated with such large widths (Sec. IIB).

B. Data analysis and results

As in our previous work on M-level widths,¹ the photoelectron spectra were smoothed with a spline-fit computer program. The background on both sides of a peak was fitted with a single fourth-degree polynomial which extends under the peak. Results were drawn with a CalComp-780 plotter. In the 46 \ll 50 region, the N₂ and N₃ photoelectron lines overlap considerably. Their background-substracted smooth spectra were deconvoluted with a DuPont 310 curve resolver, subject to the additional criterion that the N₂/N₃ photoelectron intensity ratio be 1/2. The single-component standard in the curve-resolving process was a Lorentzian shape convoluted with the Gaussian instrumental contribution, according to the procedure of Wilkinson.⁹ The combined width of the vacancy state and the incident x-ray is at least six times the instrumental width in all cases; hence the convoluted shape remains practically Lorentzian.

The application of these procedures to the $N_{2,3}$ photoelectron spectrum of Ag is illustrated in Fig. 1. The original data, smoothing and computer-fitted background are shown in Fig. 1(a). It is apparent how extensively the wide N_2 and N_3 lines overlap; this makes them difficult to resolve. Moreover, the shape of the spectrum is such that background sub-traction involves some degree of subjectivity. The deconvolution of the background-subtracted spectrum, based on Lorentzian line shapes [Fig. 1(b)], is more successful in this case than in some others with poorer signal-to-background ratios and broader peaks. These difficulties, due more to the nature of the photoelectron spectra than to the instrumental resolution, account for the rather large uncertainties attached to some of the measured widths.

Results of the measurements are listed in Table I, with theoretical predictions due to $McGuire^{10}$ and from our present calculations. The experimental vacancy-state widths were derived on the assumptions that the width of the Mg Ka_{1,2} x-ray line is 0.8 eV and that the width of the photoelectron line is the Lorentzian sum (simple addition) of the incident x-ray width and the width of the vacancy state. Instrumental broadening is neglected. We believe that it is justifiable to disregard the (Gaussian) instrumental contribution, even at degraded

resolution, because all uncorrected photoelectron lines are at least 3.5 eV wide and allowance for large experimental uncertainties is made in the assigned error limits.

The question remains to what extent non-lifetime broadening¹¹ contributes to the line widths derived from the x-ray photoelectron spectra. In view of the metallic character of the samples and the large widths (>3.5 eV) under consideration, it is expected that such non-lifetime mechanisms as charging, thermal and phonon broadening, and many-electron (conduction electron) excitations 12-14 do not contribute significantly. Plasma frequencies in the elements studied here correspond to energies 15 of >12 eV and their excitation intensities are generally much lower than those of the main photoelectron peaks, so that broadening due to this cause should be minimal. We have chosen to study only elements with filled d shells, so as to avoid significant broadening due to multiplet splitting and other related final-state effects arising from localized unpaired electrons. Although the precise nature of pronounced shakeup satellites in solids (~5-12 eV) is still somewhat ambiguous, it is an experimental fact that such satellites are observed only in metal compounds and not in the metals themselves.¹⁶⁻²⁵ Broadening contributions from shakeup or shakeoff satellites should therefore be insignificant in the pure metallic samples used in the present work.

III. DISCUSSION

A. Quasiatomic Auger Spectra

In a previous paper, we have discussed the quasiatchic character of the Auger spectra from solid Cu and Zn.¹ In particular, we showed that the $L_3-M_{4,5}M_{4,5}$ Auger transition, which involves <u>two</u> electrons from the $M_{4,5}$ (3d) band of solid Cu and Zn, does not exhibit band structure, but rather, contains fine structure similar to that in spectra of free atoms. In width and shape, these Auger spectra thus differ sharply from the x-ray photoelectron (XPS) spectra of the 3d band itself. The same fine structure has recently been observed in Auger spectra of Zn vapor which contains only free Zn atoms, ²⁶ as well as in solid germanium²⁷ and in gaseous GeH₄, ²⁸ where the 3d electrons are more core-like.

