

Outline

Surface, interface, and nanoscience—short introduction

Some surface concepts and techniques→photoemission

Synchrotron radiation: experimental aspects

Electronic structure—a brief review

**The basic synchrotron radiation techniques:
more experimental and theoretical details**

Core-level photoemission



Valence-level photoemission

Microscopy with photoemission (Later lecturers)

Outline



- Valence-band spectra: low-energy UPS limit and high-energy XPS limit
- Core-level chemical shifts: the potential model
- Core-level chemical shifts: equivalent-core ($Z+1$) and thermochemical energies
- Multiplet splittings
- Spin-orbit splitting, the Fano effect, and spin-polarized outgoing electrons
- Magnetic circular dichroism (MCD) in core-level emission
- Non-magnetic circular dichroism in core-level emission: a.k.a. circular dichroism in angular distributions (CDAD)
- Various other final state effects providing information in core-level spectra

PHOTOELECTRON EMISSION-

BASIC MATRIX ELEMENTS + SELECTION RULES:

• ATOMIC-LIKE (LOCALIZED) STATES $\Rightarrow$ CORE:

$$\psi_i(\vec{r}) = \psi_{n_i, l_i, m_i}(\vec{r}, \theta, \phi) = R_{n_i, l_i}(r) Y_{l_i, m_i}(\theta, \phi)$$

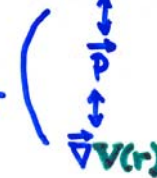


$$\psi_f(\vec{r}, \vec{k}_f) = \psi_{E_f}(\vec{r}, \vec{k}_f)$$

$$= 4\pi \sum_{l_f, m_f} i^{l_f} e^{-i\delta_f} Y_{l_f, m_f}^*(\theta, \phi) Y_{l_i, m_i}(\theta, \phi) R_{E_f, l_f}(r)$$

DIPOLE: INT. $\propto |\langle \psi_f | \hat{E} \cdot \vec{r} | \psi_i \rangle|^2 = |\hat{E} \cdot \langle \psi_f | \vec{r} | \psi_i \rangle|^2 \Rightarrow \Delta l = l_f - l_i = \pm 1$
APPROX.

EQUIVALENT
WITHIN CONSTANT
FACTOR



$\Delta m = m_f - m_i = 0, \pm 1$
TWO CHANNELS
LINEAR POLARIZ.

$\Delta m = \pm 1$, CIRCULAR POLARIZATION

VALENCE BANDS IN SOLIDS:

• BLOCH-FUNCTION (DELOCALIZED) STATES $\Rightarrow$ VALENCE:

$$\psi_i(\vec{r}) = u_{\vec{k}_i}(\vec{r}) e^{i\vec{k}_i \cdot \vec{r}}$$

$$\psi_f(\vec{r}) = u_{\vec{k}_f}(\vec{r}) e^{i\vec{k}_f \cdot \vec{r}}; E_f = \frac{p_f^2}{2m} = \frac{\hbar^2 k_f^2}{2m} \text{ USUALLY NEGLIG.}$$



$$|\langle \psi_f | \hat{E} \cdot \vec{p} | \psi_i \rangle|^2 = |\hat{E} \cdot \langle \psi_f | \vec{p} | \psi_i \rangle|^2 \Rightarrow \Delta \vec{k} = \vec{k}_f - \vec{k}_i = \vec{k}_{ph} + \vec{k}_{phonon}$$

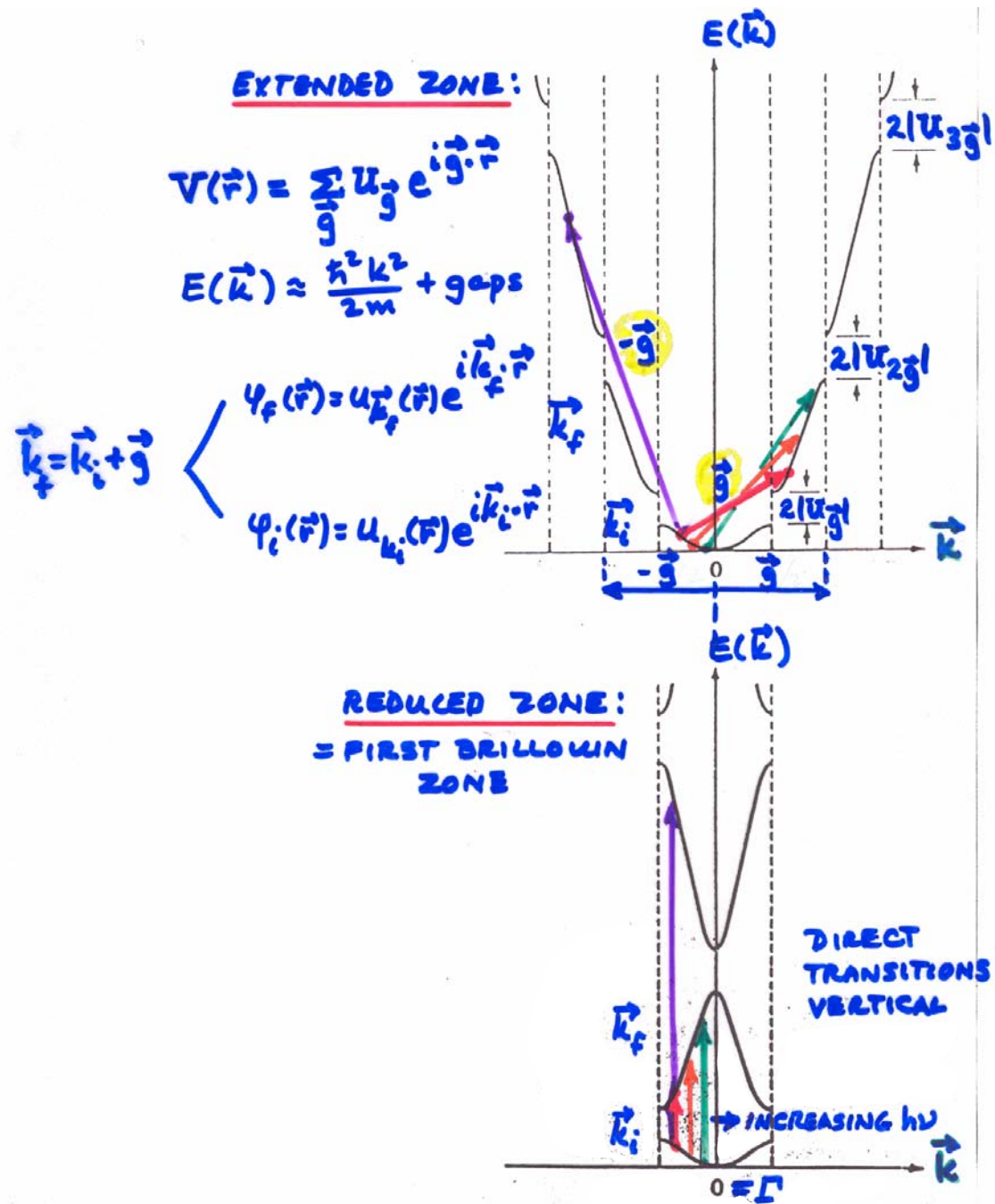
$$= \vec{g}_{BULK} \text{ (or } \vec{g}_{SURF})$$

DIRECT TRANSITIONS

BUT LATTICE VIBRATIONS $\Rightarrow$ SUM OVER $\vec{k}_{PHONON}$

$\Rightarrow$ FRACTION DIRECT $\approx$ DEBYE-WALLER FACTOR
 $= \exp[-g^2 \bar{u}^2]$

NEARLY-FREE ELECTRONS IN A WEAK PERIODIC POTENTIAL—1 DIM.

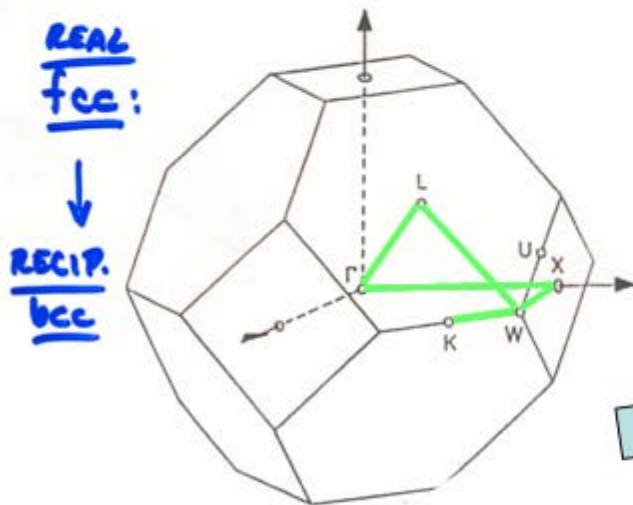


The electronic structure of a nearly free-electron metal—fcc Al

$$\phi(\vec{r}) = u_{\vec{k}}(\vec{r}) e^{i\vec{k}\cdot\vec{r}}; E(\vec{k}) \approx \frac{\hbar^2 k^2}{2m}$$

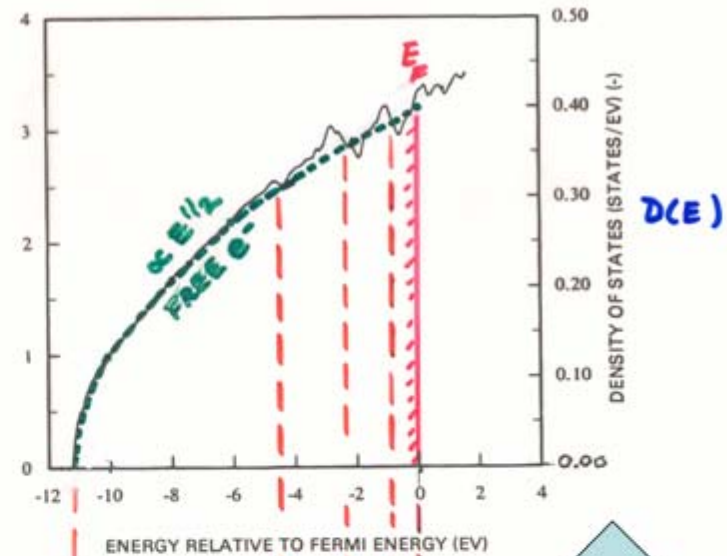
(Bloch)

3D Brillouin zone

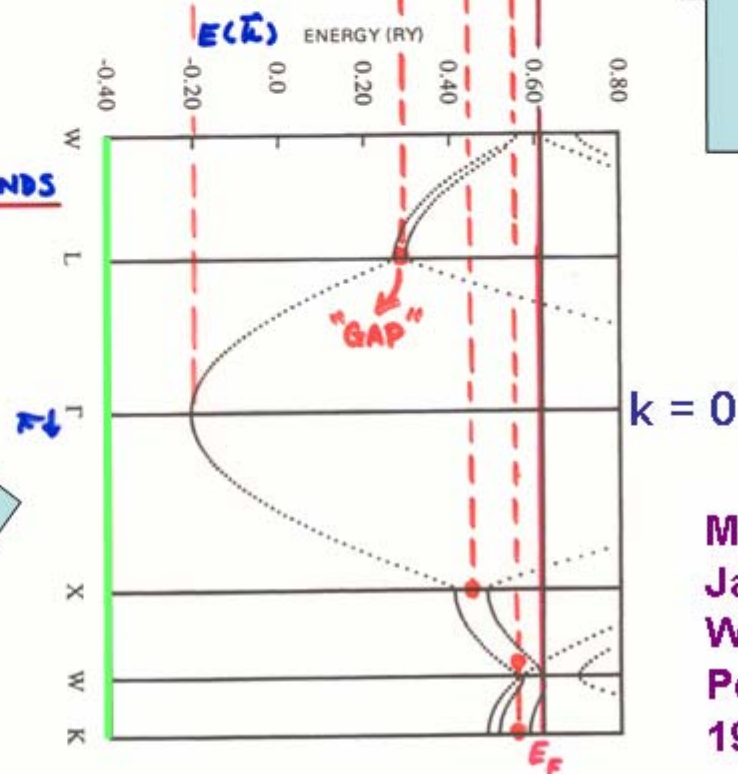


ALUMINUM - ELECTRONIC BANDS & D.O.S.

D.O.S.

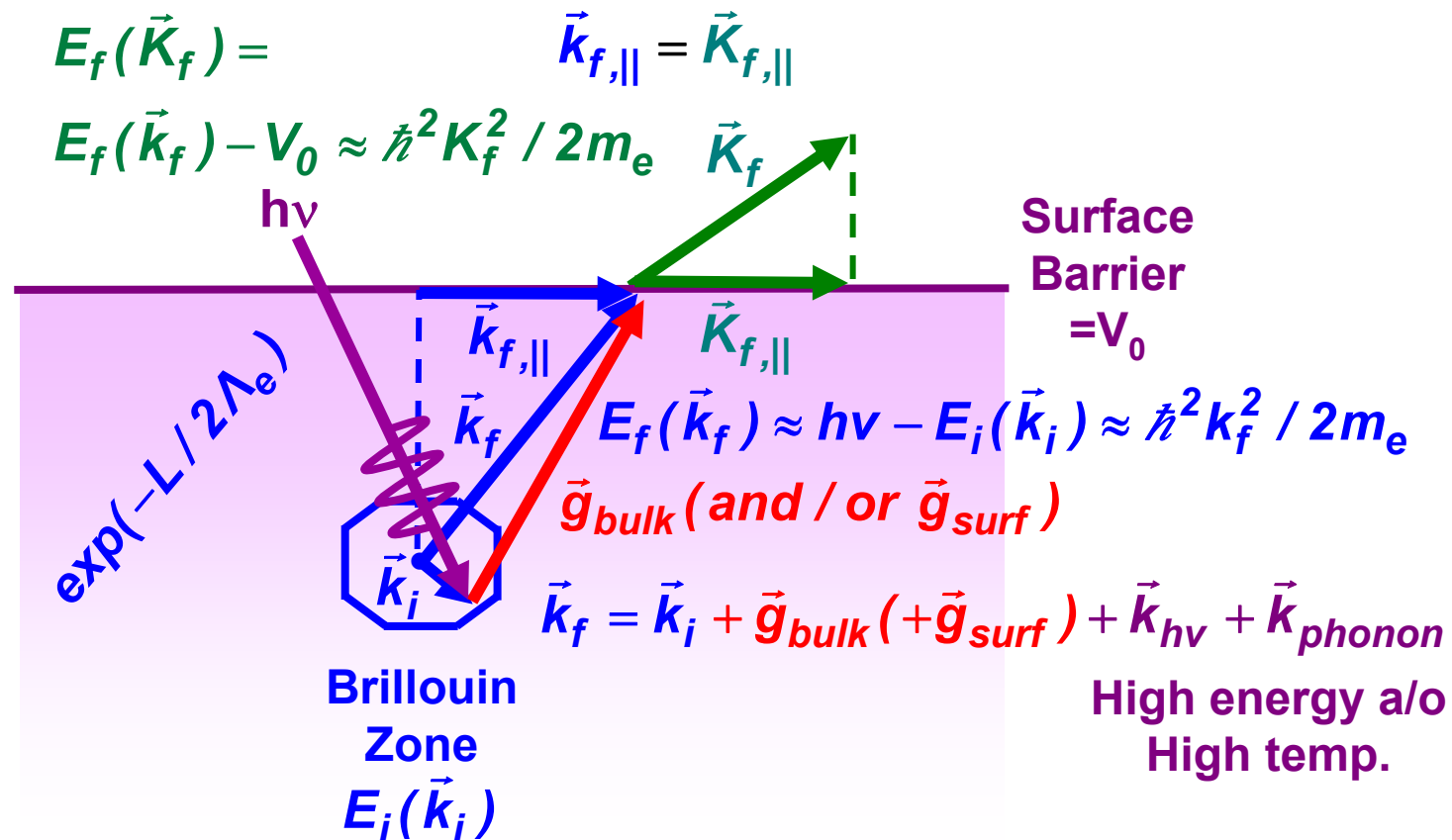


BANDS



Moruzzi,
Janak,
Williams,
Pergamon,
1978

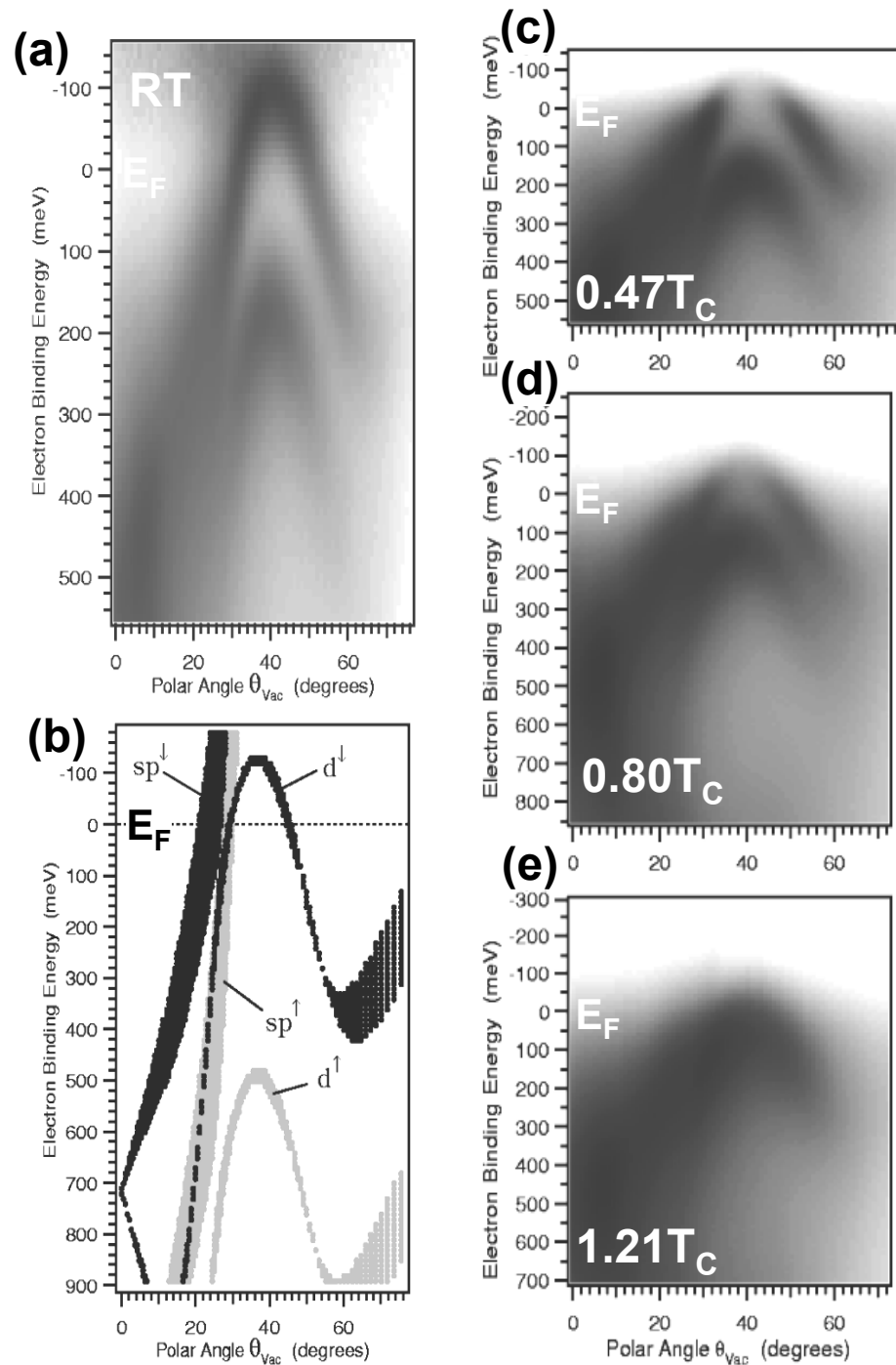
Valence-band photoemission: Angle-Resolved Photoemission (ARPES)



$$I(E_f, \vec{k}_f) \propto \left| \hat{\epsilon} \cdot \left\langle \varphi_{photoe}(E_f = h\nu + E_i, \vec{k}_f = \vec{k}_i + \vec{g}) \middle| \vec{r} \middle| \varphi(E_i, \vec{k}_i) \right\rangle \right|^2$$

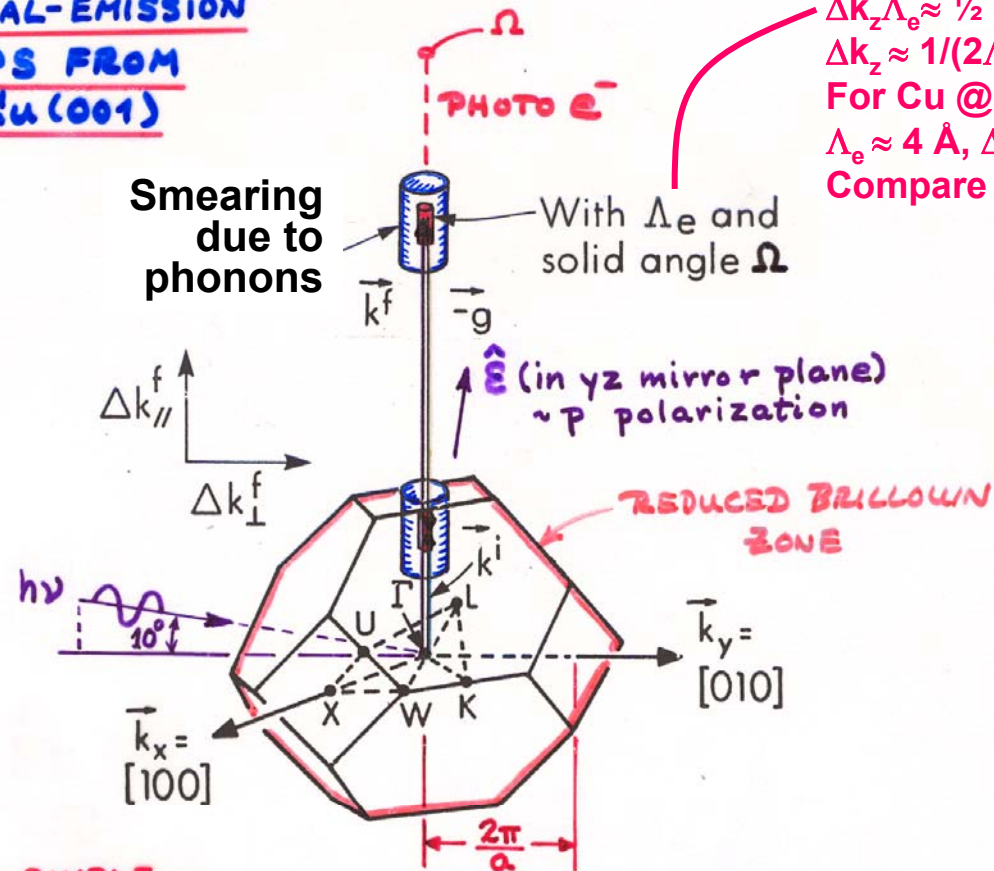
“Direct” or k-conserving transitions

**Angle-Resolved
Photoemission
from ferromagnetic
Ni(111)
 $h\nu = 21.2$ eV
Spin-split
bands**



Kreutz et al.,
Phys. Rev. B 58 (1998) 1300

EXAMPLE:
NORMAL-EMISSION
UPS FROM
Cu(001)



$$\Delta p_z \Delta z \approx \hbar/2$$

$$\Delta k_z \Lambda_e \approx 1/2$$

$$\Delta k_z \approx 1/(2\Lambda_e)$$

For Cu @ $E_{\text{kin}} \approx 80 \text{ eV}$,

$\Lambda_e \approx 4 \text{ \AA}$, $\Delta k_z \approx 0.12 \text{ \AA}^{-1}$

Compare $2\pi/a = 0.98 \text{ \AA}^{-1}$

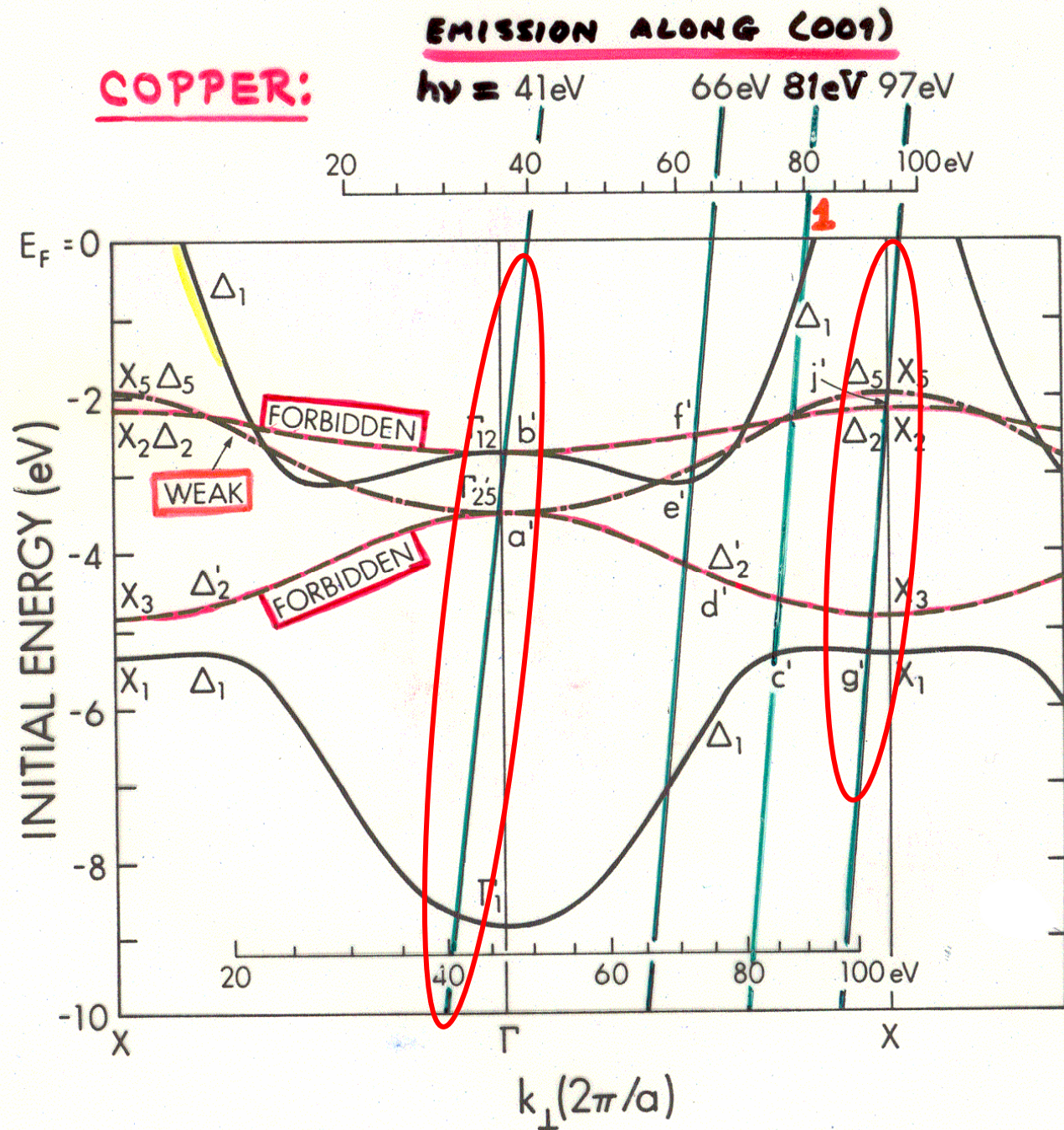
SIMPLE
DT MODEL: Direct: $\vec{k}^f = \vec{k}^i + \vec{g} + \vec{k}_{\text{ph}}$