Two reasons exist for this quasiatomic behavior of Auger spectra in contrast to the solid-state character of soft x-ray emission spectra. On the one hand, in the presence of an inner-shell photohole, the more localized valence electrons are likely to be preferentially selected by the Coulomb operator to participate in the Auger process, while the x-ray emission dipole operator has relatively long-range character. On the other hand, the doubly-ionized final state of Auger transitions is more free-atom like than the singly-ionized final state of radiative transitions.^{29,30} Our original suggestion¹ concerning a lifetime difference between radiative and radiationless processes is in error, as pointed out by Mehlhorn.³⁰

The important physical implications of the quashtomic characteristics of Auger spectra from solids are as follows:

(i) Because Coster-Kronig and Auger transitions are caused by the same Coulomb operator and both lead to doubly-ionized final states, the quasiatomic behavior of Auger transitions in solids implies similar behavior of Coster-Kronig transitions.

(ii) In the outer shells, such as the M or N shells of medium-Z elements, the width of a vacancy state is almost entirely governed by Auger and Coster-Kronig transition rates. If these transitions are quasiatomic in solids, then the vacancy-state widths as deduced from XPS of solid samples should reflect the quasiatomic transition rates, provided that other non-lifetime contributions to the XPS widths are negligible.

(iii) Whereas the fine structure of Auger spectra and the width of photoelectron lines in solids may be quasiatomic in character, their kinetic energies, and hence the measured binding energies of electrons in various shells, are definitely not free-atom like, but are influenced by solid-state effects such as extra-atomic relaxation. ^{31,32}

These points are borne out by the fact that the large discrepancies between theoretically predicted M-vacancy widths³³ and experimentally measured values could be resolved by recalculating the widths using free-atom Auger and Coster-Kronig energies and neutralatom potentials.¹ We now inquire whether similar free-atom behavior of N Auger spectra exists in metals such as Ag and Cd. Figure 2 shows

the $M_5-N_4, 5N_4, 5$ and $M_4-N_4, 5N_4, 5$ Auger spectra of Ag and Cd and the respective photoelectron spectra of the $N_{4,5}$ (4d) band. The spectra of In are included for comparison because the In $N_{4,5}$ level is more core-like. As in the case of Cu and Zn, we note a similarity among the Auger features of Ag and Cd, even though the width and shape of the photoelectron spectra are quite different. The fine structure of these Auger spectra is not well resolved. However, the fine structure is definitely similar to that in the free-atom inner-shell M4.5-N4.5N4.5 Auger spectra of gaseous Xe. 34,35 This similarity was already noted by Aksela³⁶ under coarse resolution. More recently, Powell conducted a high-resolution study of Ag and specifically emphasized the lack of band structure, and hence, the quasiatomic character of these Auger transitions.²⁹ Additional evidence for the quasiatomic characteristics is provided by the recently obtained Auger spectrum of Cd vapor, 37 which exhibits similar features as those of Cd metal, although much better resolved. We can thus anticipate that in the 46 < < 50 range the N-shell widths should be essentially free-atom like. Hence, we compare the measured widths with free-atom calculations.

B. Comparison with Theory

1. Calculation of N-subshell widths for quasifree atoms

Free-atom Auger energies, which differ by the extraatomic relaxation energy from energies measured on solid samples,³² were calculated from first principles.¹ Rela-

tivistic Hartree-Fock-Slater wave functions were used, with Slater's Xa approximation to the exchange correlation term in the expression for the statistical total energy.³⁸⁻⁴⁰ The parameter a was chosen to be 0.7 throughout. The calculated Coster-Kronig energies are listed in Table II.

Auger and Coster-Kronig transition rates were computed with Herman-Skillman⁴¹ nonrelativistic Hartree-Slater wave functions. The Latter tail correction⁴² was included, and Xa exchange was used. Radiationless transition rates were calculated in j-j coupling in the standard manner.^{5,6} The rates with which we are here concerned are exceedingly sentitive to the atomic potential. This fact is illustrated by the dependence of N-level widths on the choice of exchange potential, as plotted in Fig. 3.

Radiative transitions were disregarded, the radiative partial width of N-shell vacancy states being negligible compared with the radiationless width.¹⁰

. Total widths computed in this manner are included in Table I.

Figure 4 shows that the measured N_1 widths in the 46 < Z < 50 region agree exceedingly well with the free-atom calculations. These widths are chiefly determined by the N_1 - N_2 , 3^N_4 , 5^{N_2} super-Coster-Kronig rates. Coster-Kronig transition rates are extremely energy-sensitive, particularly near threshold.¹ This fact is well illustrated by the difference between two sets of results for the N_1 widths obtained by McGuire,¹⁰ who estimated the continuum-electron energy in two different ways; these results differ by a factof of ~ 6 at Z=47 and by ~ 25 at Z=50. The higher values (not included in Fig. 4), which increase rather than decrease with Z, were found by estimating the super-Coster-Kronig electron energy ε through the formula

$$\mathbf{e} = \mathbf{E}_{n_{1}\ell_{1}}^{(2)-(1/2)} \left[\mathbf{E}_{n_{3}\ell_{3}}^{(2)} + \mathbf{E}_{n_{3}\ell_{3}}^{(2+1)+\mathbf{E}_{n_{4}\ell_{4}}^{(2)+\mathbf{E}_{n_{4}\ell_{4}}^{(2)+\mathbf{E}_{n_{4}\ell_{4}}^{(2+1)}} \right],$$
(1)

where $E_{n\ell}(2)$, the binding energy for the $n\ell$ subshell of the neutral atom with nuclear charge Ze, was taken from the ESCA table of binding energies.⁴³ The subscript 1 refers to the hole in the initial atom, while 3 and 4 refer to the final holes. The lower set of width: found by McGuire,¹⁰ on the other hand, which agrees well with ours (Fig. 4), was calculated with super-Coster-Kronig energies estimated according to the prescription of Asaad and Burhop⁴⁴ with a work-function correction to the ESCA binding energies.⁴³ Clearly, the latter method of arriving at Coster-Kronig energies is far more realistic.

2. N2,3-level widths

Except for $_{47}$ Ag, the $4p_{1/2}$ - $4p_{3/2}$ spin-orbit splitting had notheretofore been resolved in the 46 <Z <50 region. Even though the experimentally resolved N₂ and N₃ peaks are separated by 8 to 15 eV, their widths are comparable with their set tration,

causing substantial overlap (Fig. 1). Taking experimental uncertainties into account, however, we find that the measured N2 and N2 vacancy-state widths are nearly equal, for each element. Equal N2 and N2 widths are also obtained theoretically through McGuire's calculation, 10 in which the N₂ and N₃ levels are assigned the same (average) energy, i.e., spin-orbit energy splitting is neglected. In our own, free-atom calculations, on the other hand, we take account of the difference between N, and N_3 binding energies. The widths of the N_2 and N_3 levels in this region are primarily governed by the super-Coster-Kronig transitions N2-N4,5N4,5 and N3-N4,5N4,5, which are very energysensitive (Fig. 5). Different binding energies for N_2 and N_3 electrons therefore lead to different N₂ and N₃ vacancy-state widths. Somewhat surprisingly, McGuire's calculation¹⁰ which includes a kind of averaging by treating the N_2 and N_3 levels as degenerate, agrees well with our measurements (Figs. 6 and 7; Table I). Our calculation, on the other hand, leads to N2 widths that are too large and to N, widths that are too small, in general, compared with experiment. The average of our theoretical N₂ and N₃ widths does, however, agree with the (nearly equal) measured N, and N, widths. In view of the good agreement of our calculated N_1 widths with measurement, the discrepancy in the N, and N, widths is puzzling. We note that the large (59.5-eV) N₂ width that we calculate for Xe (Table I) is not contrary to observation: in ESCA measurements, it was

found that the N₂ level of Xe is far too broad for positive identification.³⁴

4. N_{4,5}-level widths

The $N_{4,5}$ levels form the 4d band of Ag and Cd and are not split in In or Sn (Fig. 2). No effort was therefore made to measure the individual widths of these levels for comparison with atomic calculations.