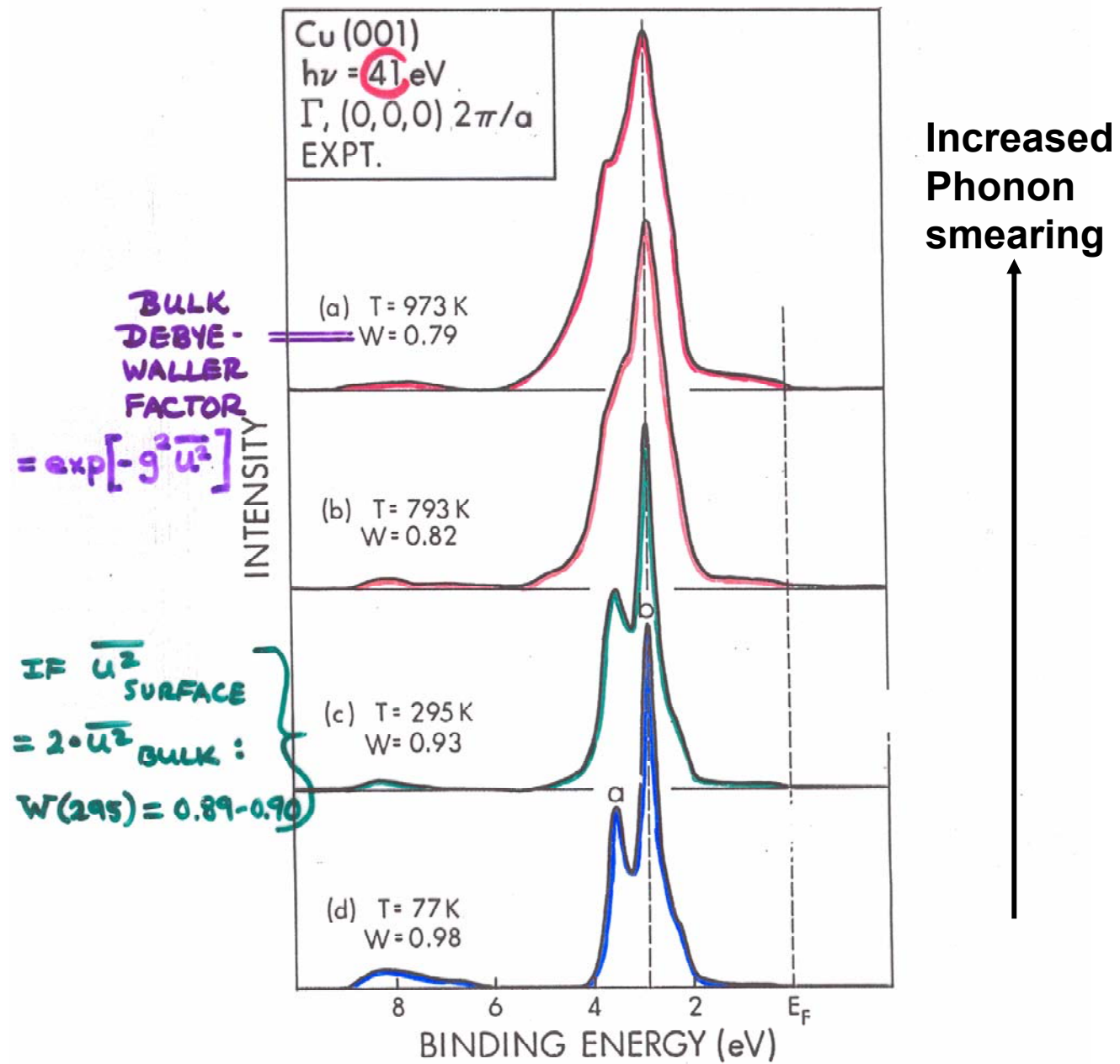
$E^i(\vec{k}^i)$ = initial band structure

$$E^f(\vec{k}^f) \approx \hbar^2 (k^f)^2 / 2m$$

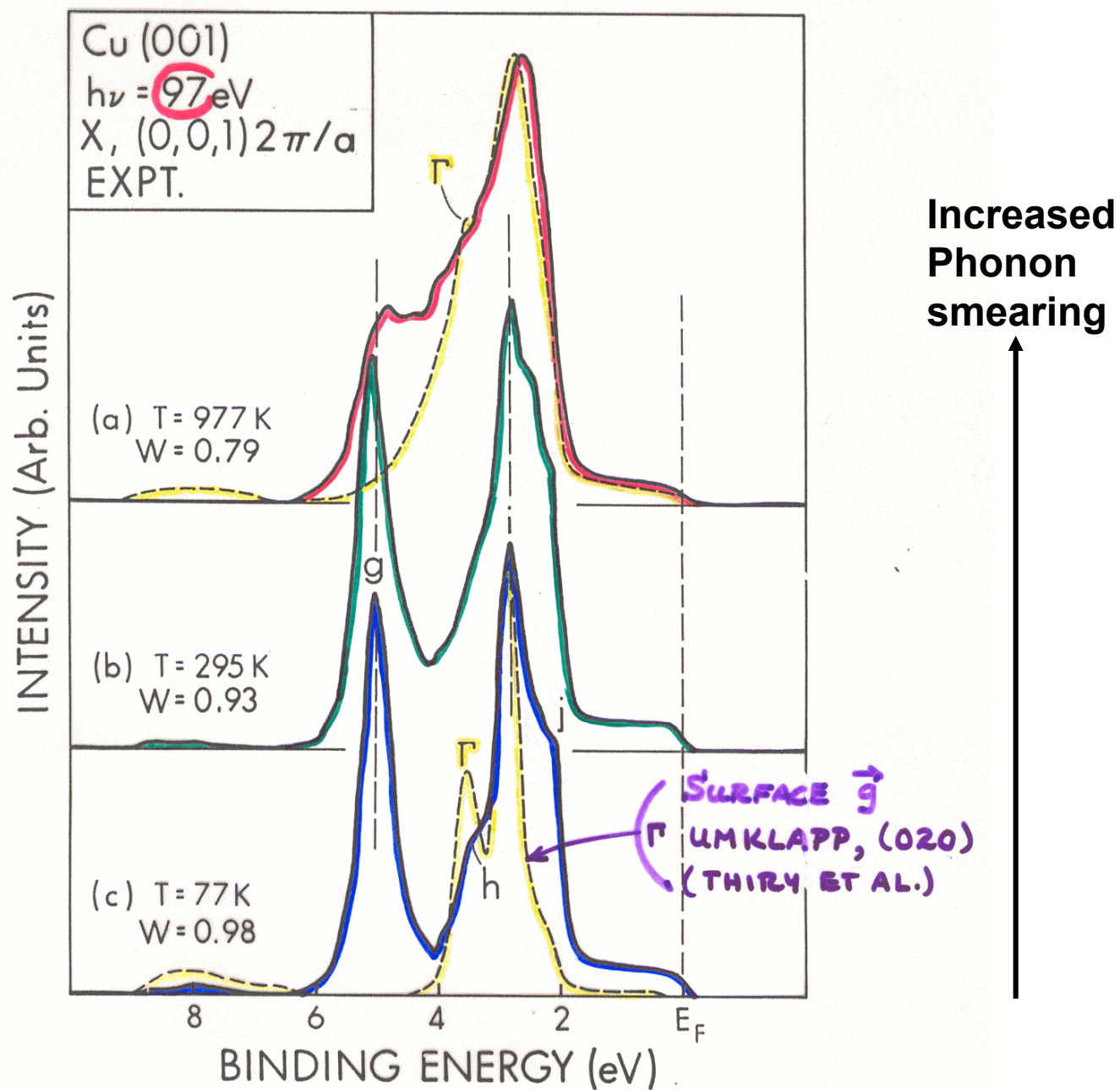
Constant matrix
 elements

Expectations
from simple
direct-
transition
theory
+ symmetry
considerations
in matrix
elements

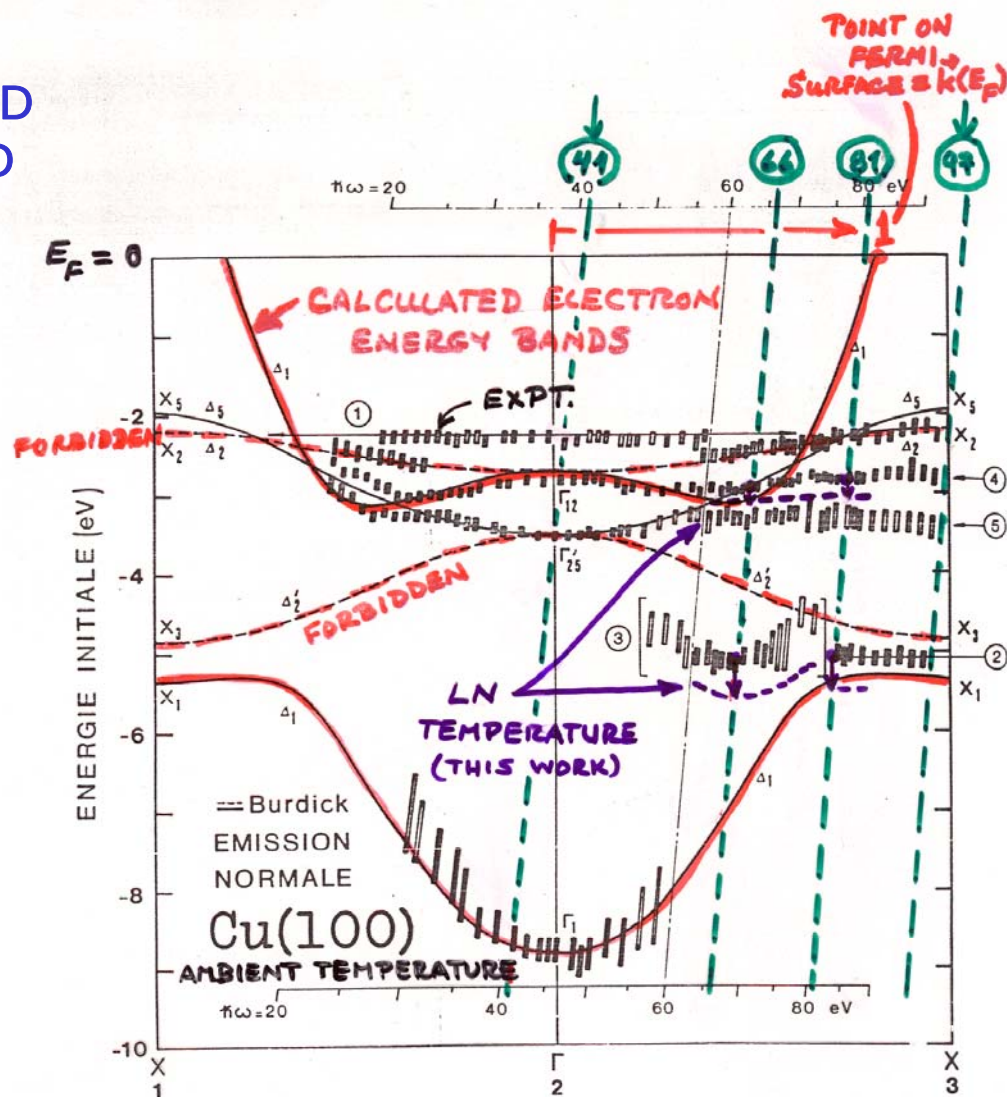




R.C. WHITE ET AL., PHYS. REV. B 35, 1147 (1987)



Cu: ANGLE-RESOLVED PHOTOEMISSION AND BAND-MAPPING ALONG (001)



P. THIRY, THESIS, UNIV. OF PARIS
(1980)

+ WHITE ET AL.
P.R. 835,1147
(1987)

FIG.56

Cu: ANGLE-RESOLVED PHOTOEMISSION AND BAND-MAPPING ALONG (110)

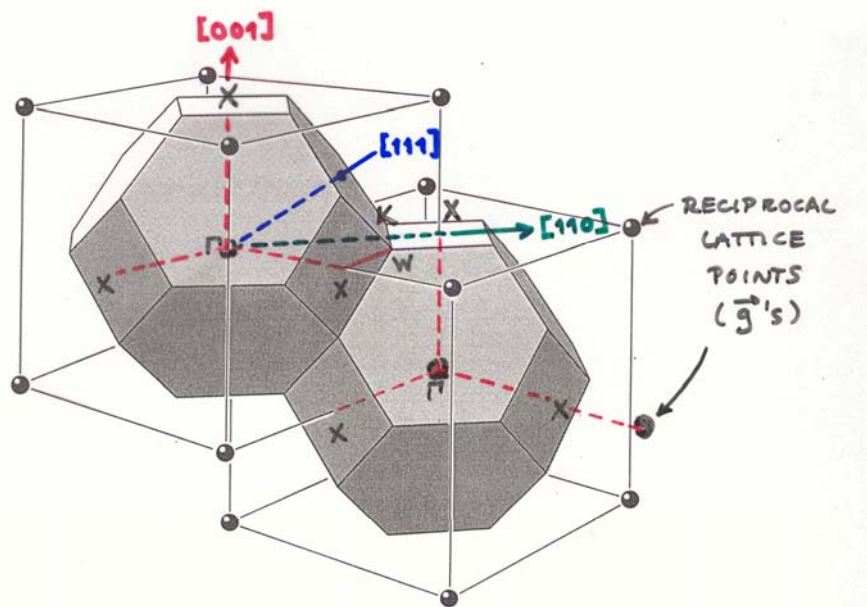
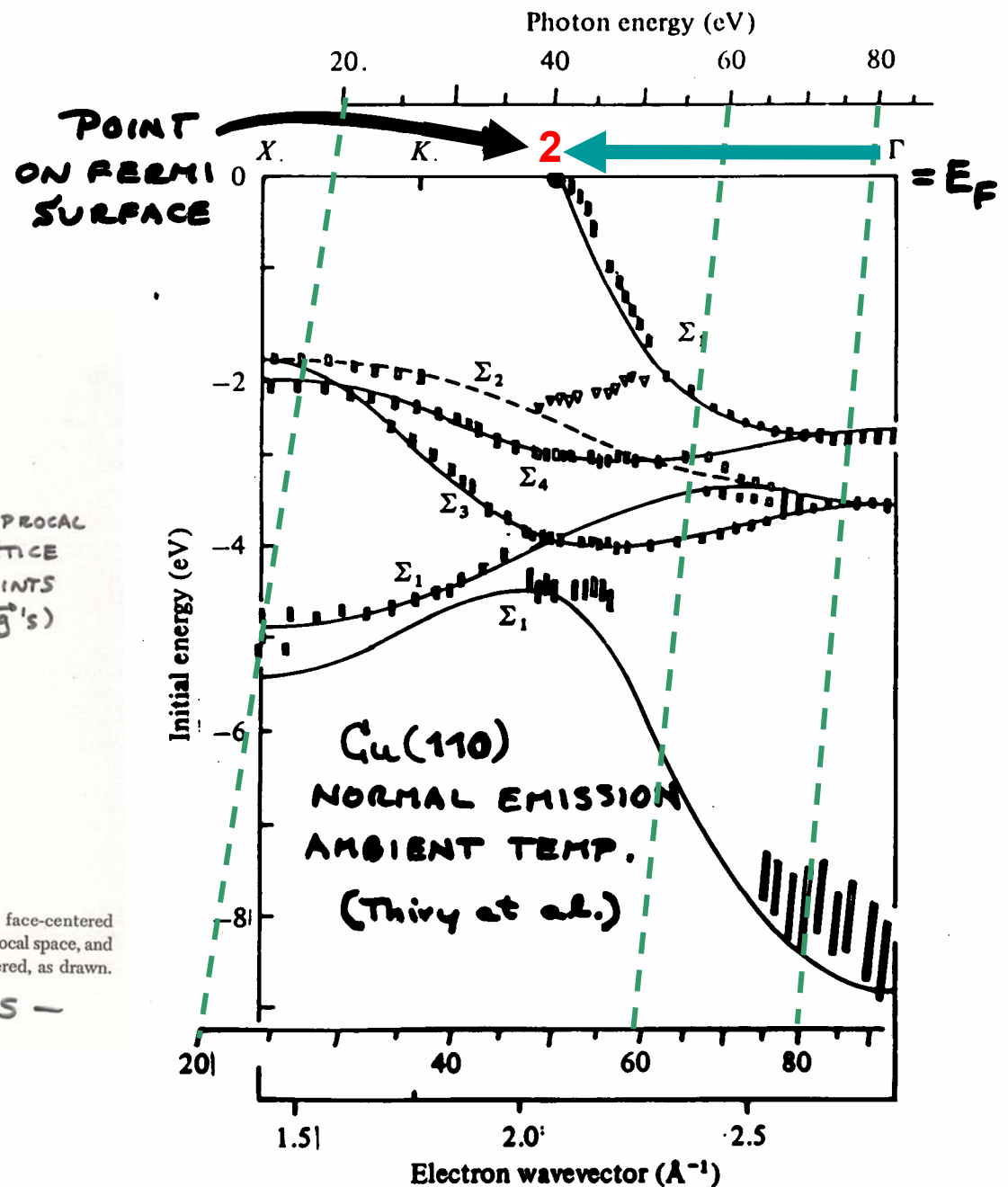
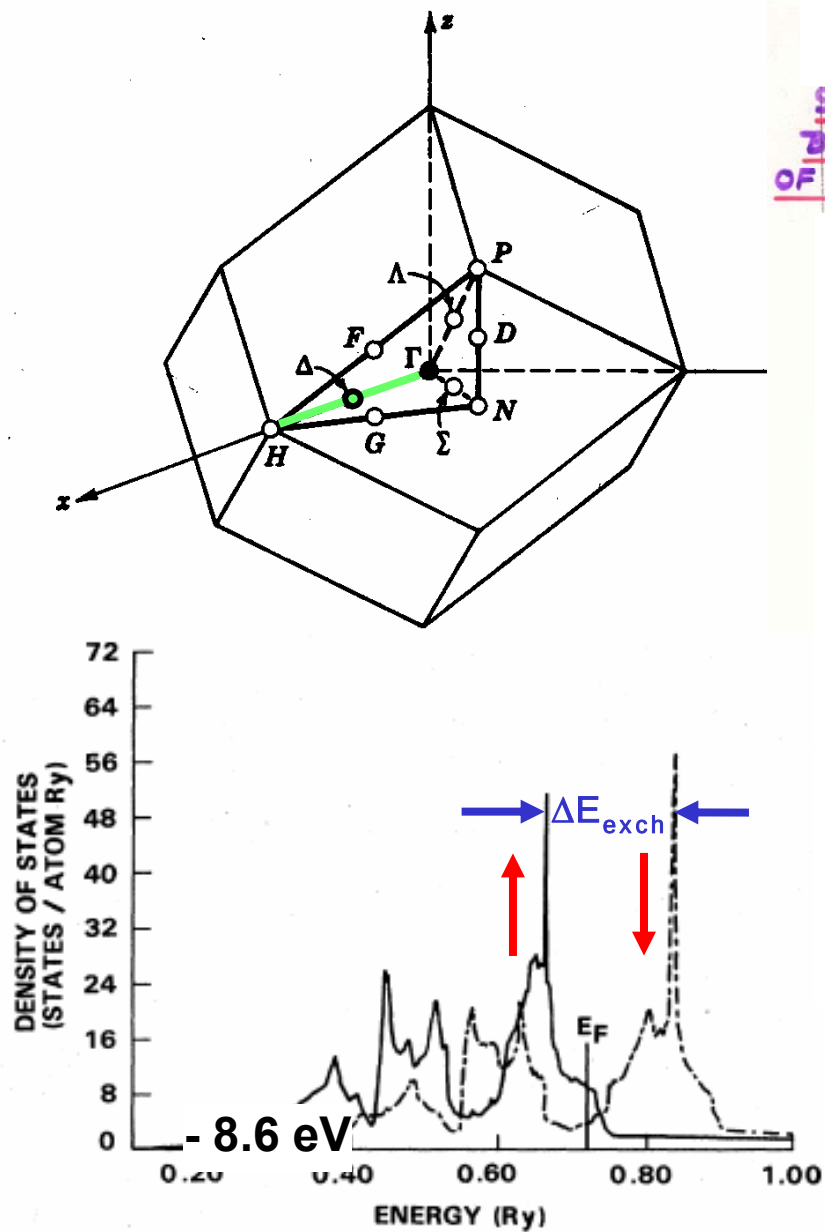


Figure 28 Brillouin zones of the face-centered cubic lattice. The cells are in reciprocal space, and the reciprocal lattice is body-centered, as drawn.