5. Concluding Remarks

The following difficulties are encountered in the calculation of theoretical N-level widths:

<u>1</u>. The super-Coster-Kronig transition rates which largely govern the widths are exceedingly energy-sensitive.

2. These radiationless transition rates are extraordinarily sensitive to the fine details of the atomic potential.

3. The very short lifetime of the $N_{2,3}$ hole states casts some doubt upon the basic validity of perturbation theory to calculate the decay of these states.

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	Ĩ	r (N ₁)		Ī	Γ (N ₂)	_	Ē	Γ (N ³)	
Element	Theory		Experiment	9u.	1	Experiment	JUT	1	
	Ø	٩		Ø	٩		ro	٩	
46 ^P d		5.25	5.0±0.5		7.79	6.5±1		6.32	6.5±1
47 ^{Ag}	41.9 (7.0)	4.13	4.2±0.5	9.69	10.55	8.5±1	9.69	5.44	7.5±1
48 ^{cd}		4.07	4.0±0.5		13.38	12±1.5		8.65	10±1.5
49 ^{In}		3.03	3.5±0.5		17.59	15±2		10.55	14±2
50 ^{Sn}	78.2 (3.1)	2.89	2.8±0.5	16.2	20.81	17±2	16.2	10.39	17±2
54 ^{Xe}		5.87			59.48			2.38	

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^a McGuire (Reference 10). Values in brackets are based on an alternative energy estimate for the N₁-N_{2,3}N_{4,5} super-Coster-Kronig transitions. See text, Sec. III B2.

b_{Present} free-atom calculations.

Element N2N45 46Pd 11.45 47A9 7.82 48Cd 4.60	N3N45			- ^t N	- ^T N	N1-	N2 ⁻	л ³⁻
11.45 7.82 4.60		N201	N2 ⁰ 23	N ₃ 01	N ₃ 0 ₂₃	N45N45	N45N45	N45 ^N 45
7.82 4.60	15.74					63.17	30.53	26.24
4.60	11.57	18.20		21.95		64.27	29.74	25.99
	10.60	18.21		24.21		66.16	29.38	23.38
	5.02	16.77	24.48	23.70	31.41	65.91	27.64	20.71
		15.01	23.48	22.84	31.31	63.10	24.27	16.47
		21.10	32.52	34.66	46.08	61.23	6.23	

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Figure Captions

Fig. 1. Mg Ka_{1,2} x-ray excited photoelectron spectrum of Ag. (a) Original spectrum with smoothing and computer-fitted background. (b) Deconvolution of the background-subtracted spectrum into Lorentzian shapes produced with a DuPont 310 curve resolver, subject to the criterion that the N_2/N_3 intensity ratio be 1/2. Units on the abscissa are 0.4 eV/channel.

Fig. 2. $M_4 - N_4, 5N_4, 5$ and $M_5 - N_4, 5N_4, 5$ Auger spectra (left) and Mg Ka x-ray excited $N_{4,5}$ photoelectron spectra (right) of Ag, Cd, and In. The small peak in the $N_{4,5}$ photoelectron spectra is caused by the Mg Ka_{3,4} satellite radiation from the x-ray source.

Fig. 3. Dependence of the N_1 - and N_3 -level widths on the choice of exchange used in the calculation, illustrating extreme sensitivity of the dominant radiationless transition rates to the atomic potential.

Fig. 4. Comparison of theoretical and experimental widths of N_1 vacancy states. Experimental data and free-atom calculations are from the present work; triangles represent the lower of two sets of results obtained by McGuire (Reference 10) under different energy assumptions (see text, Sec. III B 2).

Fig. 5 $N_2 - N_4 N_5$ and $N_3 - N_4 N_5$ partial widths of Sn as functions of Auger electron energy, illustrating steep energy dependence of these super-Coster-Kronig transition rates.

Fig. 6. Comparison of measured N_2 -level widths with theoretical results from the present work and of McGuire (Reference 10).

Fig. 7. Comparison of measured N_3 -level widths with theoretical results from this work and of McGuire (Reference 10)

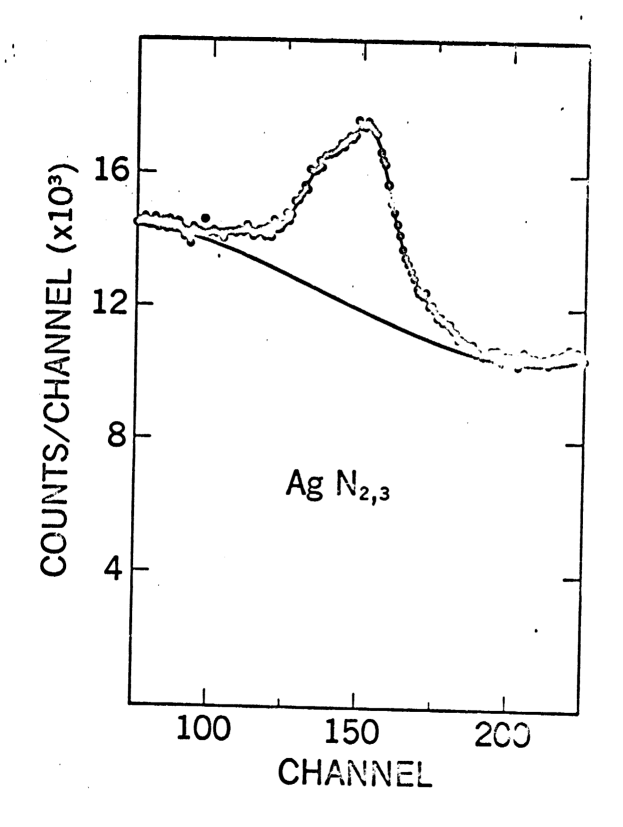