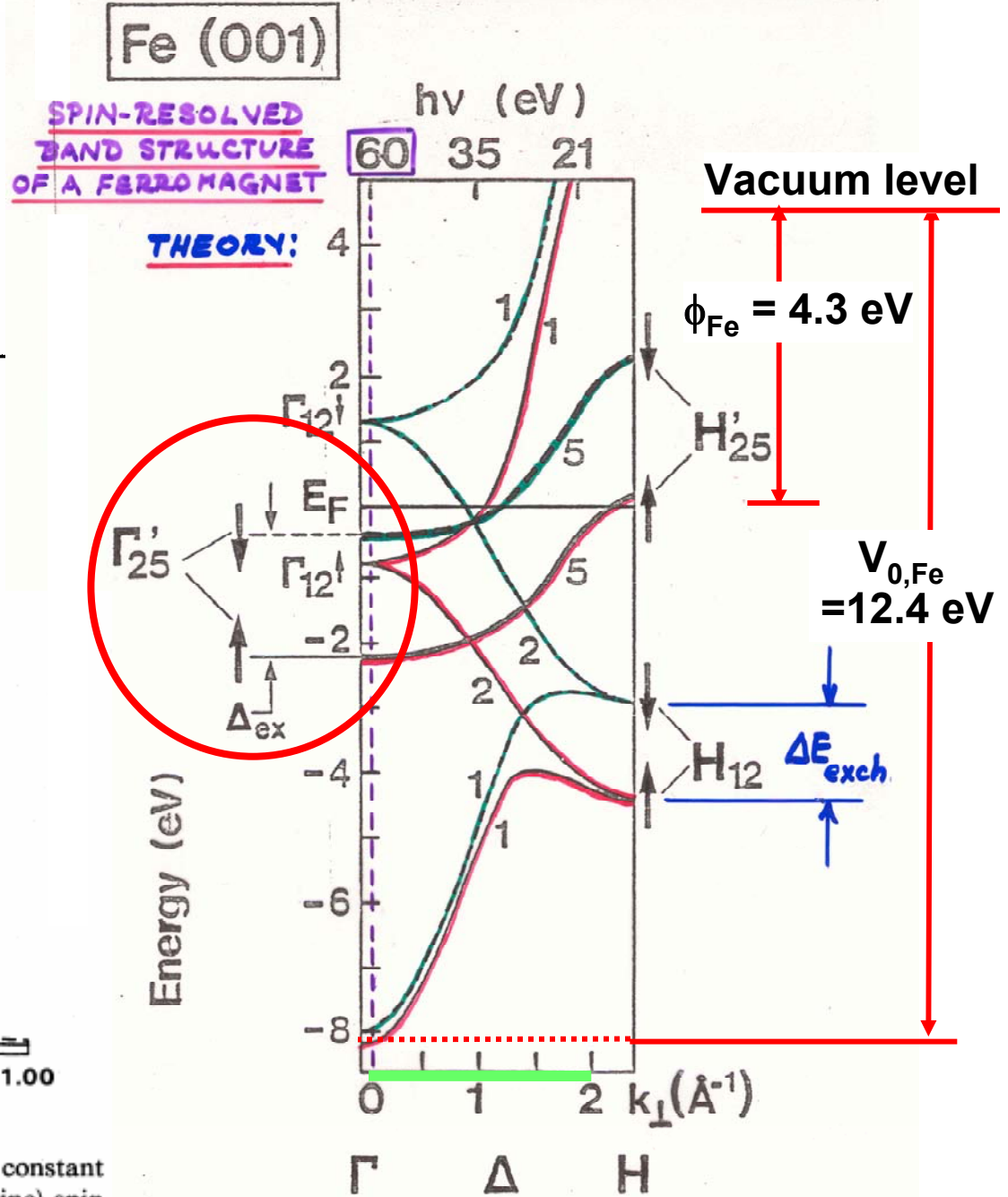
— STACKING OF fcc BRILLOUIN ZONES —

P.Thiry, Ph.D.
thesis, Univ.
of Paris (1980)



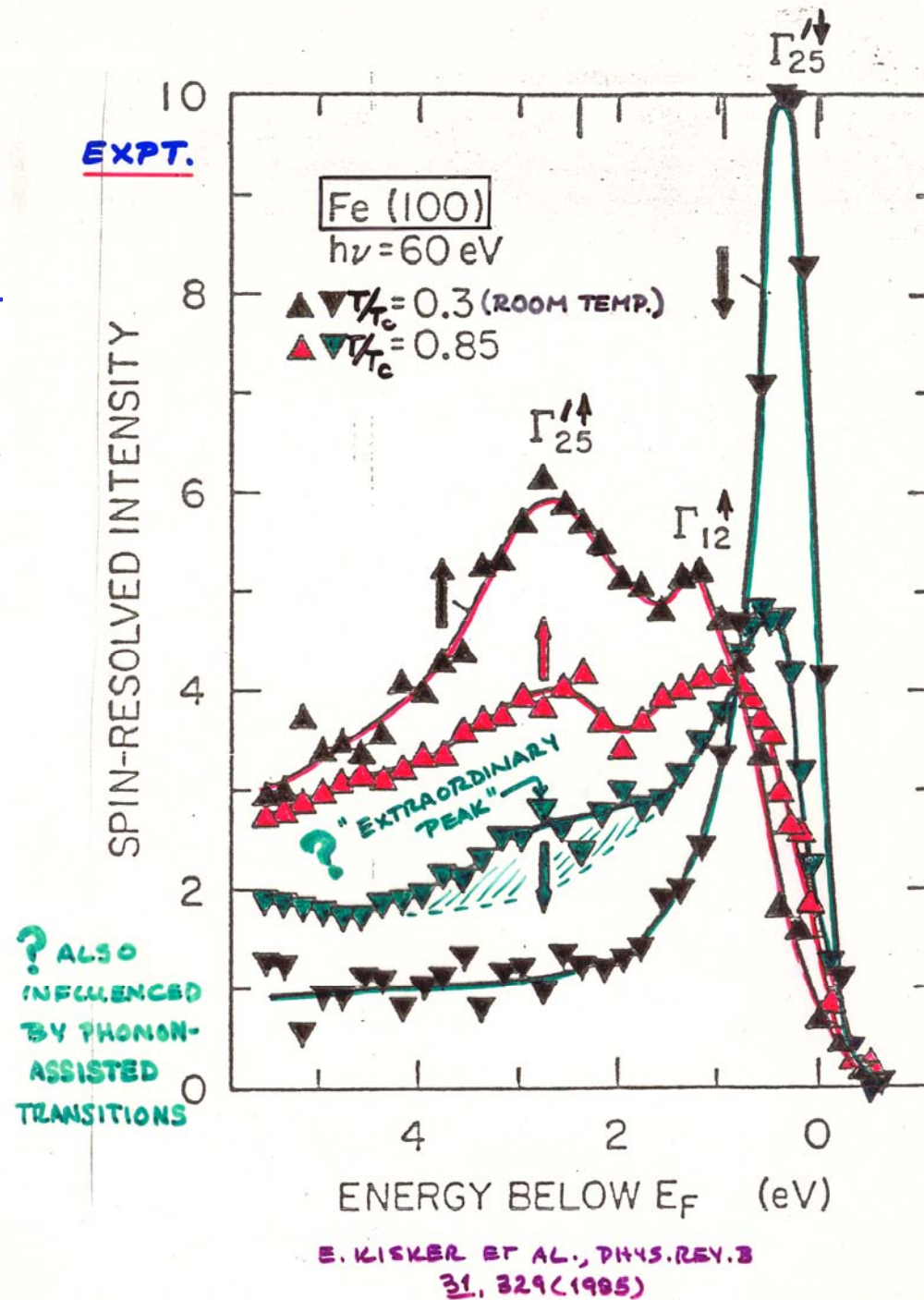


Hathaway et al., Phys. Rev. B 31, 7603 ('85)



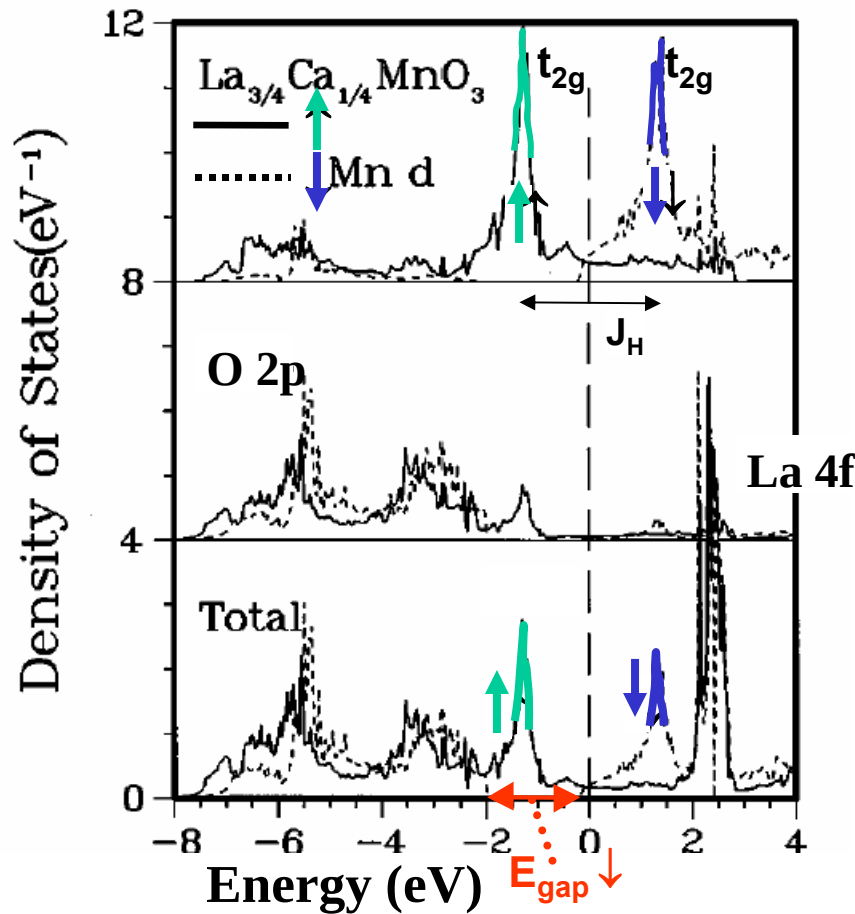
E. KISKER ET AL., PHYS. REV. B 31, 329 (1985)

Fe: ANGLE AND SPIN-RESOLVED SPECTRA AT Γ POINT



Half-Metallic Ferromagnetism

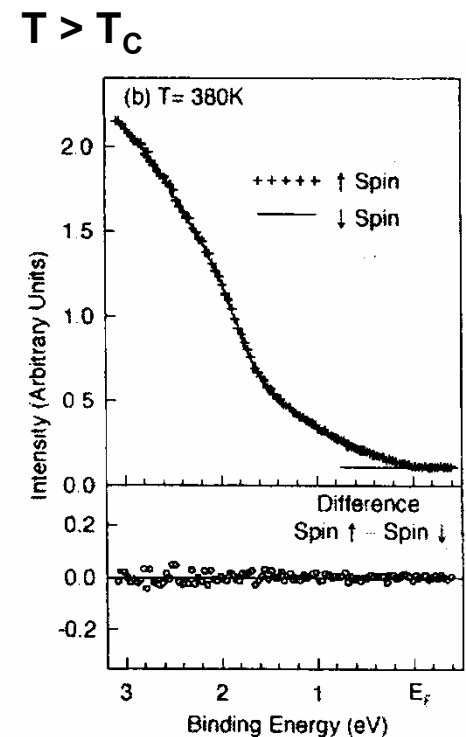
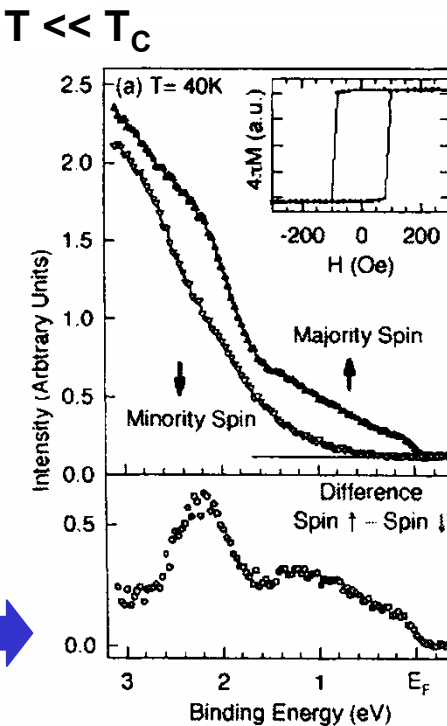
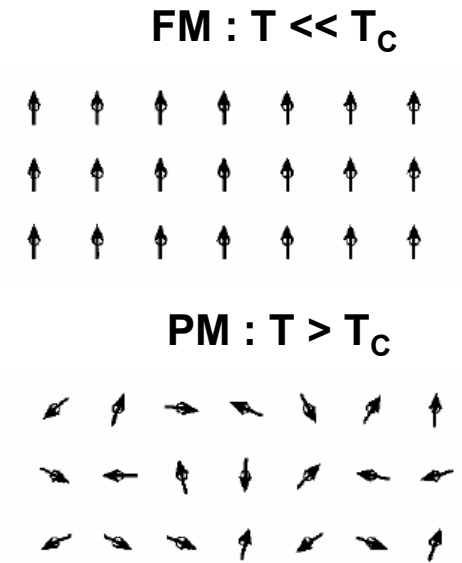
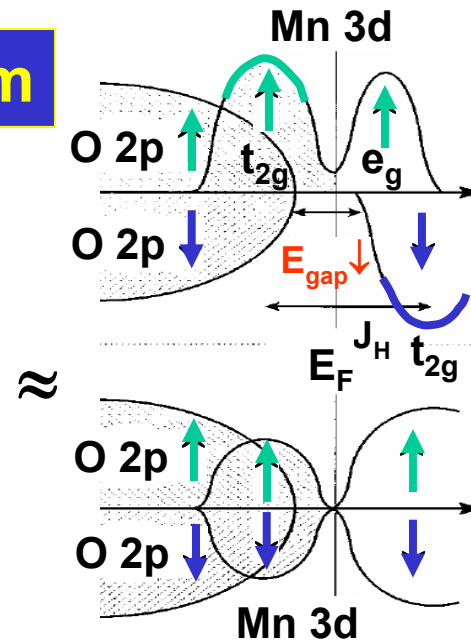
LDA theory- FM $\text{La}_{0.75}\text{Ca}_{0.25}\text{MnO}_3$



Pickett and Singh, PRB **53**, 1146 (1996)

Experiment- spin-resolved PS
 $\text{La}_{0.70}\text{Sr}_{0.30}\text{MnO}_3$ as thin film

Park et al., Nature, PRB **392**, 794 (1998)

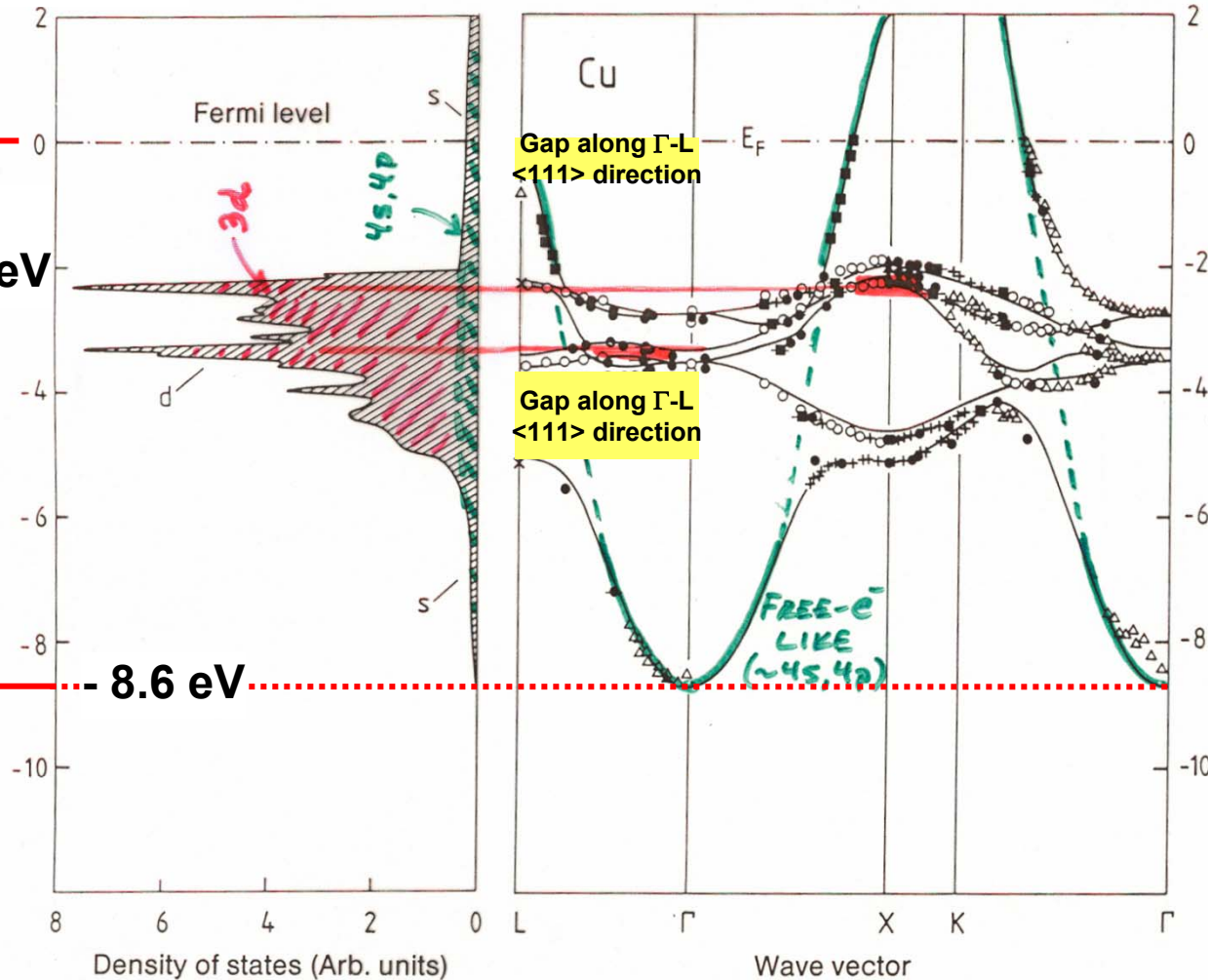


Vacuum level

The electronic structure of a transition metal—fcc Cu

$$\phi_{\text{Cu}} = 4.4 \text{ eV}$$

$$V_{0,\text{Cu}} = 13.0 \text{ eV}$$



Cu $1s^2 \dots 3d^{10} 4s^1$
ELECTRONIC BANDS
+ DENSITY OF STATE

} MIXING
 } 3d LIKE
 } MIXING

Experimental points from angle-resolved photoelectron spectroscopy

Fig. 7.12. Bandstructure $E(k)$ for copper along directions of high crystal symmetry (right). The experimental data were measured by various authors and were presented collectively by Courths and Hüfner [7.4]. The full lines showing the calculated energy bands and the density of states (left) are from [7.5]. The experimental data agree very well, not only among themselves, but also with the calculation

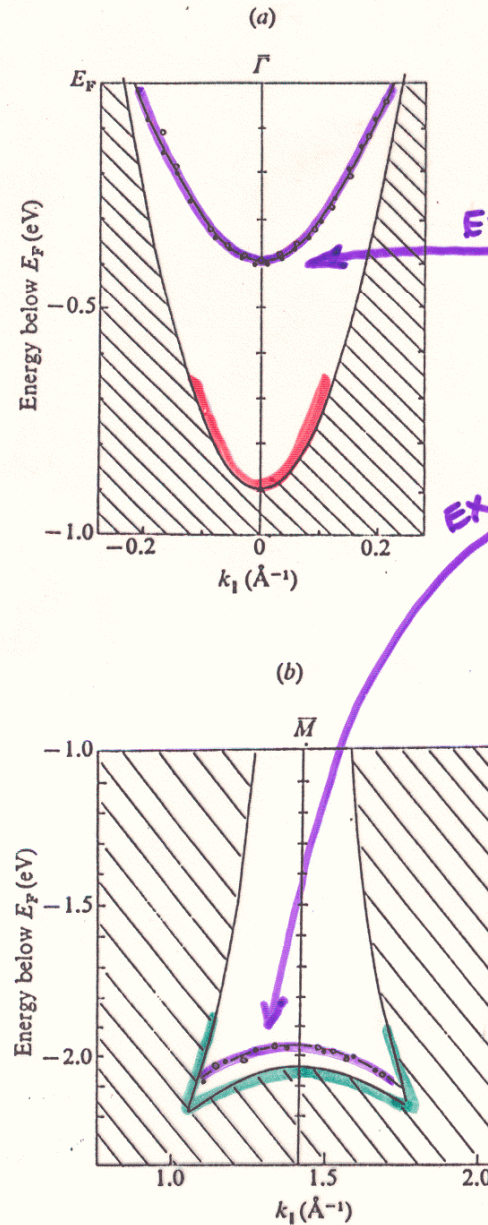
Surface states on Cu(111)

Shockley
surface
state

Tamm
surface
state

Zangwill,
Surface Physics,

Fig. 4.21. Experimental dispersion of Cu(111) surface states plotted with a projection of the bulk bands: (a) Shockley state near the zone center (Γ point, Kevan, 1983); (b) Tamm state near the zone boundary ($\bar{M}$ point, Heimann, Hermanson, Miosga and Neddermeyer, 1979). Compare with Fig. 4.17.



THEORY

Fig. 4.17. Surface states (dashed curves) and bulk projected bands of Cu(111) surface according to a six-layer surface band structure calculation (Euceda, Bylander & Kleinman, 1983).

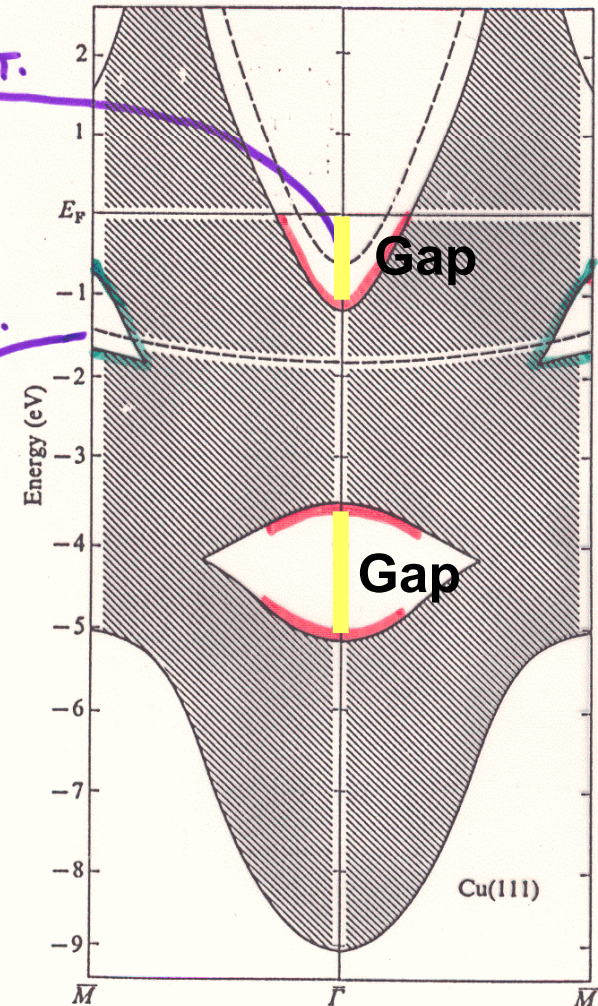
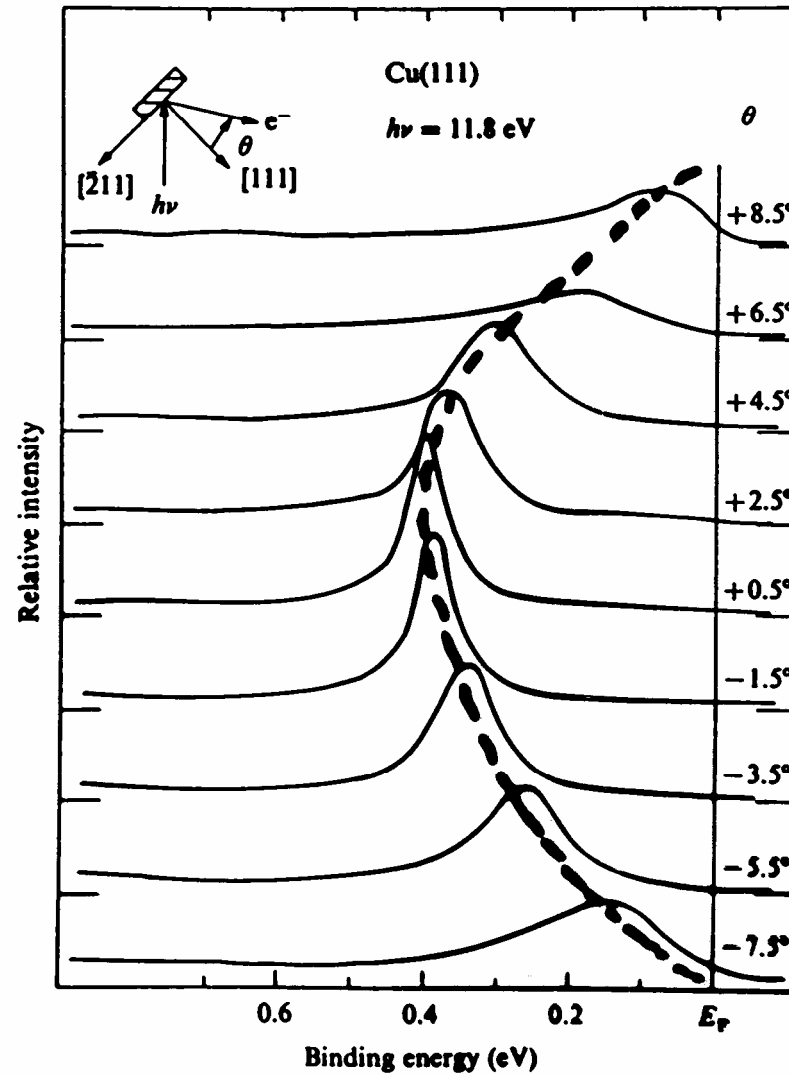
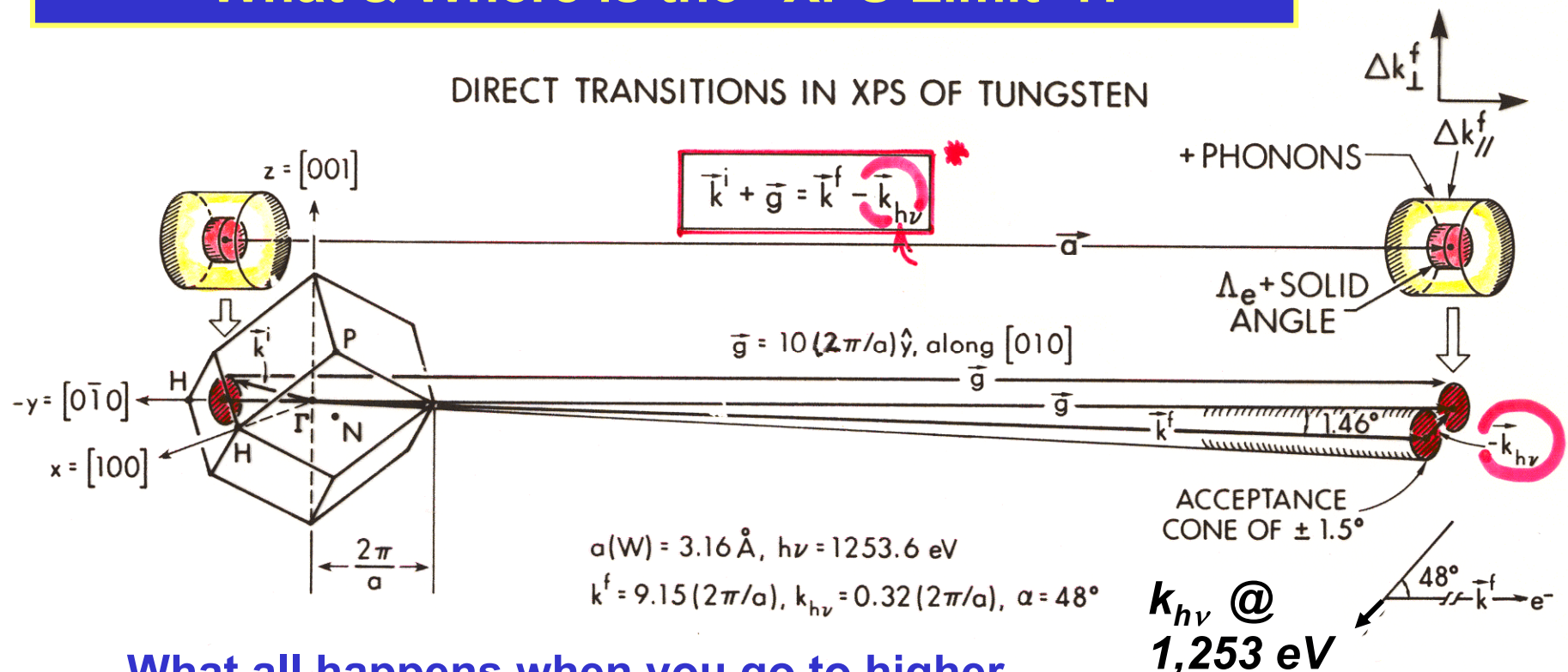


Fig. 4.20. Photoemission energy distribution curves from Cu(111) at different collection angles. Equation (4.32) has been used to express the electron kinetic energy in terms of the binding energy of the electron state (Kevan, 1983).



Zangwill,
Surface Physics,

Valence-Band Photoemission at High Energy-- What & Where is the “XPS Limit”?:



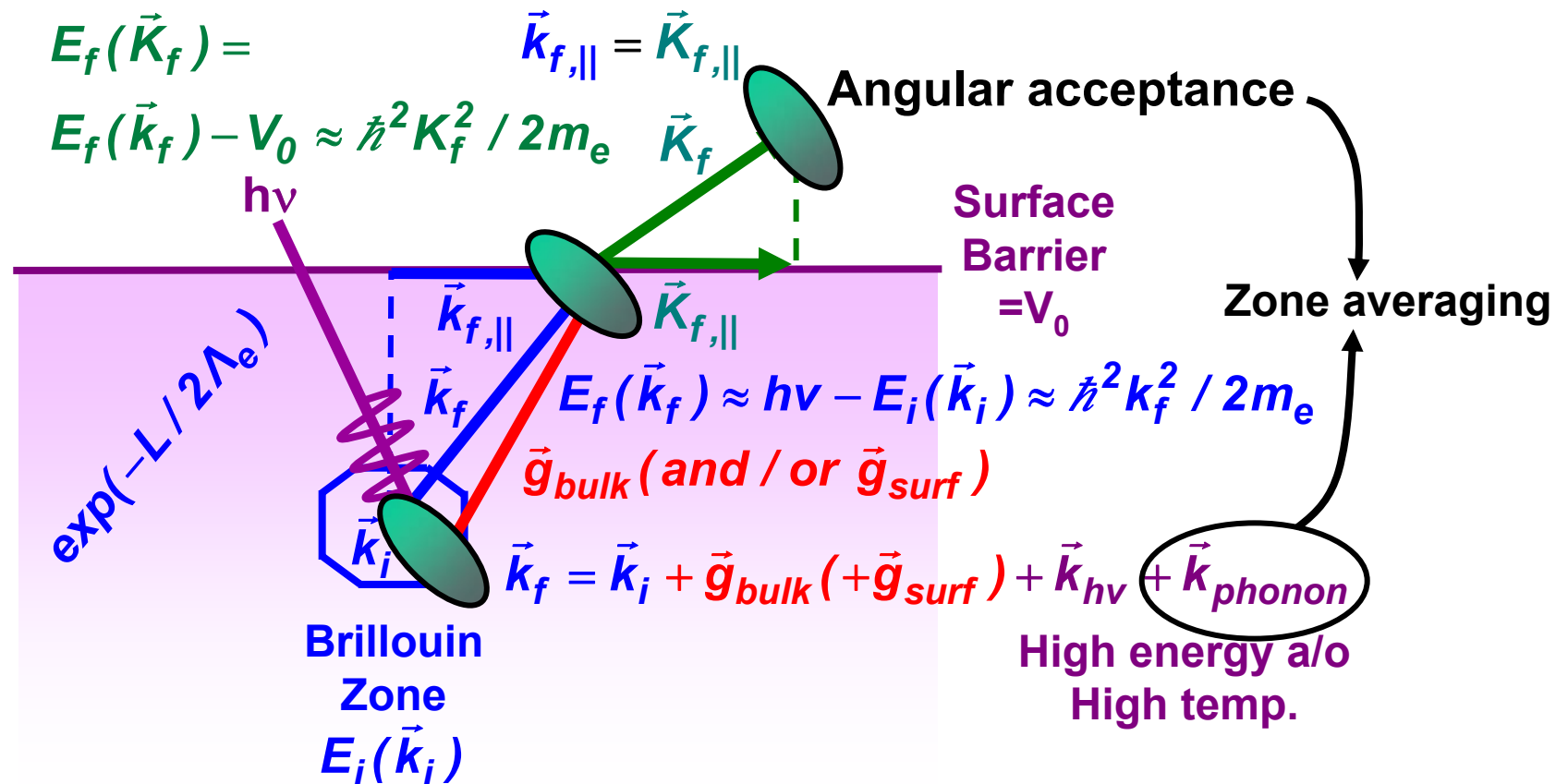
What all happens when you go to higher photon energies?

- non-dipole effect → the photon momentum
- angular acceptance → B.Z. averaging
- lattice recoil, phonon creation → more Brillouin Zone averaging

→ The XPS limit of full B.Z. averaging and D.O.S. sensitivity

Hussain et al., Phys. Rev. B 22, 3750 ('80)

Valence-band photoemission—at higher energy



$$I(E_f, \vec{k}_f) \propto \left| \hat{\epsilon} \cdot \left\langle \varphi_{photoe}(E_f = h\nu + E_i, \vec{k}_f = \vec{k}_i + \vec{g}) \middle| \vec{r} \middle| \varphi(E_i, \vec{k}_i) \right\rangle \right|^2$$

$$h\nu = 1487 \text{ eV}$$

Estimating phonon effects: 1st approx.

$$W(T) = \text{Debye-Waller factor} = \exp\left(-\frac{1}{3}g^2\langle U^2(T)\rangle\right)$$

$$I(E, T) = W(T)I_{DT} + [1 - W(T)]I_{NDT}$$

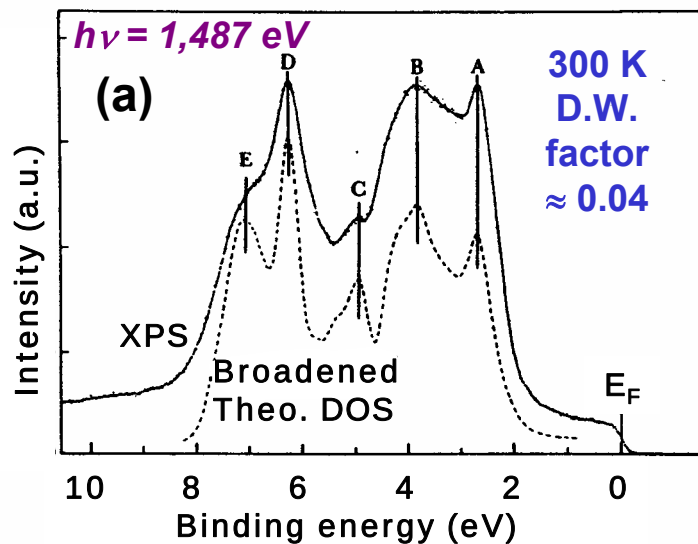
Shevchik, Phys. Rev. B 20, 3020 ('79); Hussain et al., Phys. Rev. 22, 3750 ('80)

TABLE I. Tabulation of thermal displacements $\langle U^2 \rangle$ and Debye-Waller factors $[W(T) = \exp(-\frac{1}{3}\langle U^2 \rangle g^2)]$ in the XPS regime ($E_{\text{kin}} = 1482 \text{ eV}$) for various elements. The Debye temperatures are taken from Ref. 30. The solid line divides the Debye-Waller factors so as to indicate when $\approx \frac{1}{2}$ of the transitions will be direct. Elements are in order of Z .

Element	Θ_D (K)	A (amu)	$T = 4 \text{ K}$		$T = 77 \text{ K}$		$T = 300 \text{ K}$		$T = 1000 \text{ K}$	
			$\langle U^2 \rangle$ (10^{-12} cm^2)	W	$\langle U^2 \rangle$ (10^{-12} cm^2)	W	$\langle U^2 \rangle$ (10^{-12} cm^2)	W	$\langle U^2 \rangle$ (10^{-12} cm^2)	W
Bc	1440	9.012	0.54	0.34	0.86	0.33	1.07	0.25	2.47	0.04
C	2230	12.001	0.41	0.33	0.41	0.53	0.74	0.33	0.83	0.34
Mo	400	24.212	1.13	0.22	1.48	0.16	2.52	0.01		
Al	428	26.981	0.95	0.20	1.15	0.22	2.79	0.03		
Si	643	28.086	0.60	0.46	0.66	0.42	1.26	0.19	3.07	0.02
Ca	230	40.08	1.19	0.21	1.94	0.08	6.24	0.0		
Ti	420	47.9	0.54	0.49	0.66	0.42	1.63	0.12	3.16	0.0
V	380	50.942	0.56	0.48	0.72	0.39	1.85	0.09	5.93	0.0
Cr	630	51.996	0.33	0.68	0.37	0.58	0.71	0.40	2.15	0.06
Nb	410	94.938	0.48	0.50	0.60	0.46	1.49	0.14	4.72	0.0
Fe	470	55.847	0.42	0.52	0.43	0.53	1.13	0.23	3.58	0.01
Co	443	58.933	0.42	0.52	0.40	0.52	1.19	0.21	3.74	0.0
Ni	450	58.71	0.41	0.53	0.43	0.50	1.17	0.22	3.67	0.01
Cu	343	63.54	0.30	0.62	0.67	0.42	1.82	0.09	5.84	0.0
Zn	327	65.37	0.31	0.61	0.70	0.40	1.92	0.08		
Ge	374	72.59	0.40	0.50	0.50	0.51	1.34	0.17	4.28	0.0
As	282	74.922	0.32	0.61	0.77	0.37	2.25	0.05	7.32	0.0
Zr	291	91.22	0.41	0.50	0.60	0.46	1.74	0.10	5.65	0.0
Nb	275	92.906	0.43	0.57	0.65	0.43	1.90	0.08	6.21	0.0
Mo	450	95.94	0.25	0.72	0.30	0.65	0.71	0.32	2.25	0.05
Ru	600	101.07	0.13	0.78	0.28	0.77	0.40	0.60	1.21	0.21
Rh	480	102.905	0.22	0.75	0.26	0.81	0.59	0.46	1.84	0.09
Pd	274	106.4	0.27	0.61	0.57	0.48	1.68	0.11	5.46	0.0
Ag	225	107.87	0.45	0.56	0.75	0.38	2.44	0.04	5.00	0.0
Cd	209	112.40	0.47	0.55	0.81	0.35	2.70	0.03		
Sa	200	118.69	0.46	0.55	0.83	0.34	2.55	0.04		
Sb	211	121.75	0.43	0.57	0.79	0.35	2.46	0.04		
Hf	252	178.49	0.24	0.73	0.38	0.60	1.18	0.22	3.85	0.01
Ta	240	180.948	0.25	0.72	0.42	0.59	1.23	0.19	4.19	0.0
W	400	183.83	0.12	0.82	0.15	0.79	0.47	0.53	1.48	0.14
Re	630	186.2	0.15	0.80	0.17	0.81	0.40	0.60	1.23	0.19
Os	500	190.2	0.11	0.86	0.15	0.84	0.30	0.69	0.92	0.30
Ir	420	192.2	0.14	0.84	0.17	0.81	0.41	0.59	1.29	0.19
Pt	240	195.09	0.23	0.74	0.33	0.60	1.19	0.21	3.85	0.01
Au	165	196.967	0.34	0.65	0.52	0.35	2.46	0.04	5.14	0.0
Pb	105	207.19	0.51	0.52	1.54	0.13	3.75	0.0		

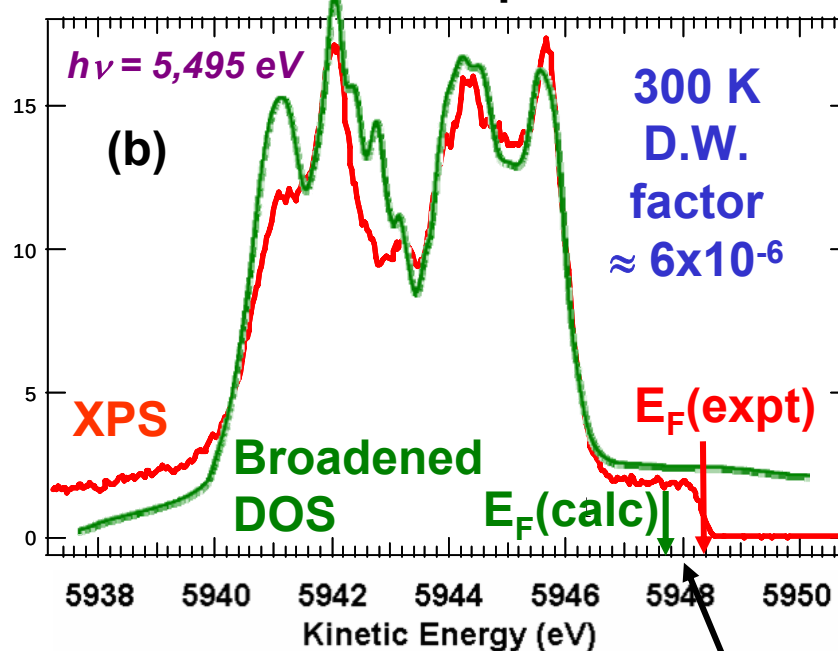
Greater than 50% direct transitions

Gold Valence Spectrum



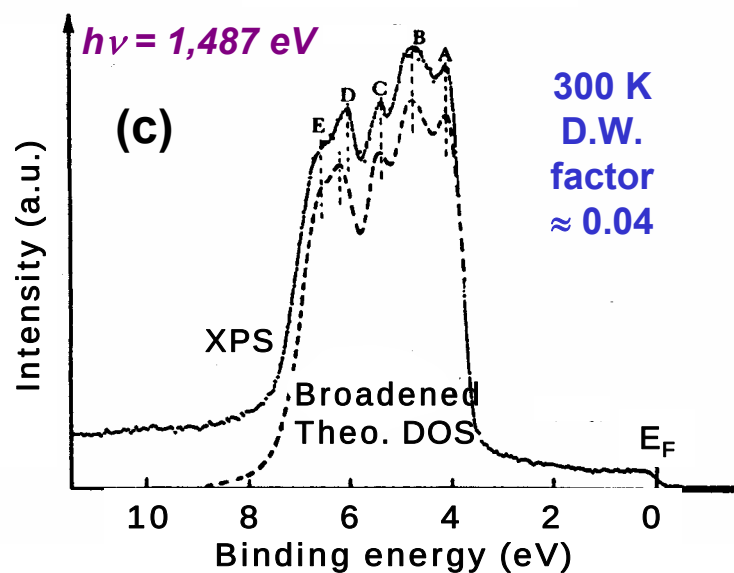
Valence spectra in the XPS Limit

Gold Valence Spectrum

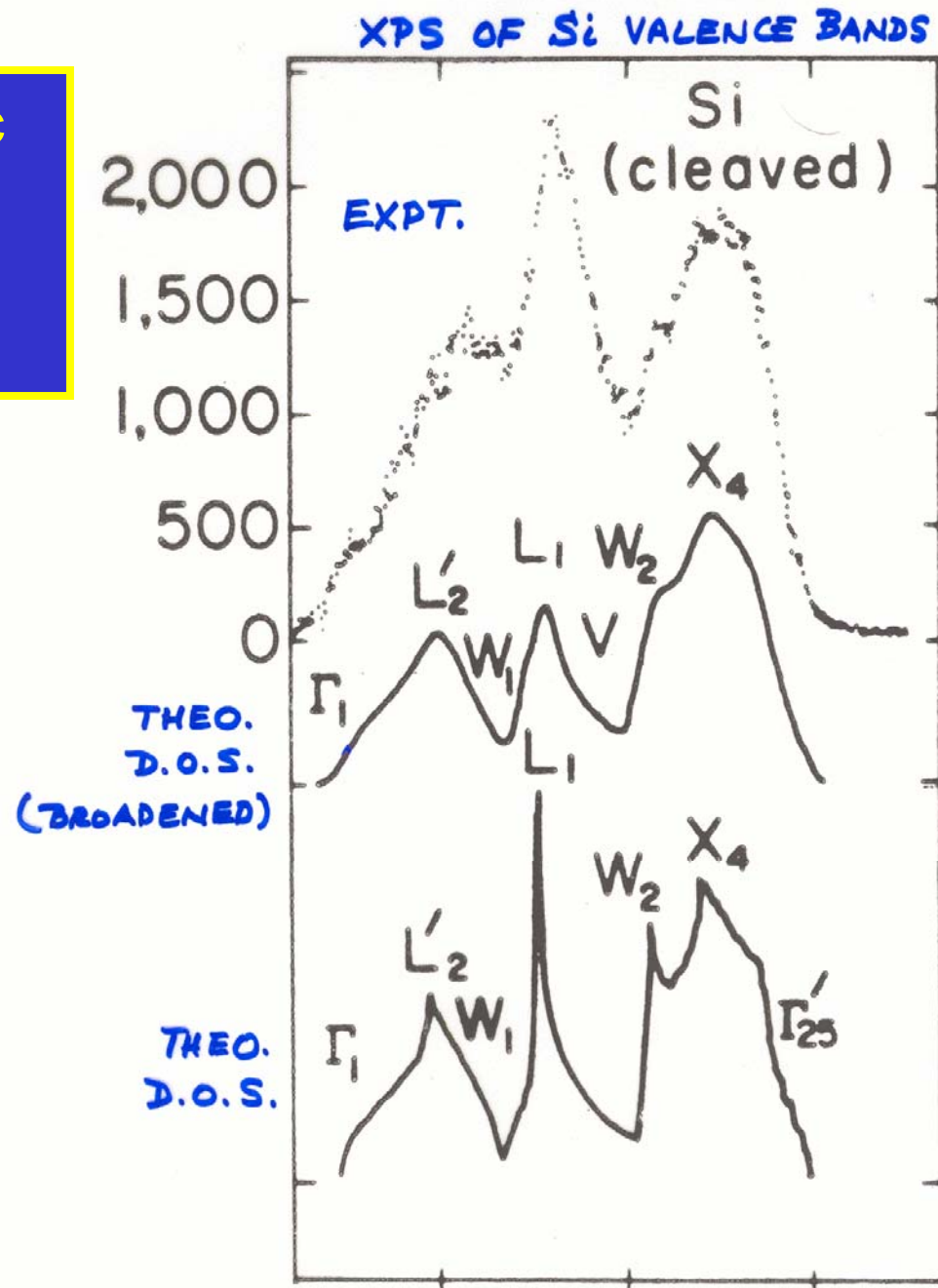


Screening/
self-energy
correction

Silver Valence Spectrum

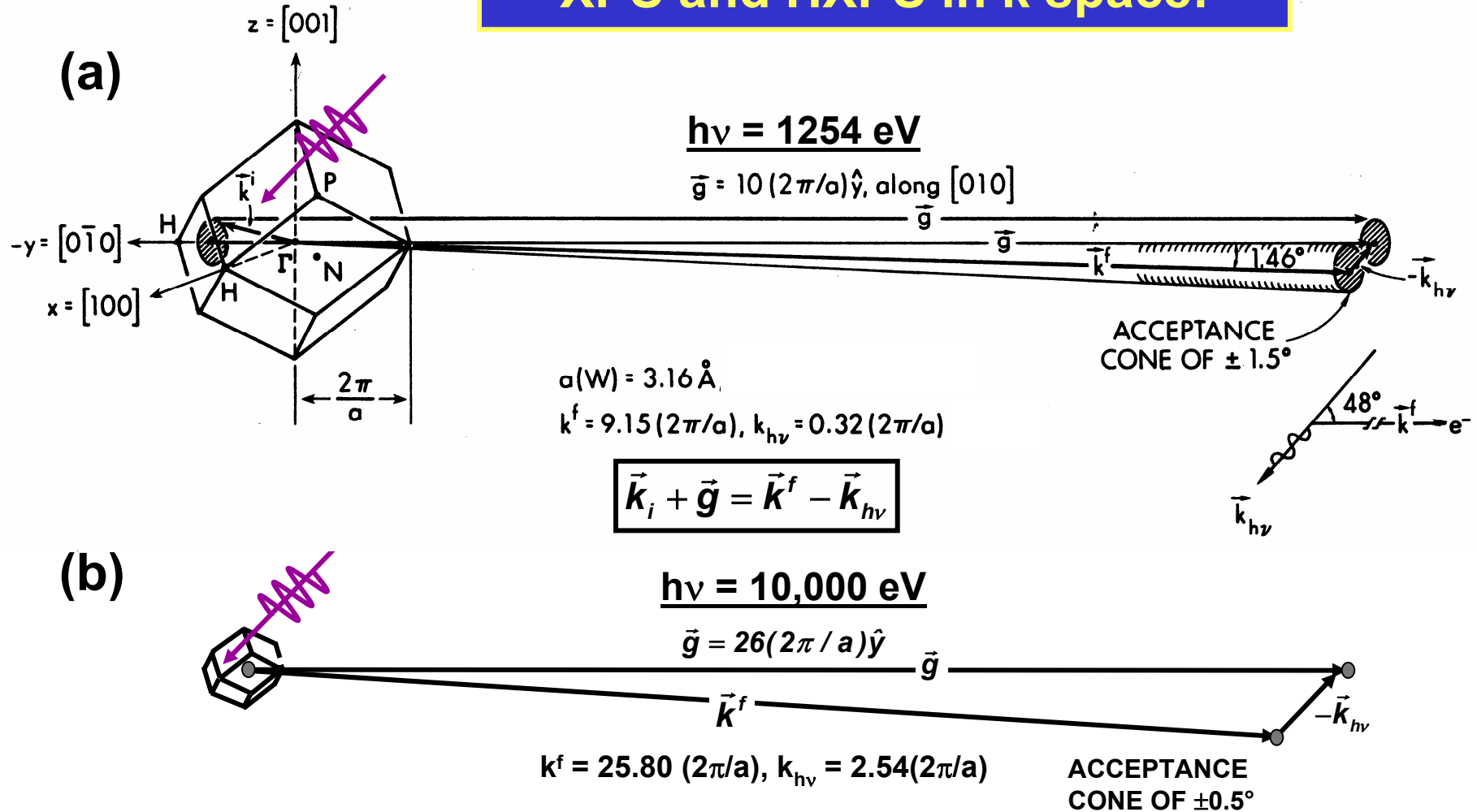


Some classic cases in the XPS limit:



"Basic Concepts of XPS"
Figure 14

XPS and HXPS in k space:

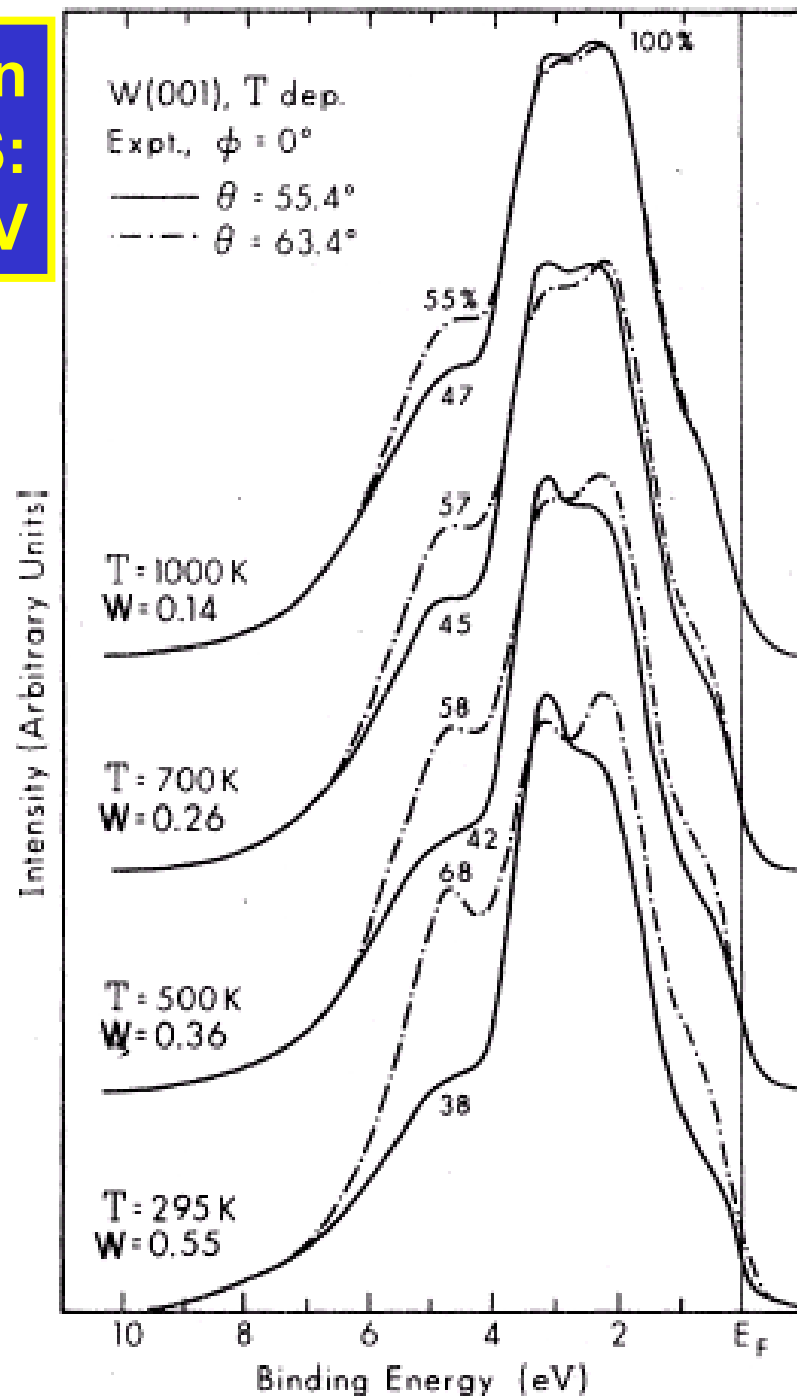


Phonon effects: Approximate fraction of “good” direct transitions

$$\approx \text{Debye-Waller factor} = W(T) \approx \exp[-g^2 \langle u^2(T) \rangle]$$

= Mean-squared
vibrational displacement

**Direct-transition
effects in XPS:
W(110) at 1253.6 eV**

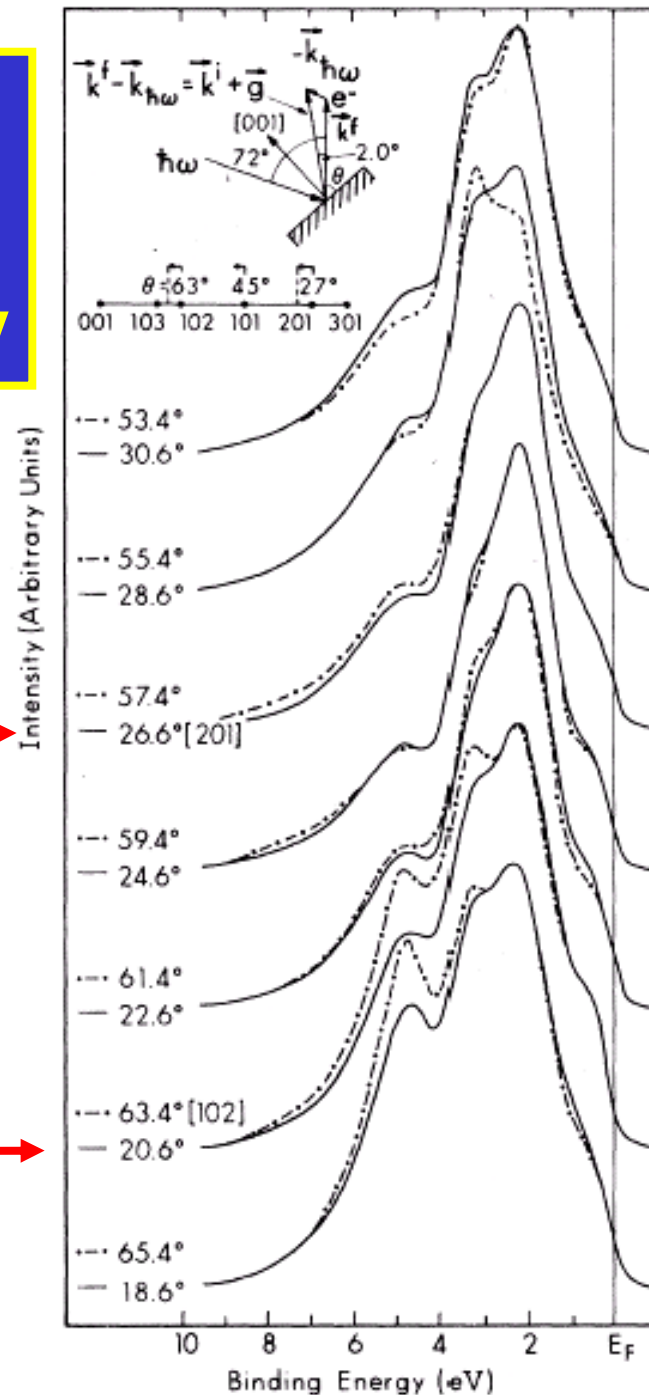


Present if
vibrations stiff
enough (Debye
T high enough),
but suppressed
as temperature is
raised.

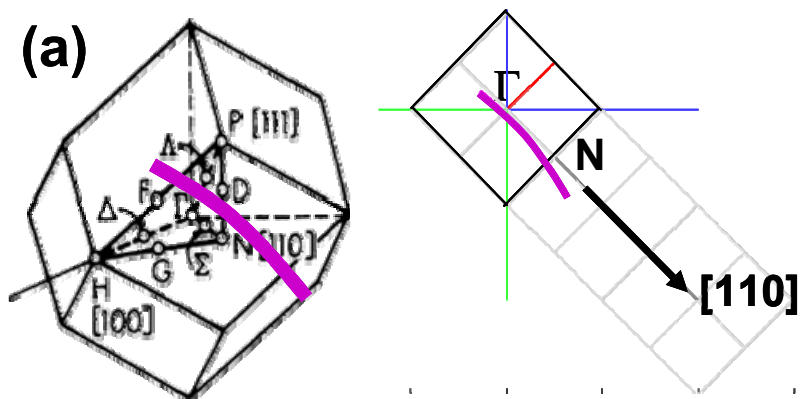
Hussain et al.,
Phys. Rev. 22,
3750 (1980)

**Effect of photon momentum on k conservation:
W(110) at 1253.6 eV**

Symmetry-related spectra shifted by 6.0° for best match. Theoretical 4.8° due to k_{hv}

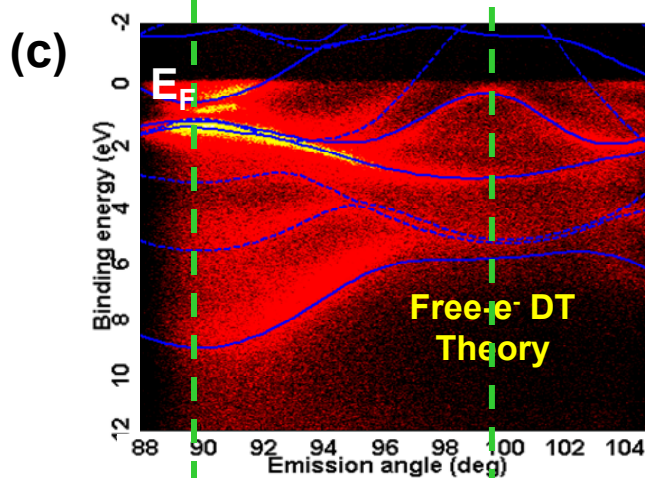
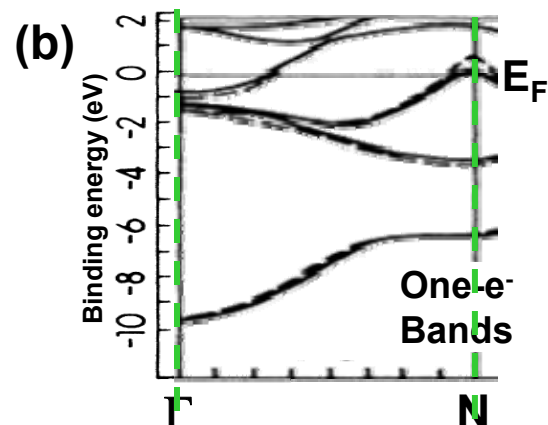


Hussain et al.,
Phys. Rev. 22,
3750 (1980)

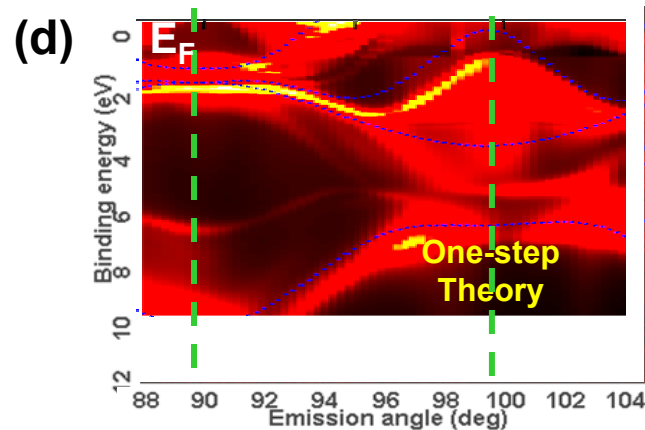


Angle-Resolved
Photoemission
from W(110)
 $h\nu = 260 \text{ eV}$
 $90^\circ = \text{normal}$
 $\Theta_{\text{Debye}} = 400\text{K}$

Plucinski et al., Phys.
Rev. B, submitted

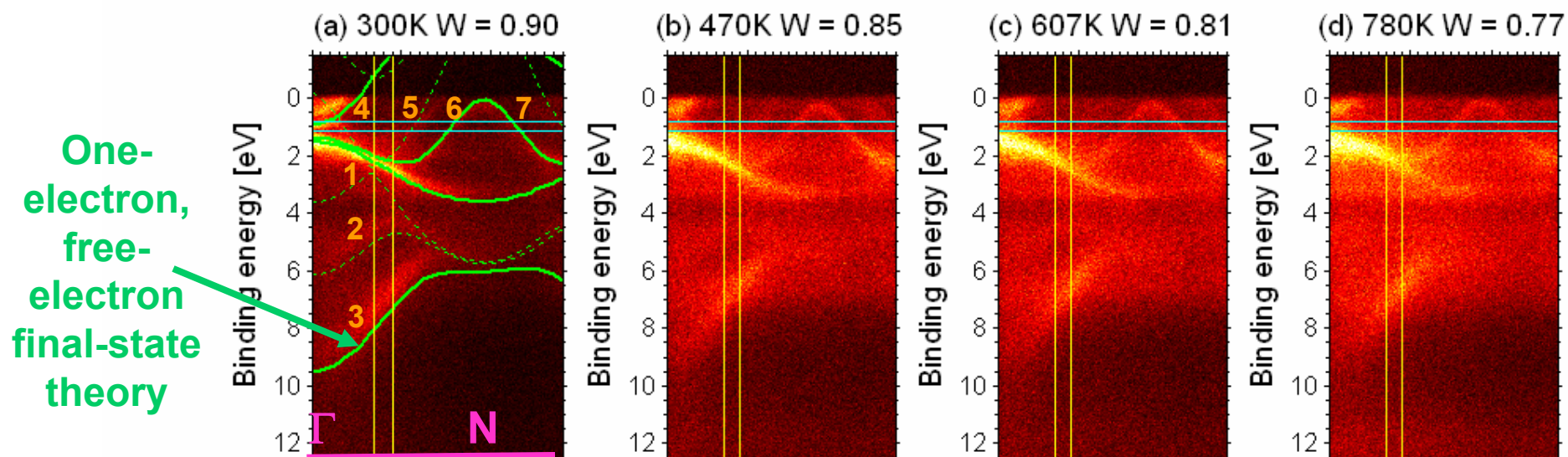


Plucinski

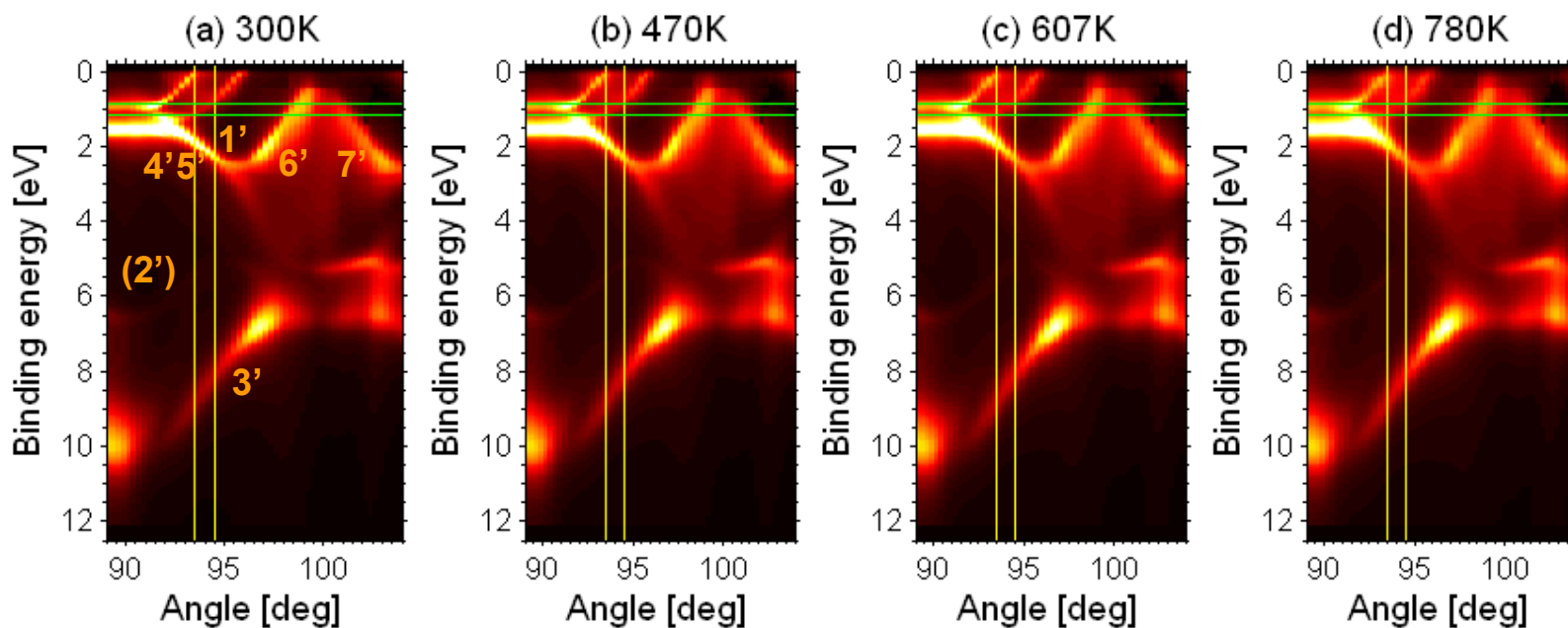


Braun, Minar,
Ebert

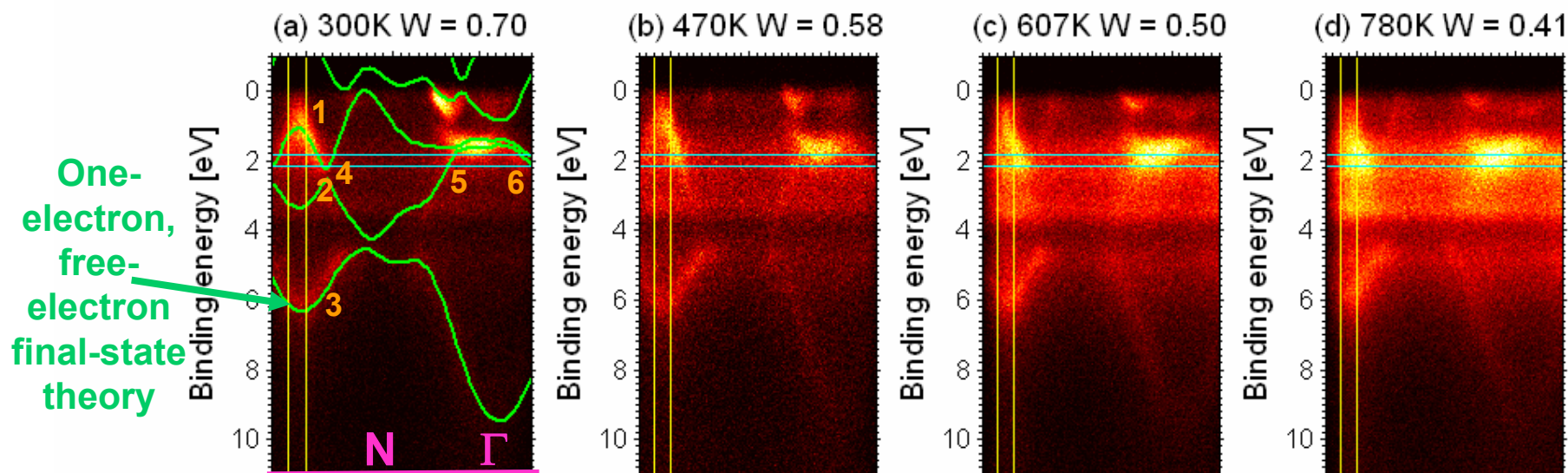
ARPES from W(110): Expt, $h\nu = 260$ eV, near-normal emission, $\Theta_{\text{Debye}} = 400\text{K}$



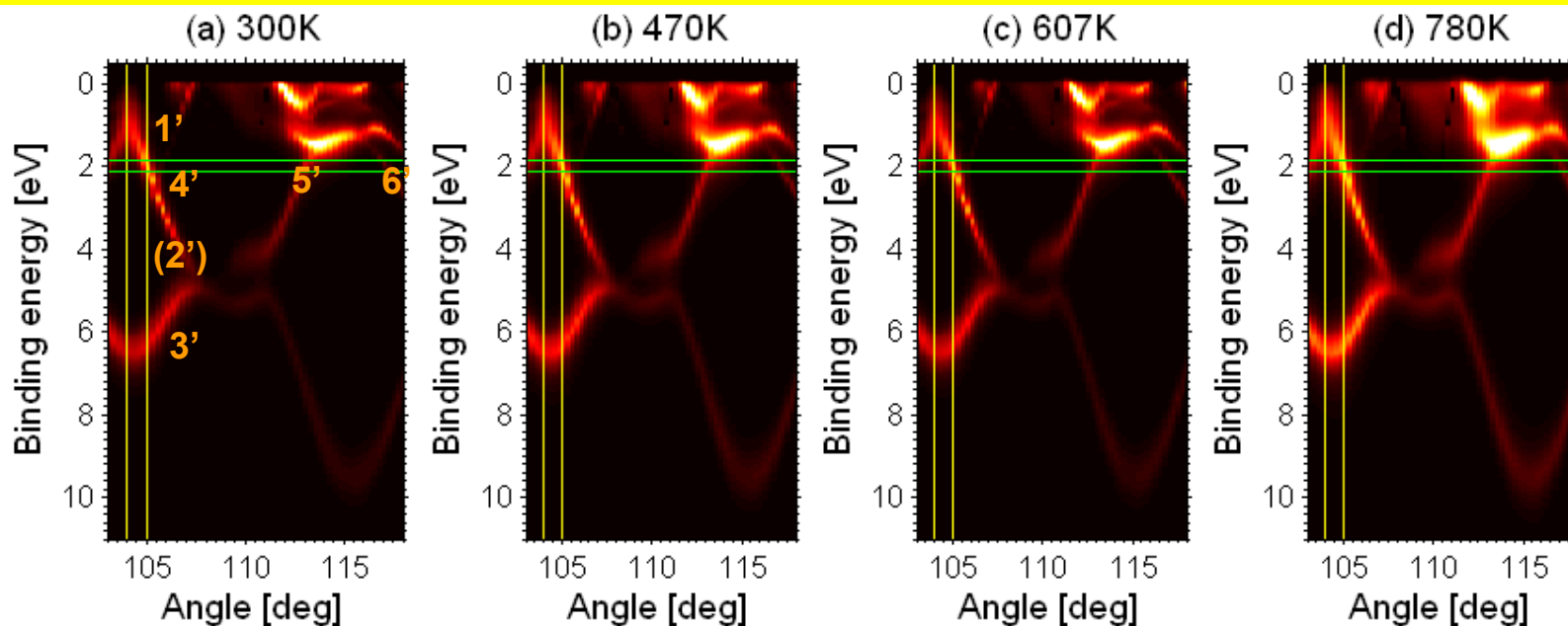
ARPES from W(110): 1-Step Theory, $h\nu = 260$ eV, near-normal emission, $\Theta_{\text{Debye}} = 400\text{K}$