Fig. 1A

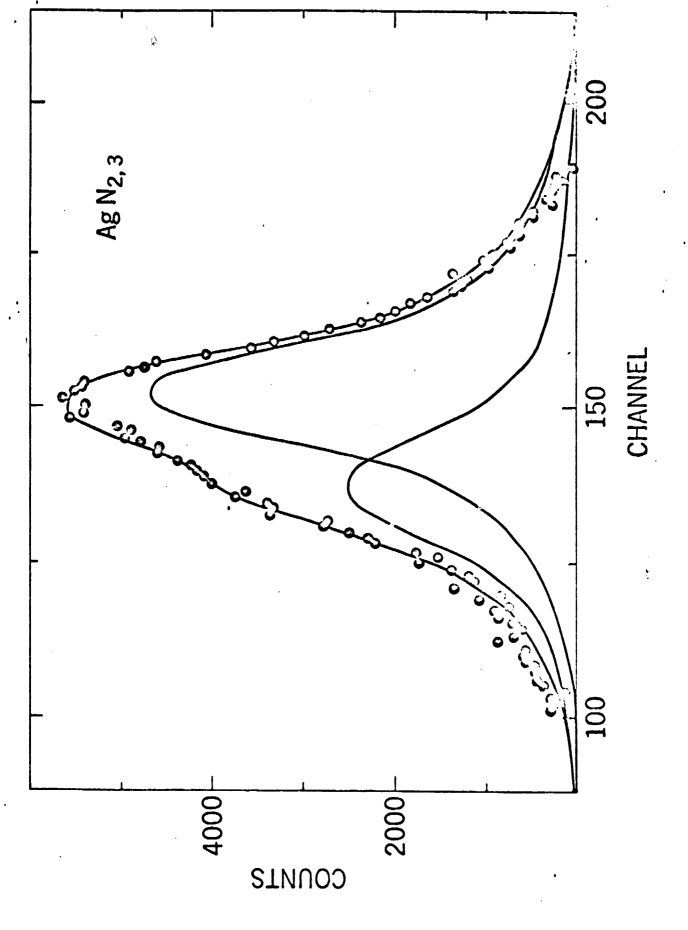
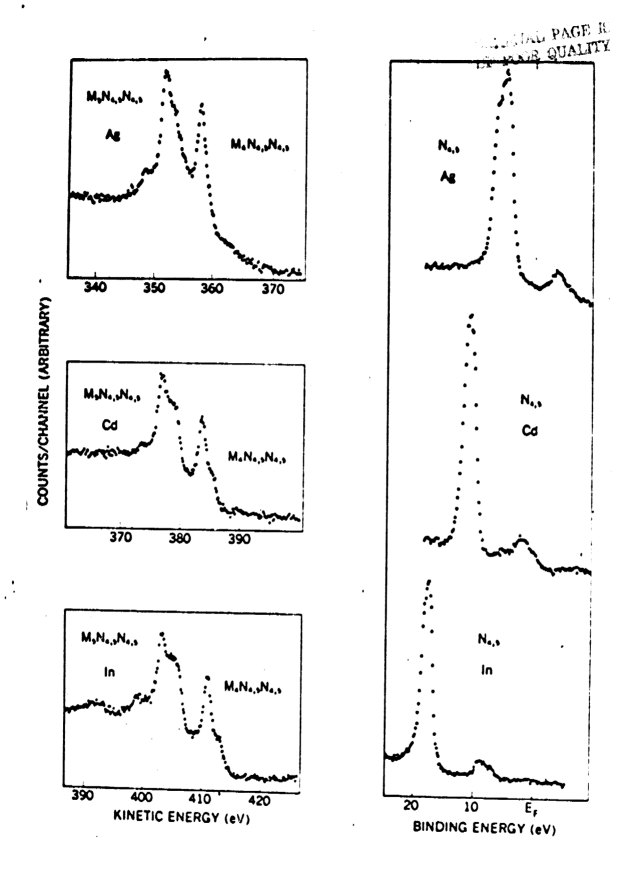
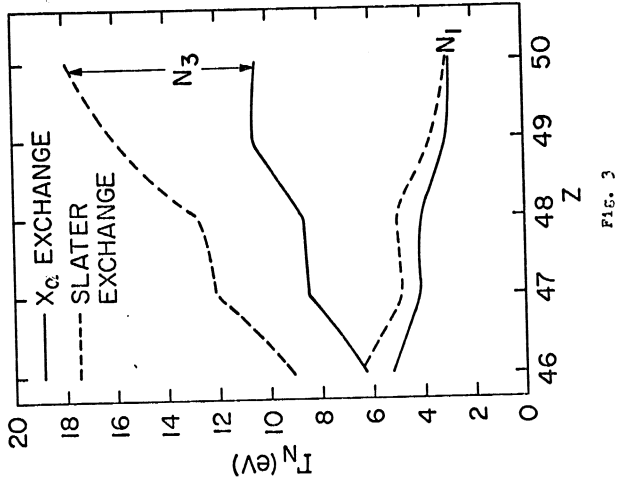


Fig. 1 (b)