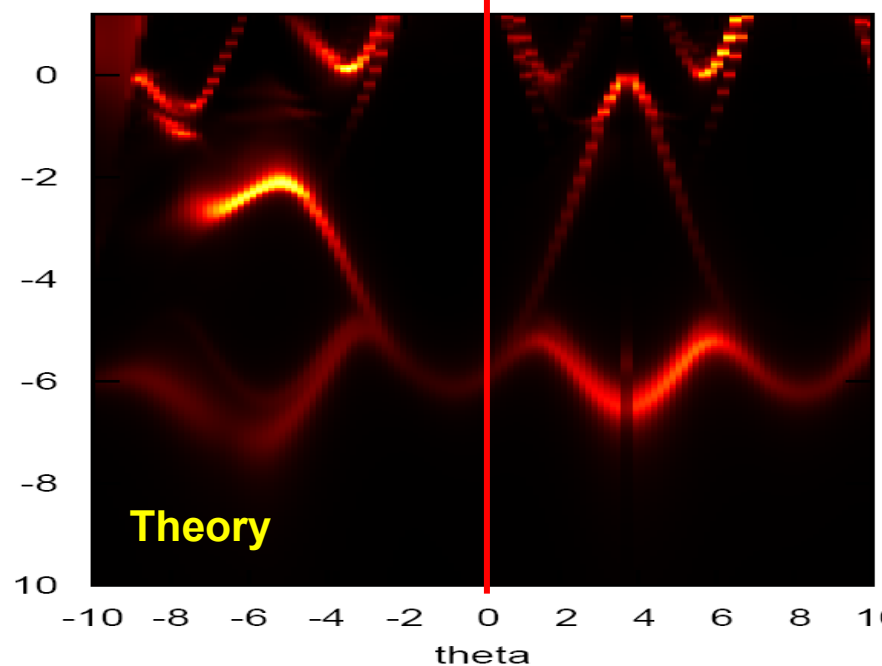
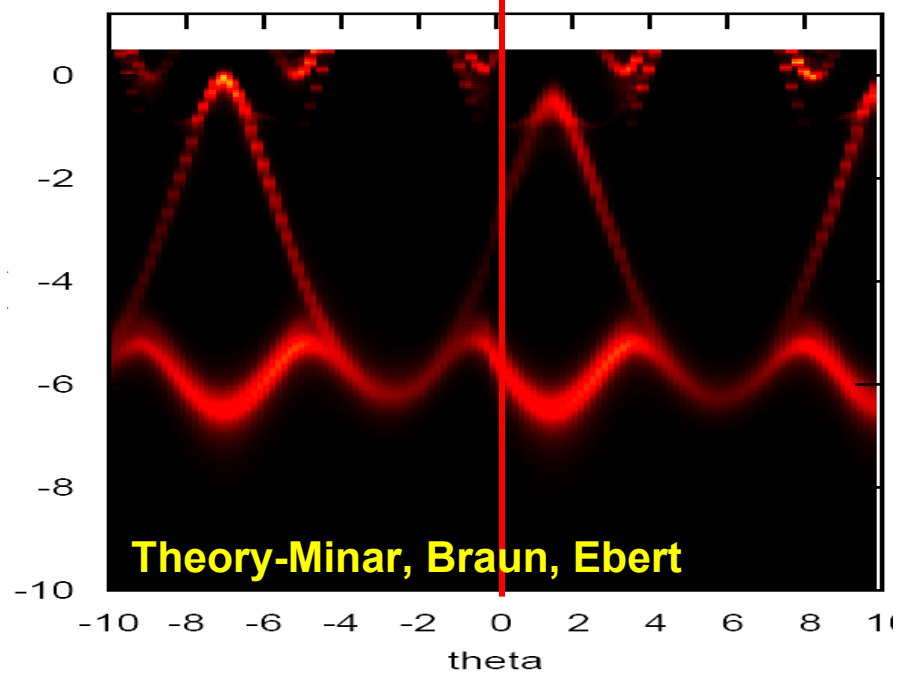
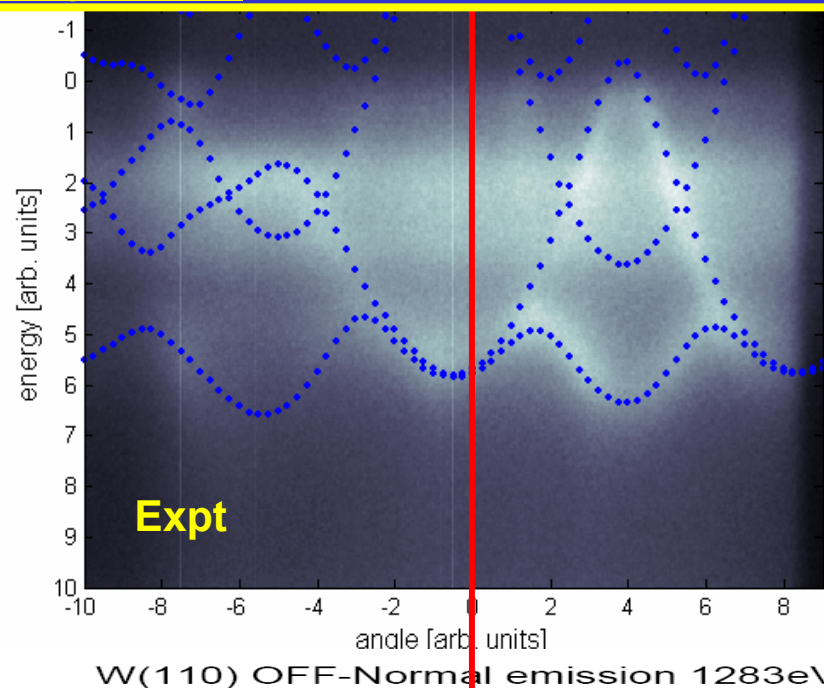
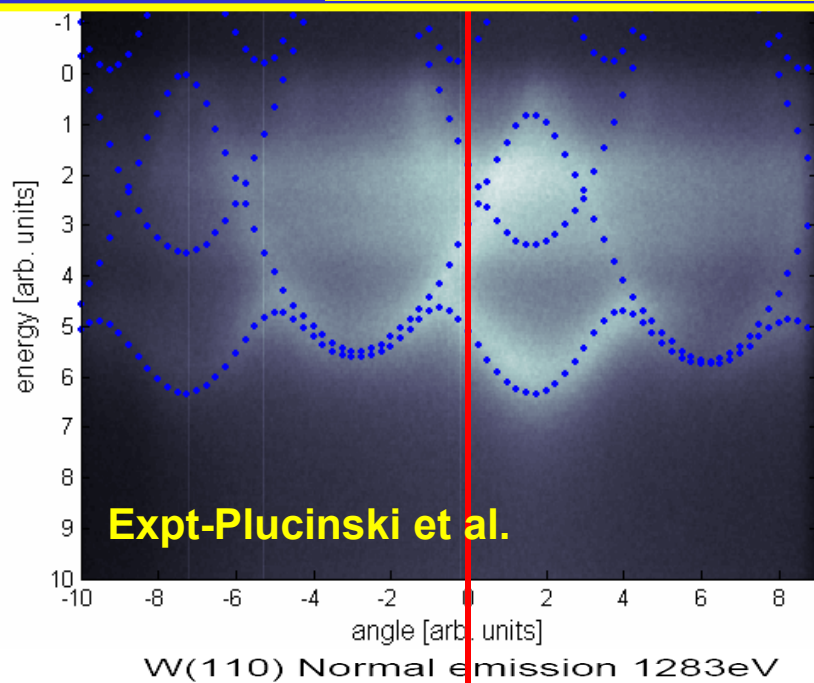
ARPES from W(110): Expt, $h\nu = 860$ eV, near-normal emission, $\Theta_{\text{Debye}} = 400\text{K}$



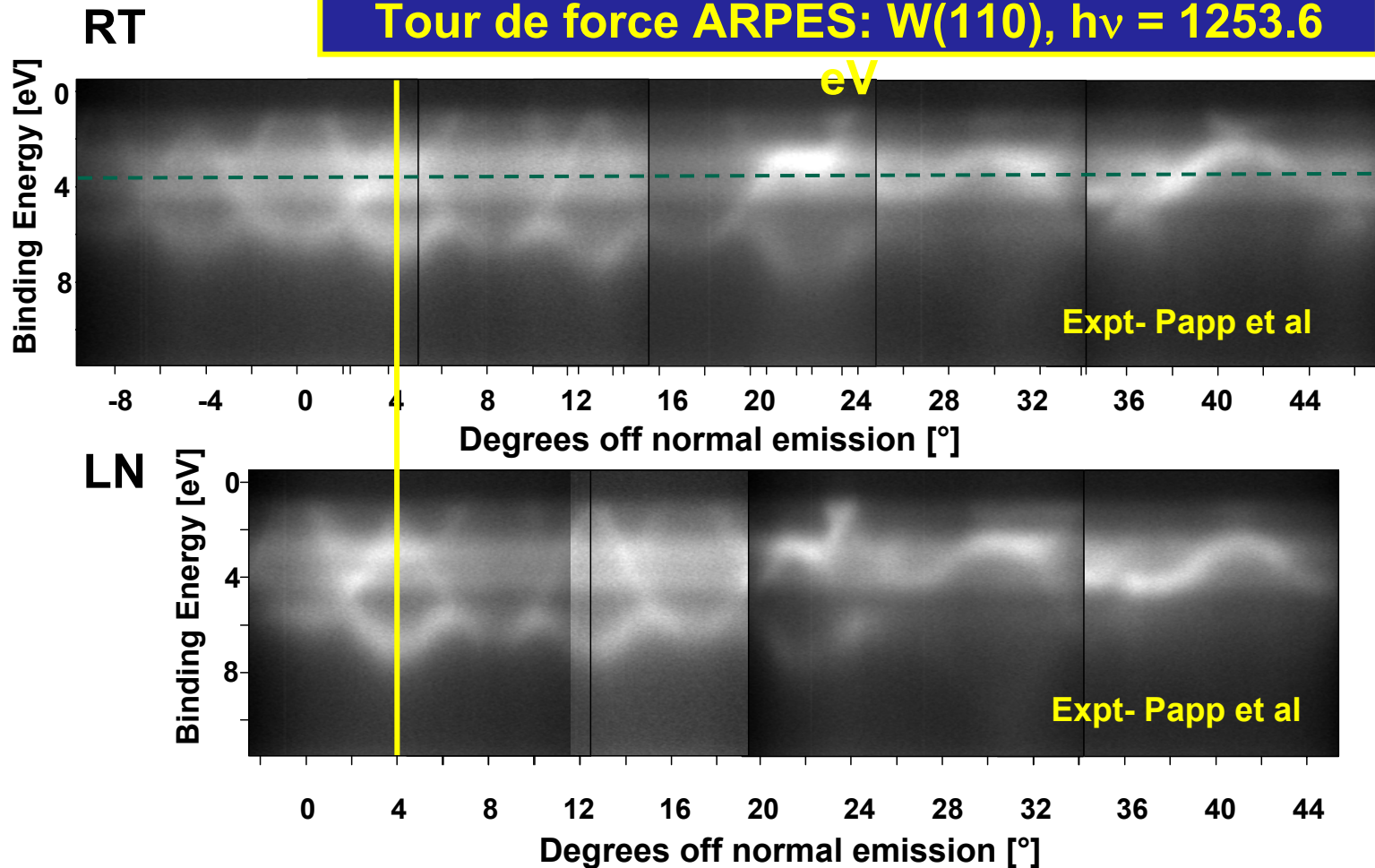
ARPES from W(110): 1-Step Theory, $h\nu = 870$ eV, near-normal emission, $\Theta_{\text{Debye}} = 400\text{K}$



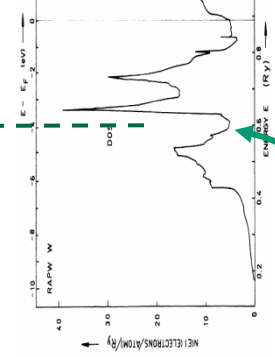
ARPES with a non-monochromatized lab. x-ray source: $h\nu = 1253.6$ eV, $T = \sim 77$ K



Tour de force ARPES: W(110), $h\nu = 1253.6$



Density of States

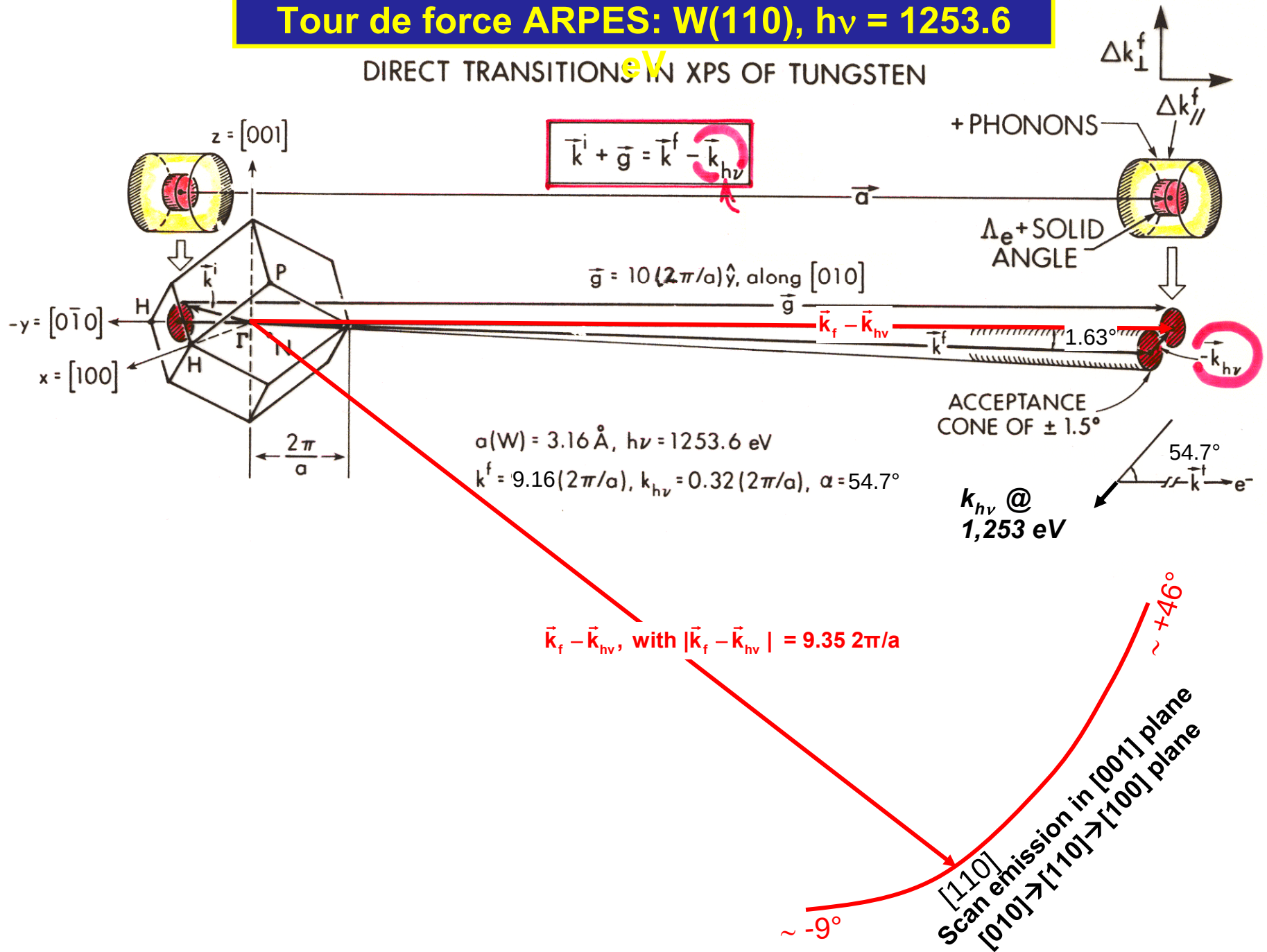


“Pseudo Gap”

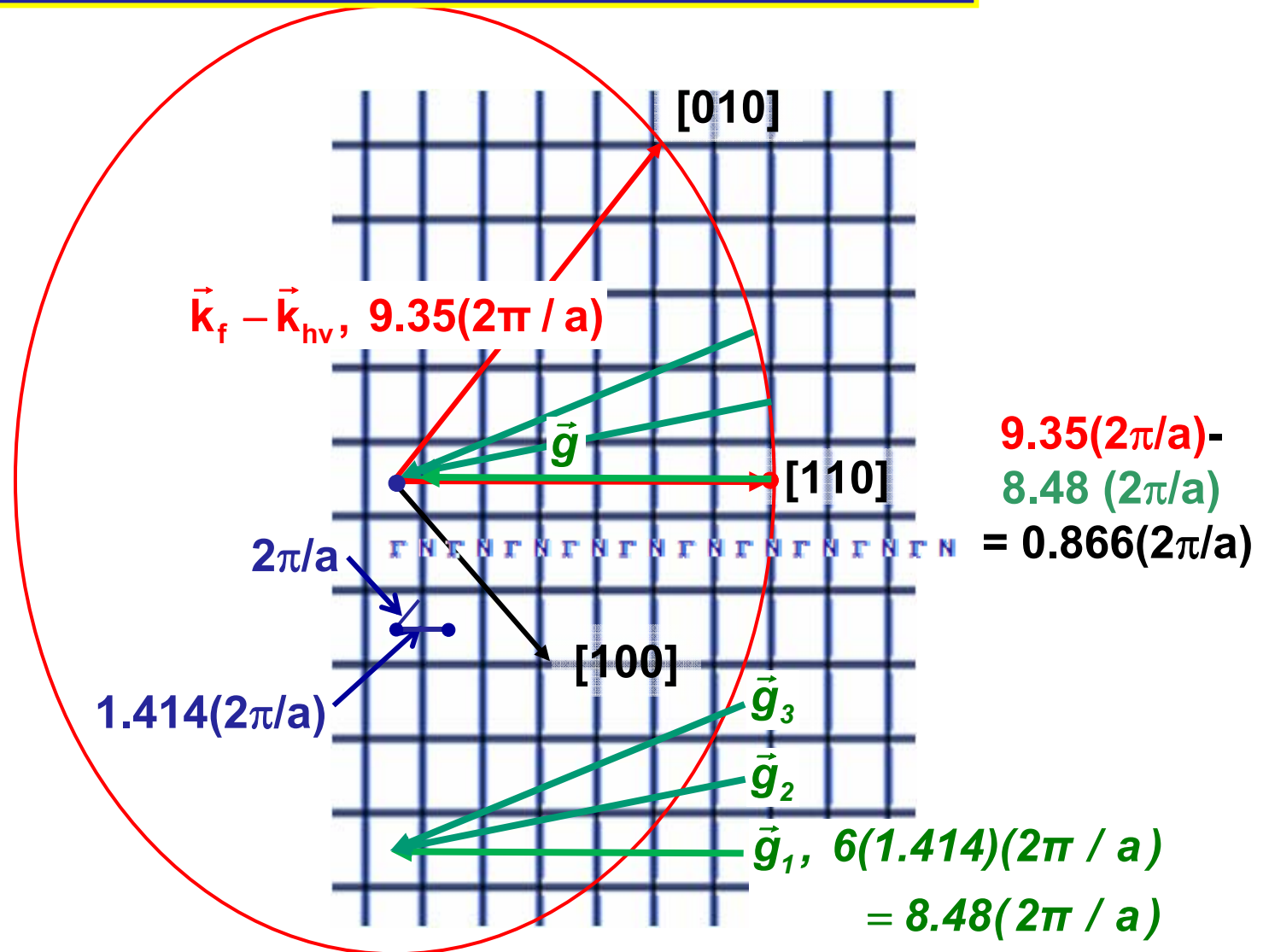
Christensen &
Feuerbacher, PRB
10, 2349 ('74)

Tour de force ARPES: W(110), $h\nu = 1253.6$

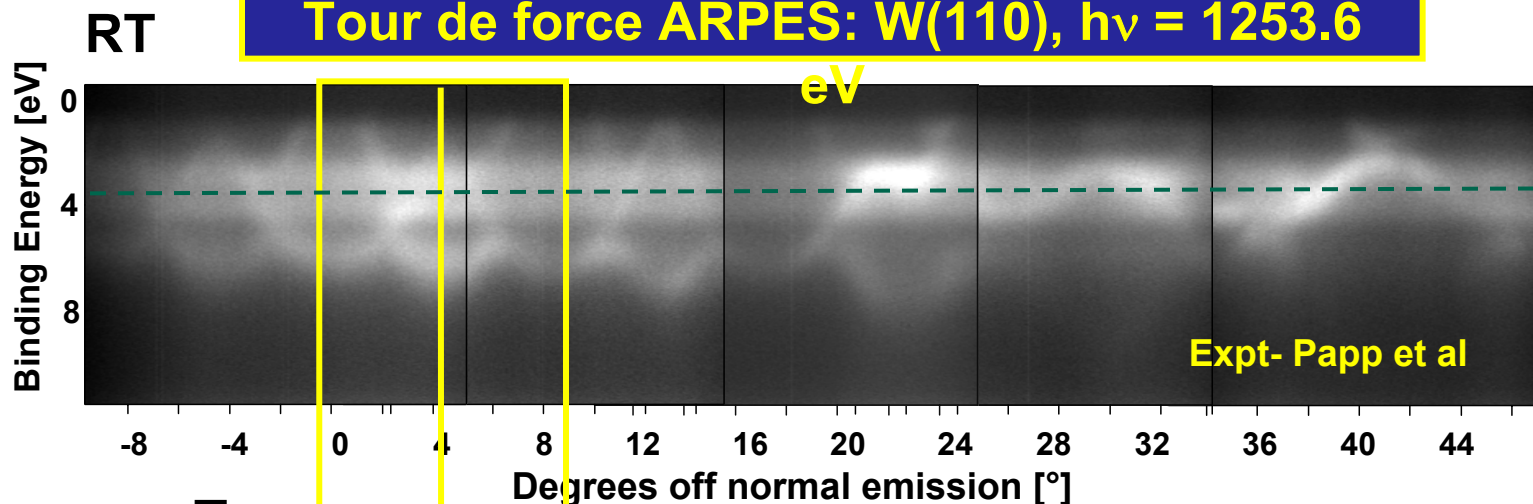
DIRECT TRANSITIONS IN XPS OF TUNGSTEN



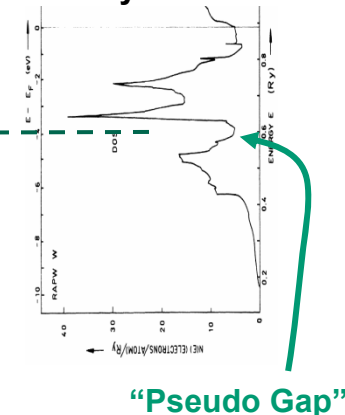
**Tour de force ARPES in an extended zone
scheme: W(110), [001] plane, $h\nu = 1253.6$ eV**



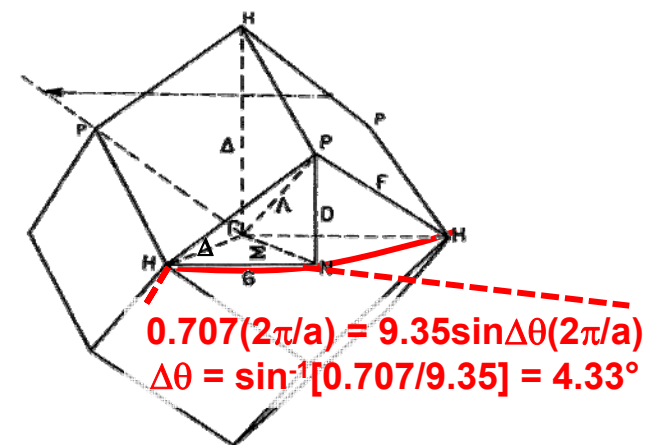
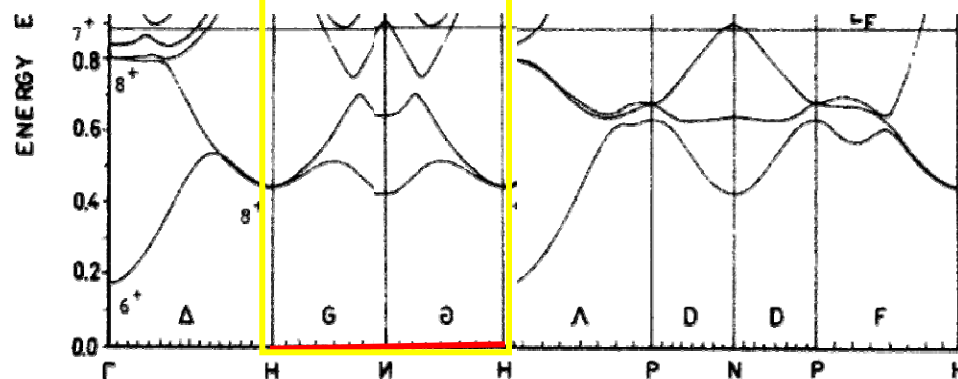
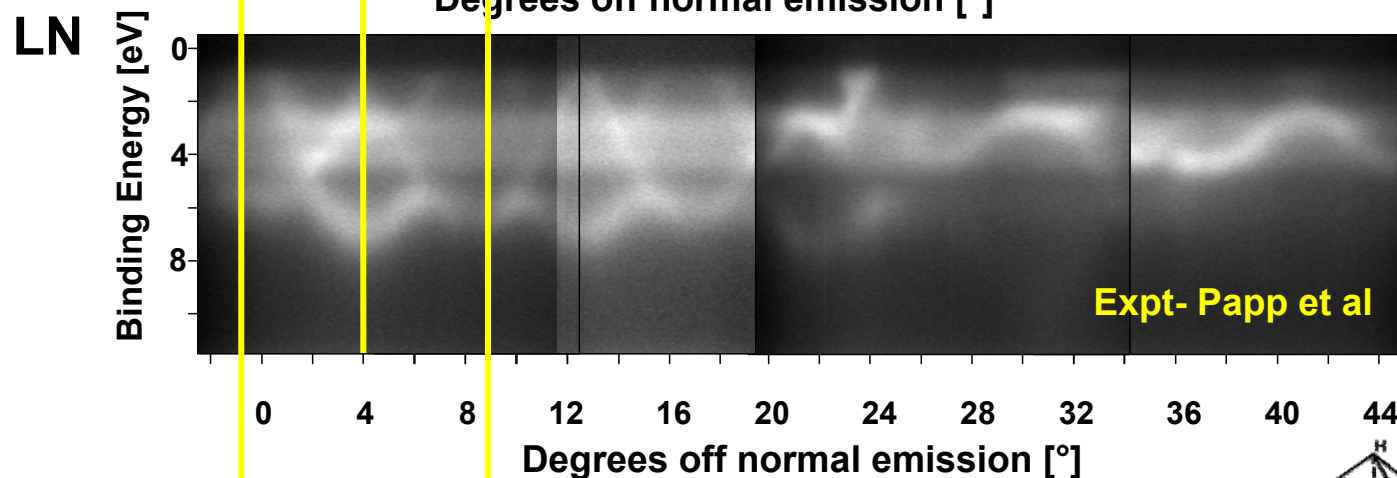
Tour de force ARPES: W(110), $h\nu = 1253.6$



Density of States



Christensen &
Feuerbacher, PRB
10, 2349 ('74)

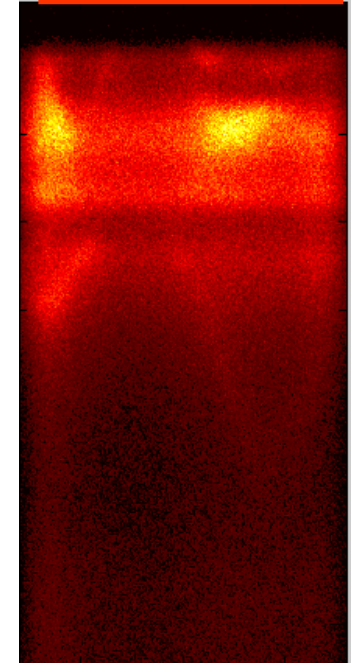
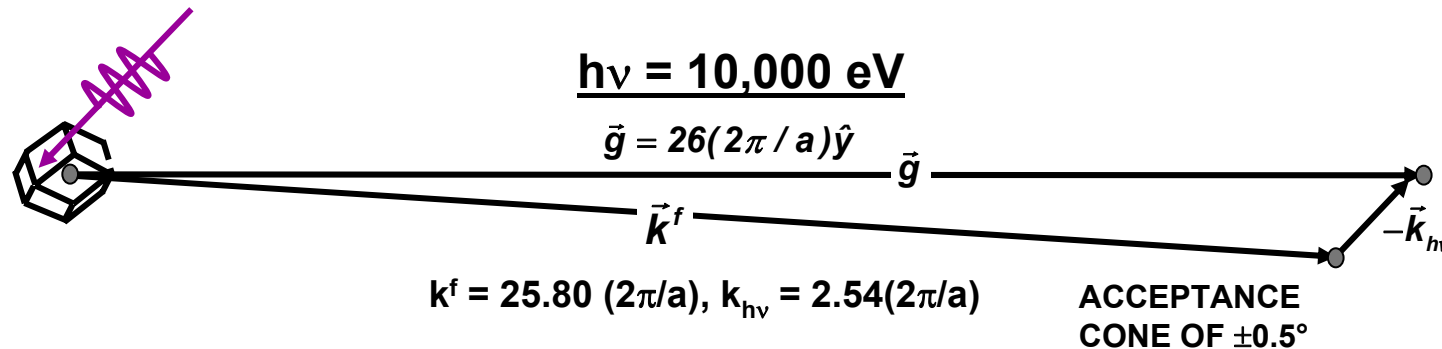


And what would happen at 10 keV?:

$$h\nu = 870 \text{ eV}$$

$$T = 780\text{K}$$

$$W \approx 0.41$$



$$\Theta_{\text{Debye}} = 310\text{K}, \langle u^2 \rangle (10^{-20} \text{ cm}^2) = 5.34 + 0.0583T \xrightarrow{\text{High } T} 0.0583T$$

$$\text{Debye-Waller Factor} = W(T) \approx \exp(-k_e^2 \langle u^2(T) \rangle)$$

$$= \exp(-C_1 E_{\text{kin}} \langle u^2(T) \rangle) \xrightarrow{\text{High } T} \exp(-C_2 E_{\text{kin}} T)$$

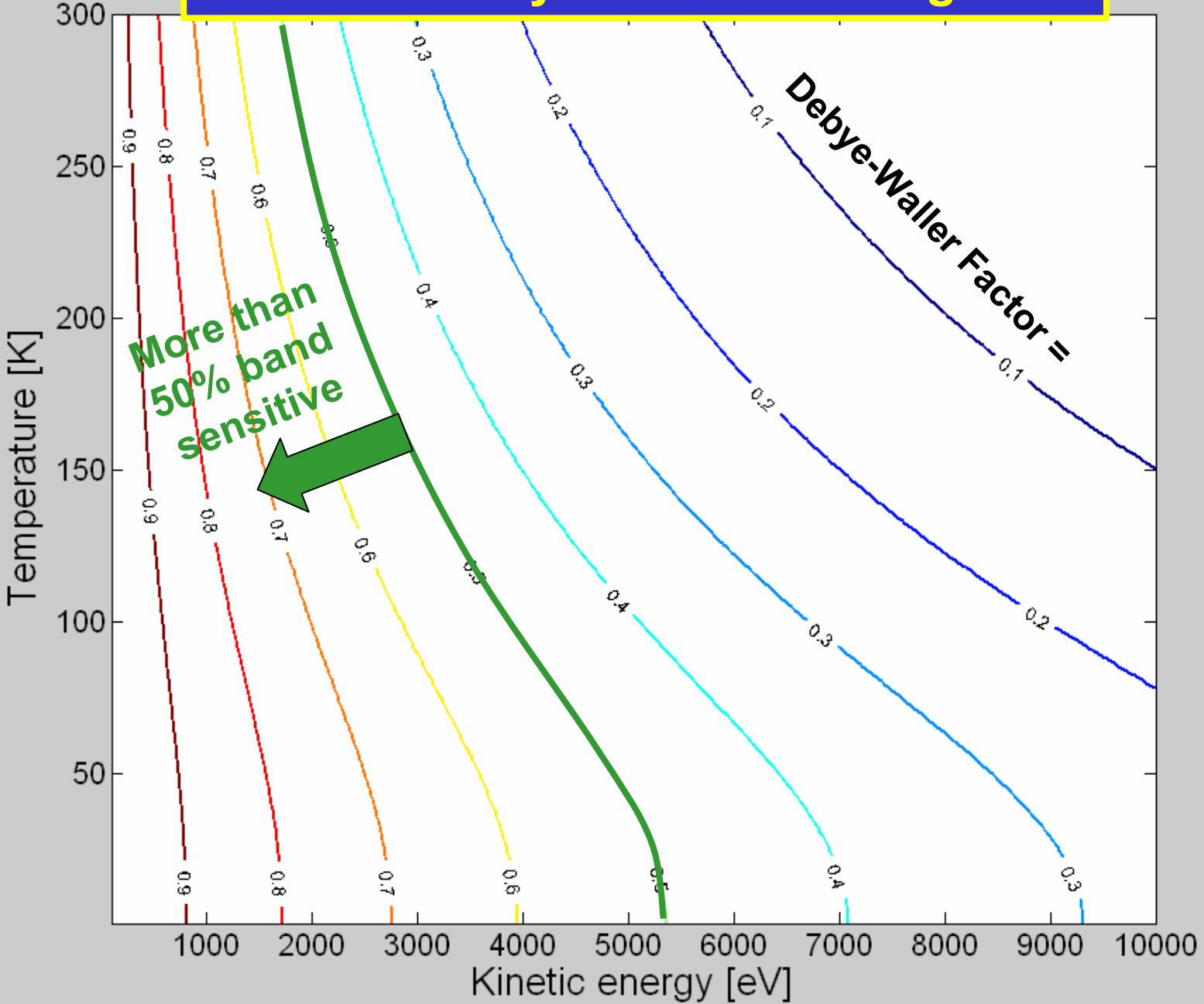
$$W \text{ at } 4\text{K}: W \approx 0.27, \approx 27\% \text{ direct}$$

$$\text{at } 77\text{K}: W \approx 0.20, \approx 20\% \text{ direct}$$

$$\text{at } 300\text{K}: W \approx 0.017, \approx 2\% \text{ direct}$$

Correlated vibrations and better theory (e.g. Phys. Rev. B 35, 1147 ('87) and 53, 7524 ('96) + 54, 14703 ('96)) may yield different DT percentages, but needs further experimental and theoretical study

Quantitative systematics—Tungsten :



Outline

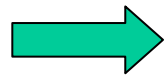
Surface, interface, and nanoscience—short introduction

Some surface concepts and techniques→photoemission

Synchrotron radiation: experimental aspects

Electronic structure—a brief review

**The basic synchrotron radiation techniques:
more experimental and theoretical details**



Core-level photoemission

Valence-level photoemission

Microscopy with photoemission

Outline

- Valence-band spectra: low-energy UPS limit and high-energy XPS limit

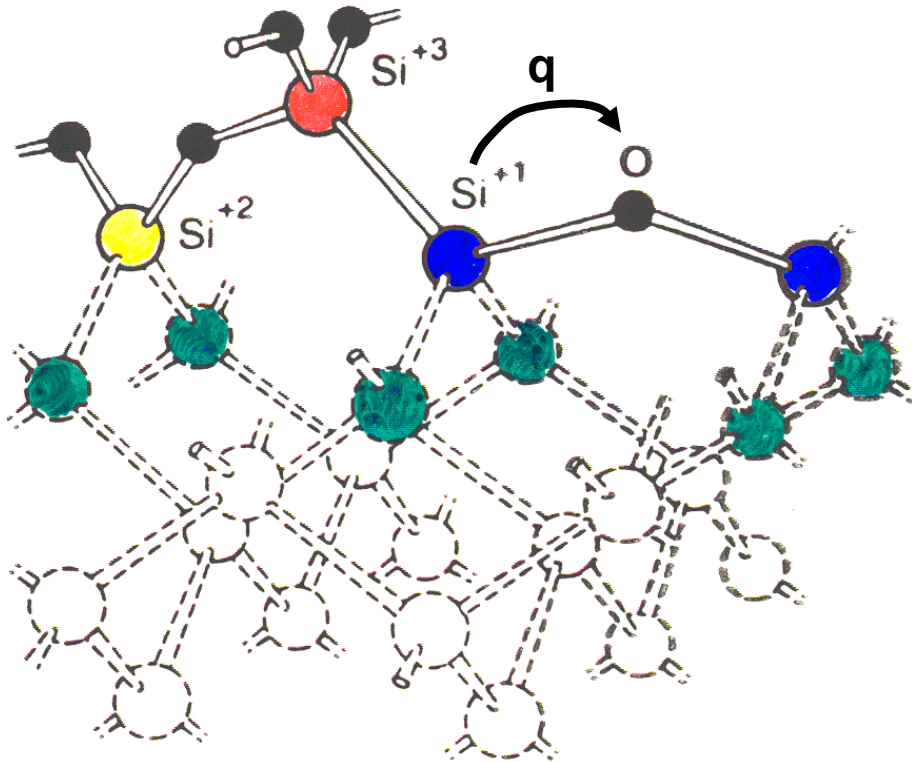


- Core-level chemical shifts: the potential model
- Core-level chemical shifts: equivalent-core ($Z+1$) and thermochemical energies
- Multiplet splittings
- Spin-orbit splitting, the Fano effect, and spin-polarized outgoing electrons
- Magnetic circular dichroism (MCD) in core-level emission
- Non-magnetic circular dichroism in core-level emission: a.k.a. circular dichroism in angular distributions (CDAD)
- Various other final state effects providing information in core-level spectra

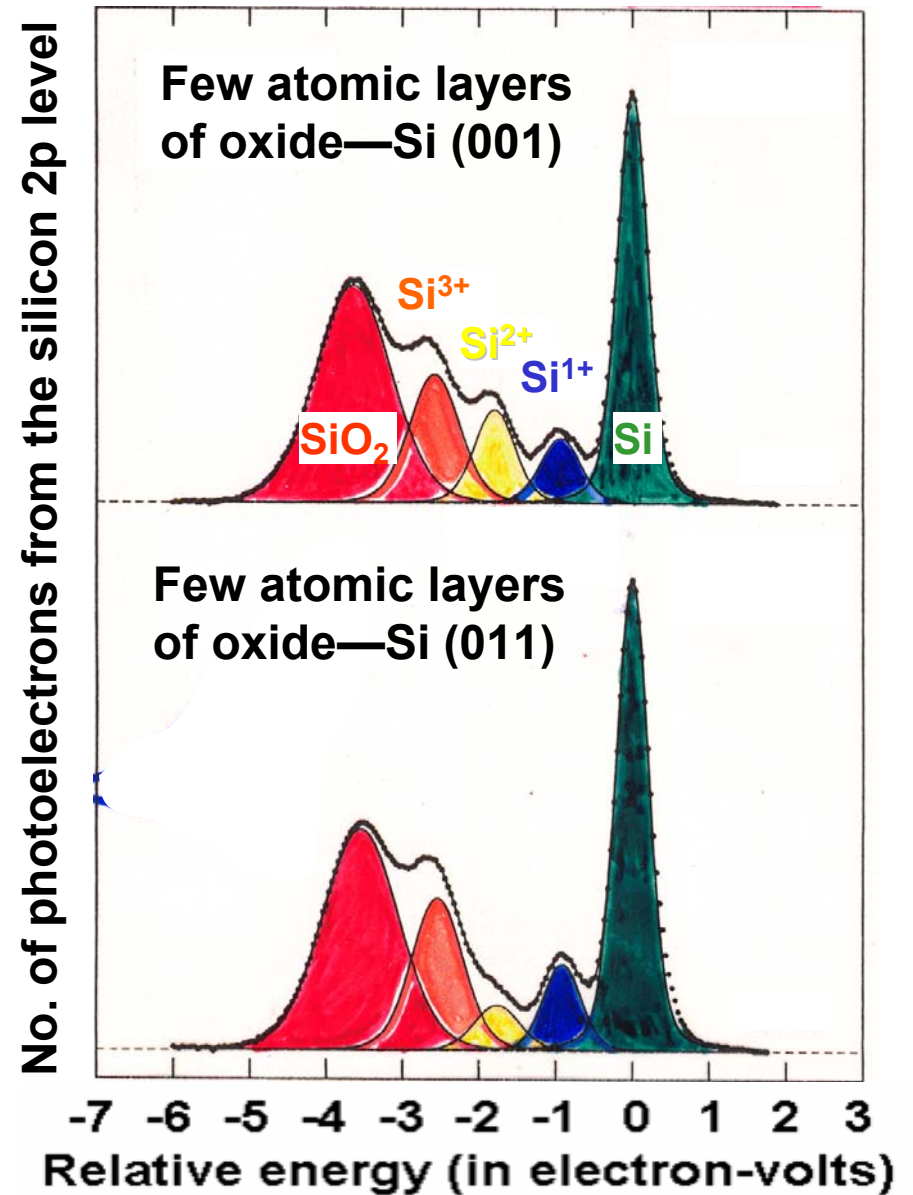
Looking into the silicon dioxide layer with photoelectron spectroscopy

Charge transfer, e^- - e^- coulomb integral:

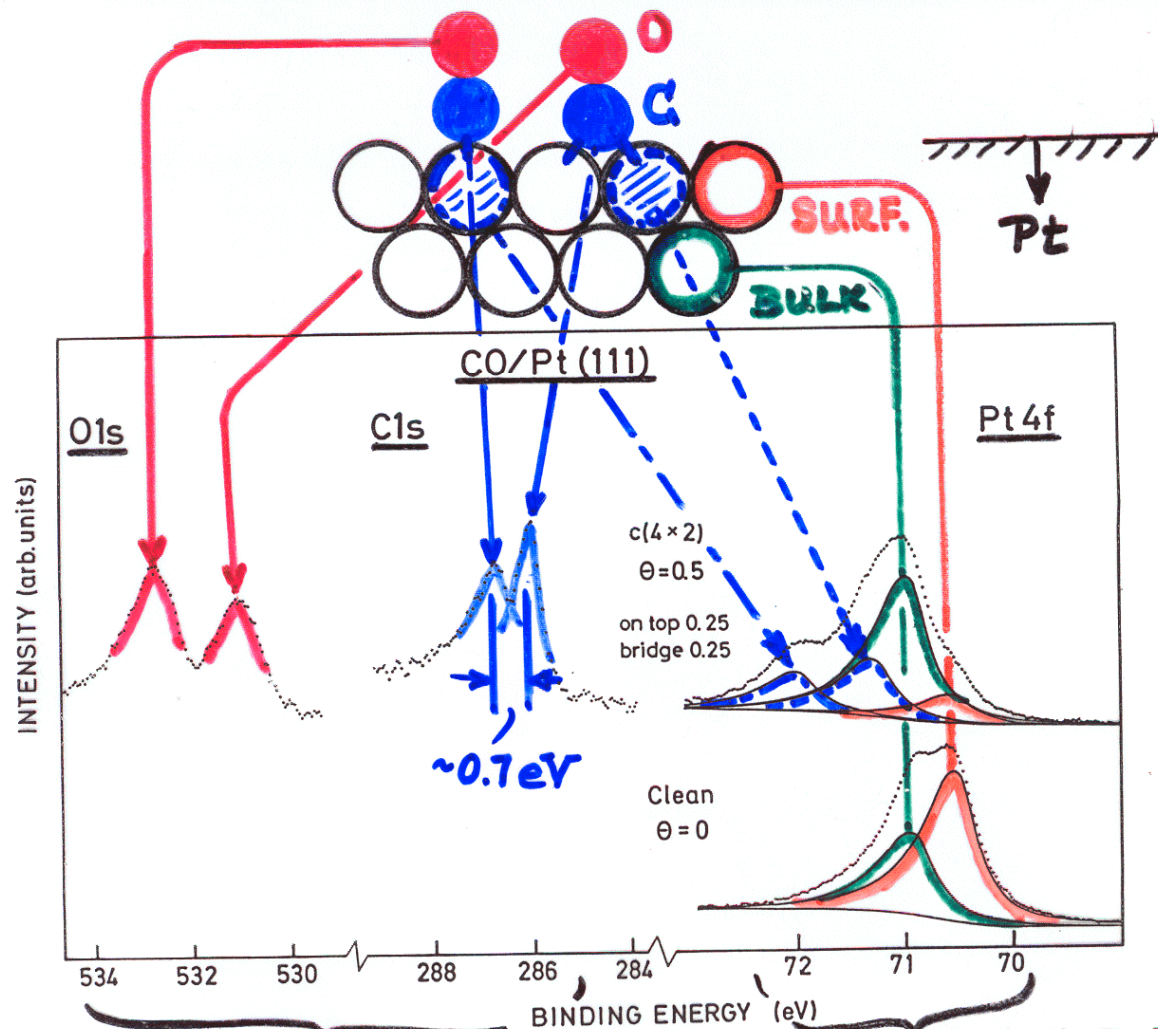
$$\text{Shift} \approx q_{\text{Si}} J_{\text{Si}2p, \text{Si}3p} = \int \varphi_{2p}^*(\vec{r}_1) \varphi_{3p}^*(\vec{r}_2) \frac{e^2}{r_{12}} \varphi_{2p}(\vec{r}_1) \varphi_{3p}(\vec{r}_2) dV_1 dV_2$$



Himpsel et al., Phys. Rev. B 38, 6086 ('88)