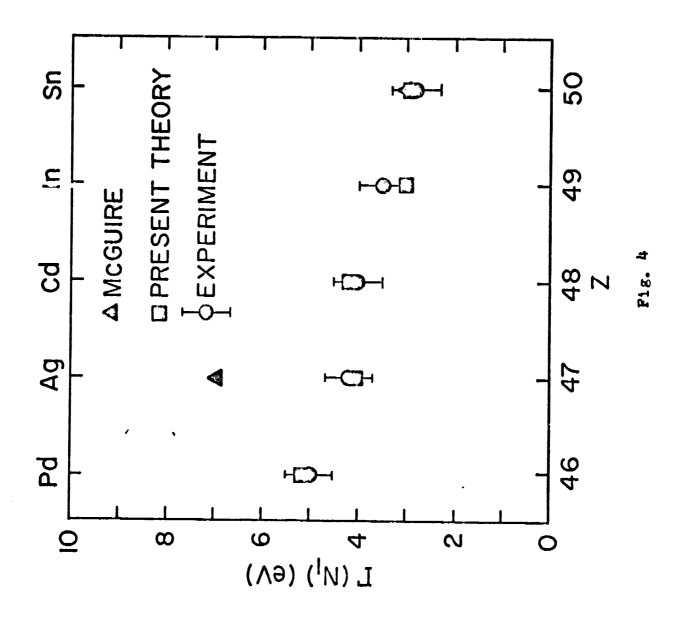


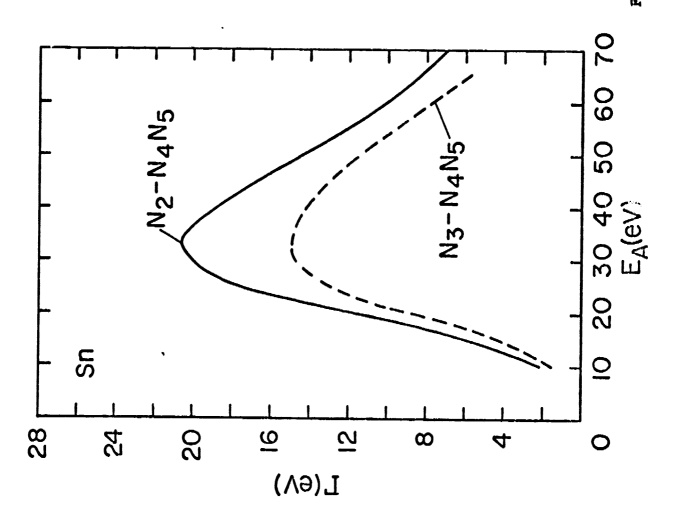
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Fig. 2

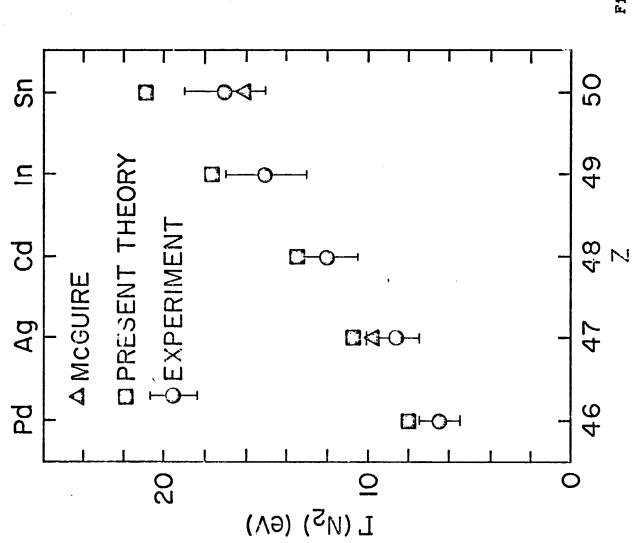


ORIGINAL OF POOR QUALITY



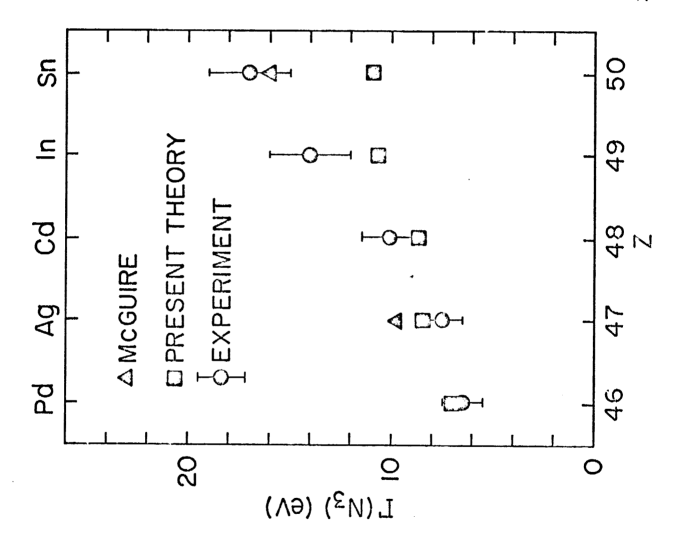


P18. 5



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F18. 6



F16. 7