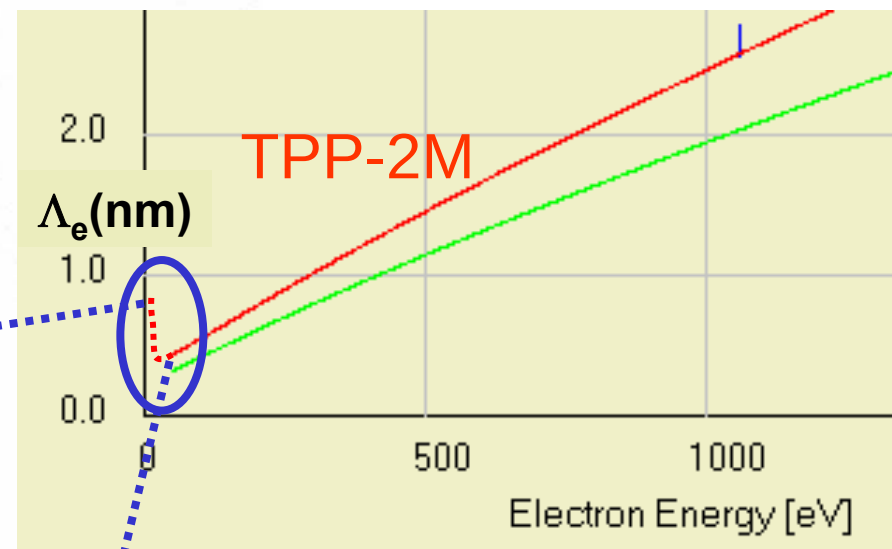
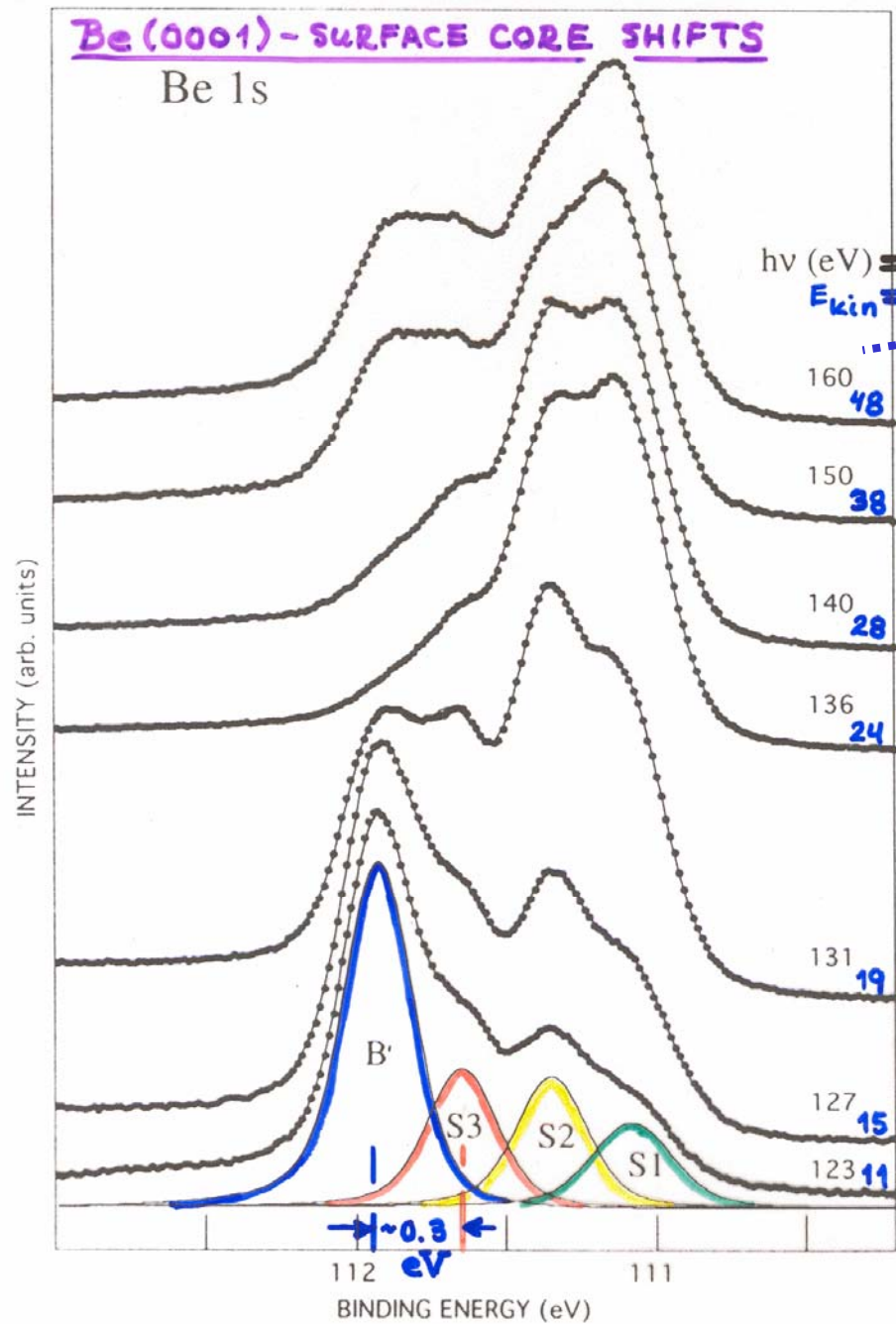


CHEMICAL SHIFTS IN ADSORBATE & SUBSTRATE

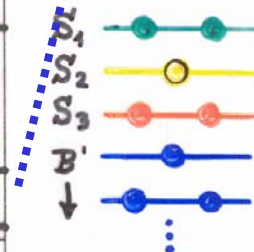


MONOCH. XPS,
 $\Delta E \sim 0.3 \text{ eV}$

SYNCH. RAD., $\Delta E \sim 0.15 \text{ eV}$
BJÖRNEHOLM
ET AL.
(NILSSON GRP.,
UPPSALA)



MINIMUM
IN e^-
ESCAPE
DEPTH



JOHANSSON ET.
AL., P.R.L.
71, 2453 (1993)

What does the hole do?

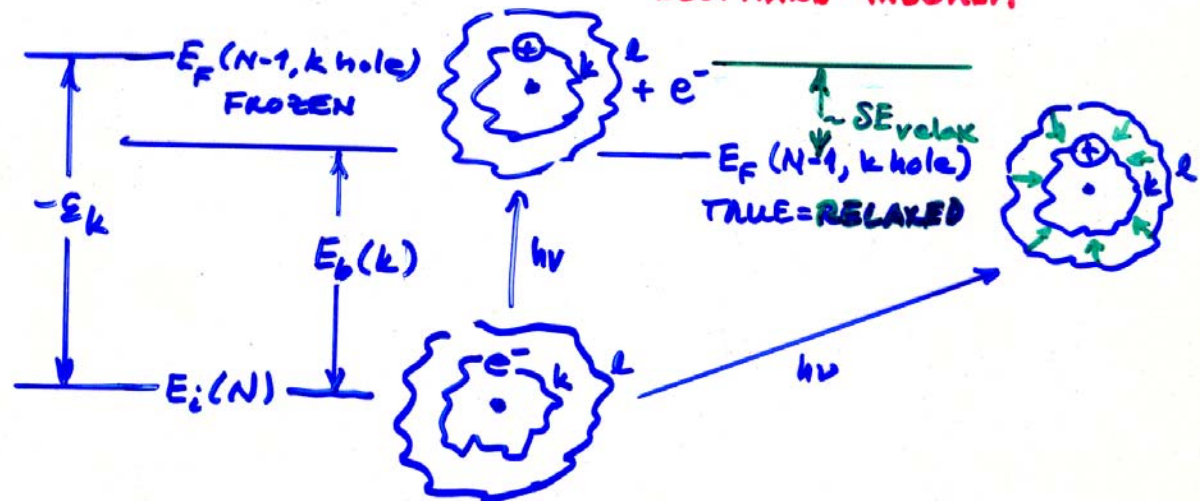
BINDING ENERGIES + KOOPMANS' THEOREM:

N - e^- SCH. EQN. — $\hat{H}(N)\Psi_j(N) = E_j(N)\Psi_j(N)$, $j=1,2,\dots$
 MINIMIZE $E_j(N)$ $\left\{ \begin{array}{l} \Psi_j \approx \Phi_j = \text{SLATER DET.} \end{array} \right.$

N 1- e^- HARTREE-FOCK EQNS. — $\hat{H}(1)\varphi_k(1) = \varepsilon_k(1)\varphi_k(1)$
 • COUPLED INTEGRO-DIFF.
 • COULOMB + EXCHANGE

$E_b(k) = k^{\text{th}} \text{ BINDING ENERGY} = E_f(N-1, k \text{ hole}) - E_i(N)$
 (+) EXACT
 OR $E_b(k) = -\varepsilon_k$ IF $\varphi_{ki} = \varphi_{kf}$ (FROZEN ORBITAL)

KOOPMANS' THEOREM



⇒ RELAXATION, SCREENING, CONFIGURATION INTERACTION, SELF-ENERGY EFFECTS ALWAYS PRESENT; ANDERSON IMPURITY MODEL ETC.

KOOPMANS' THEOREM CALCULATION OF SHIFTS

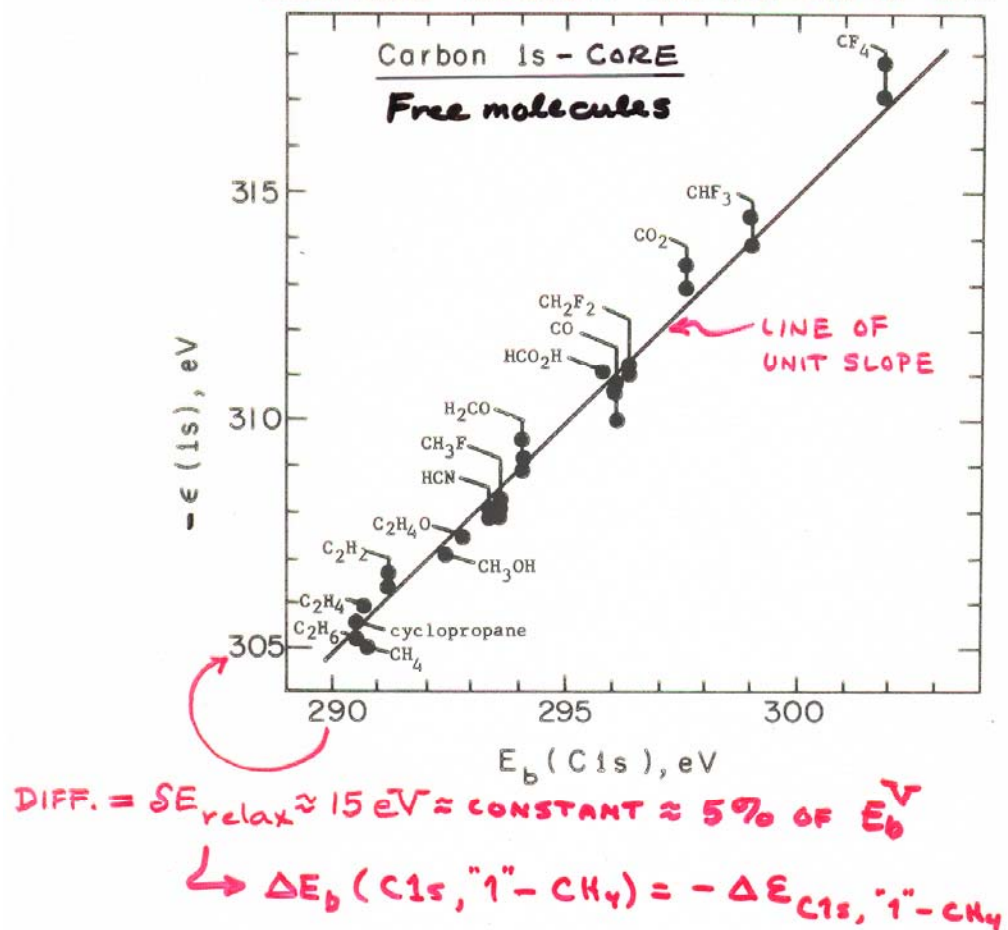
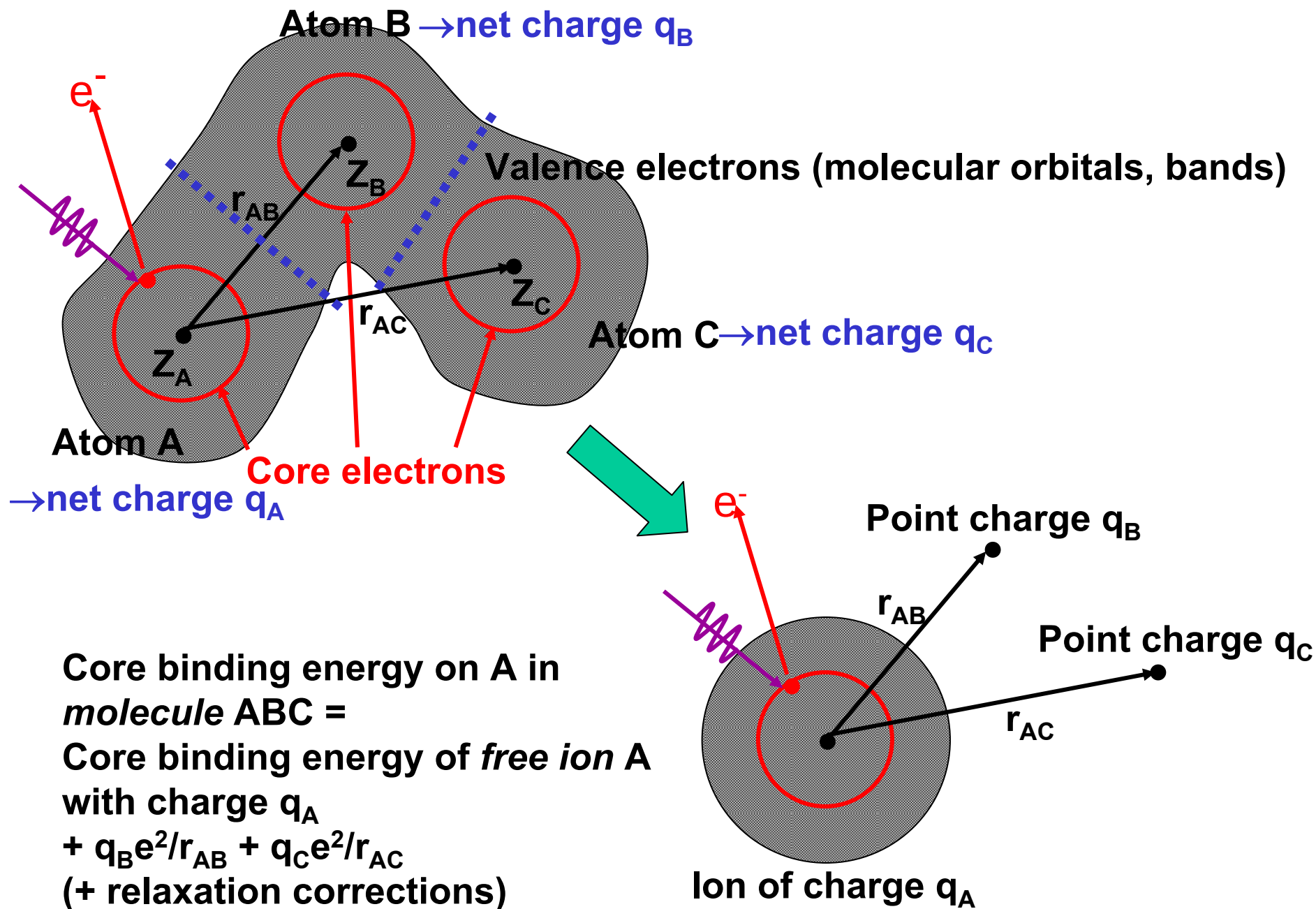
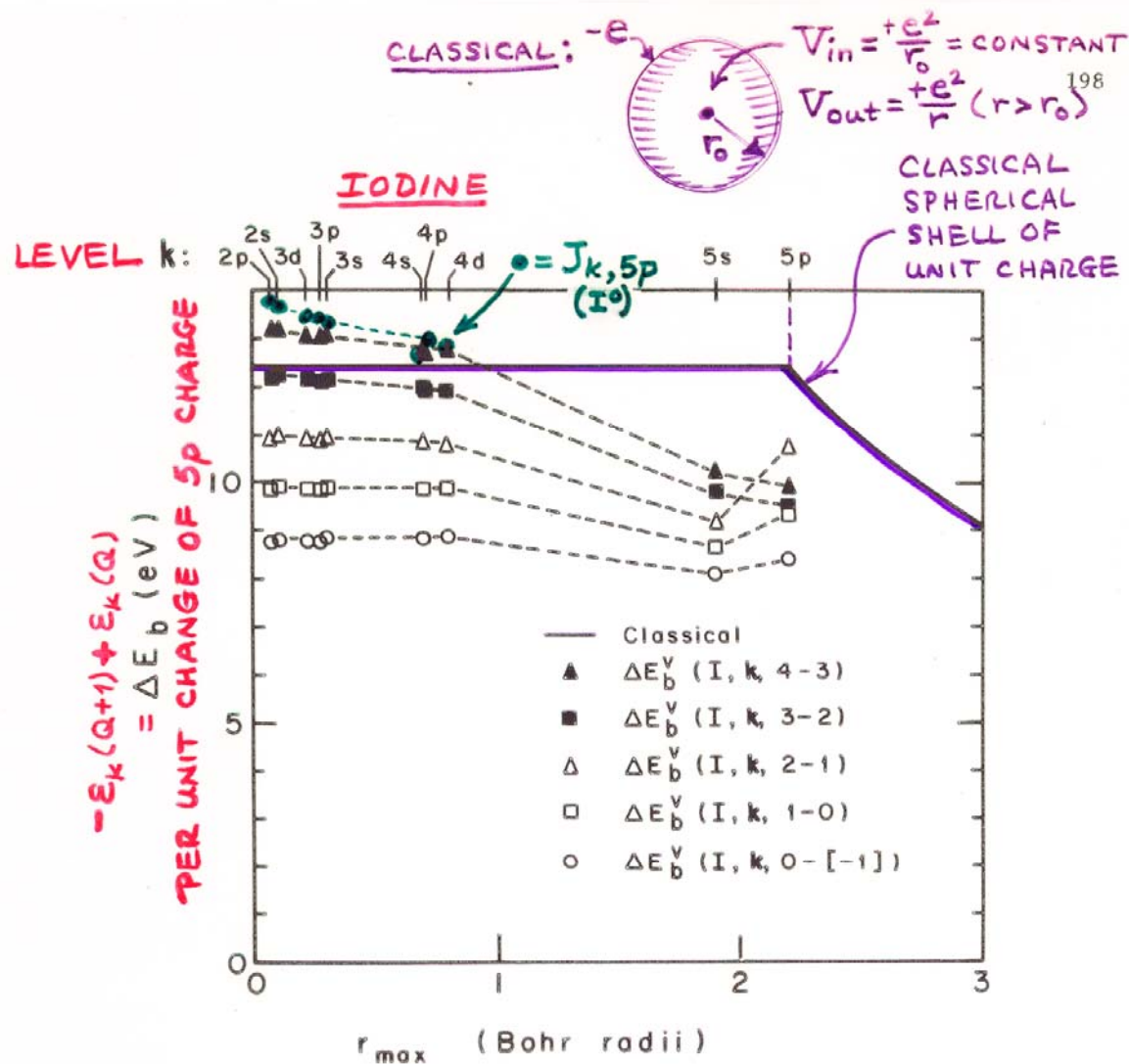


Figure 18 -- Plot of carbon 1s binding energies calculated via Koopmans' Theorem against experimental binding energies for several carbon-containing gaseous molecules. For some molecules, more than one calculated value is presented. The slope of the straight line is unity. The two scales are shifted with respect to one another by 15 eV, largely due to relaxation effects. All of the theoretical calculations were of roughly double-zeta accuracy or better. (From Shirley, reference 7.)

POTENTIAL MODEL FOR CORE-LEVEL CHEMICAL SHIFTS



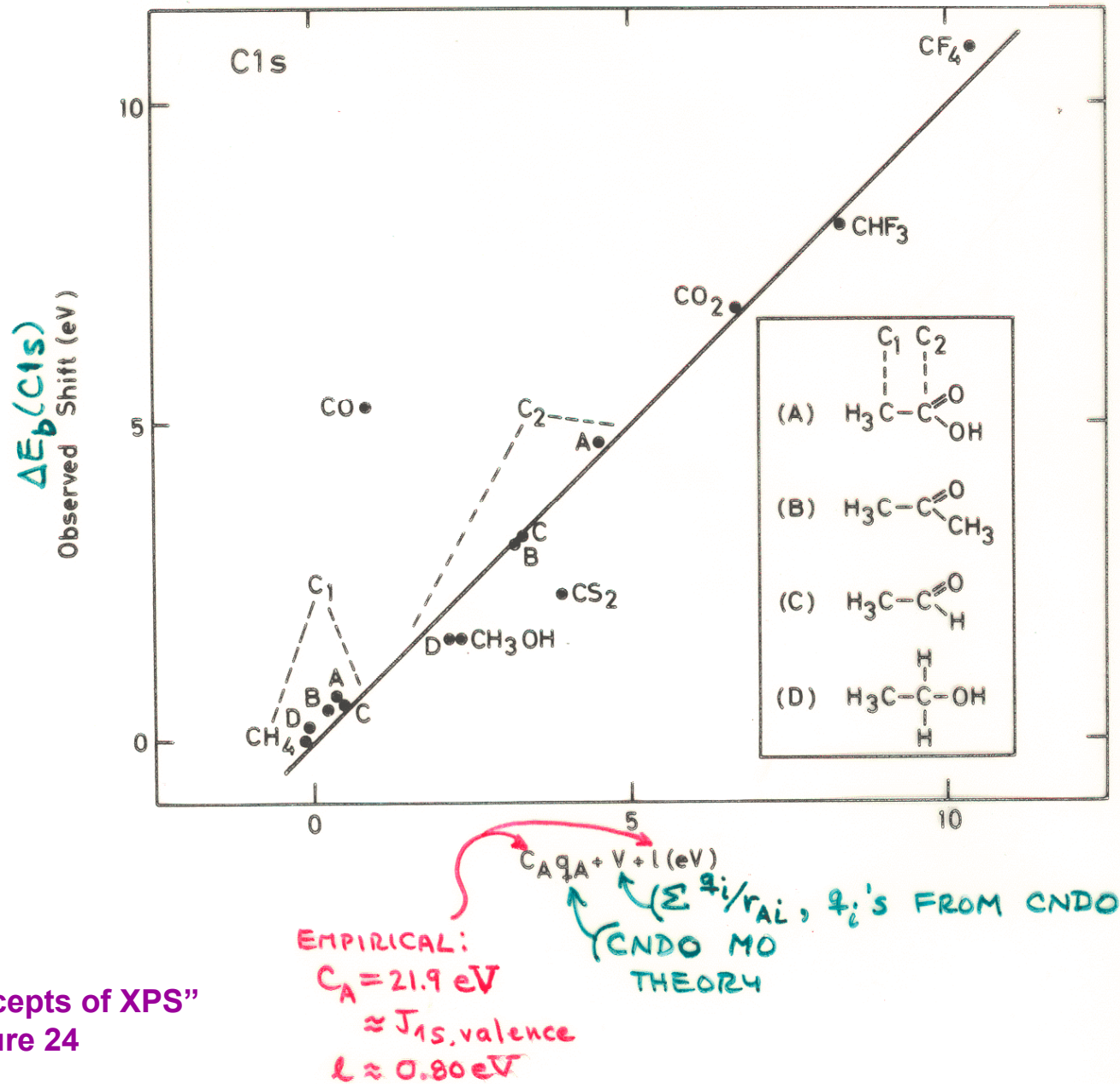
FREE-ION (INTRAAATOMIC) ASPECTS OF SHIFTS: Koopmans' Theorem & Classical Charged Shell



⇒ REMOVAL/ADDITION OF VALENCE e^- CHARGE IN BONDING SHIFTS ALL INNER e^- E_b 's $\approx E_c$'s BY SAME AMOUNT

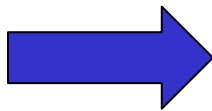
“Basic Concepts of XPS”

POTENTIAL MODEL CALCULATION OF CARBON CHEMICAL SHIFTS



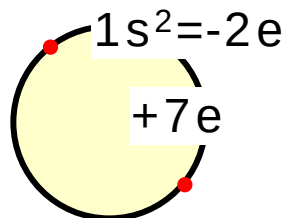
“Basic Concepts of XPS”
Figure 24

Outline

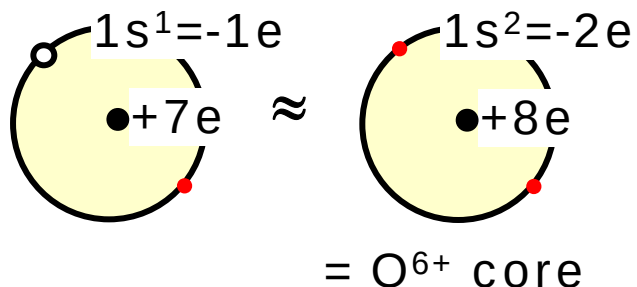
- Valence-band spectra: low-energy UPS limit and high-energy XPS limit
- Core-level chemical shifts: the potential model
-  • Core-level chemical shifts: equivalent-core ($Z+1$) and thermochemical energies
- Multiplet splittings
- Spin-orbit splitting, the Fano effect, and spin-polarized outgoing electrons
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- Non-magnetic circular dichroism in core-level emission: a.k.a. circular dichroism in angular distributions (CDAD)
- Various other final state effects providing information in core-level spectra

CORRELATION OF THERMOCHEMICAL DATA WITH CHEMICAL SHIFTS: EQUIVALENT-CORE OR (Z+1) MODEL

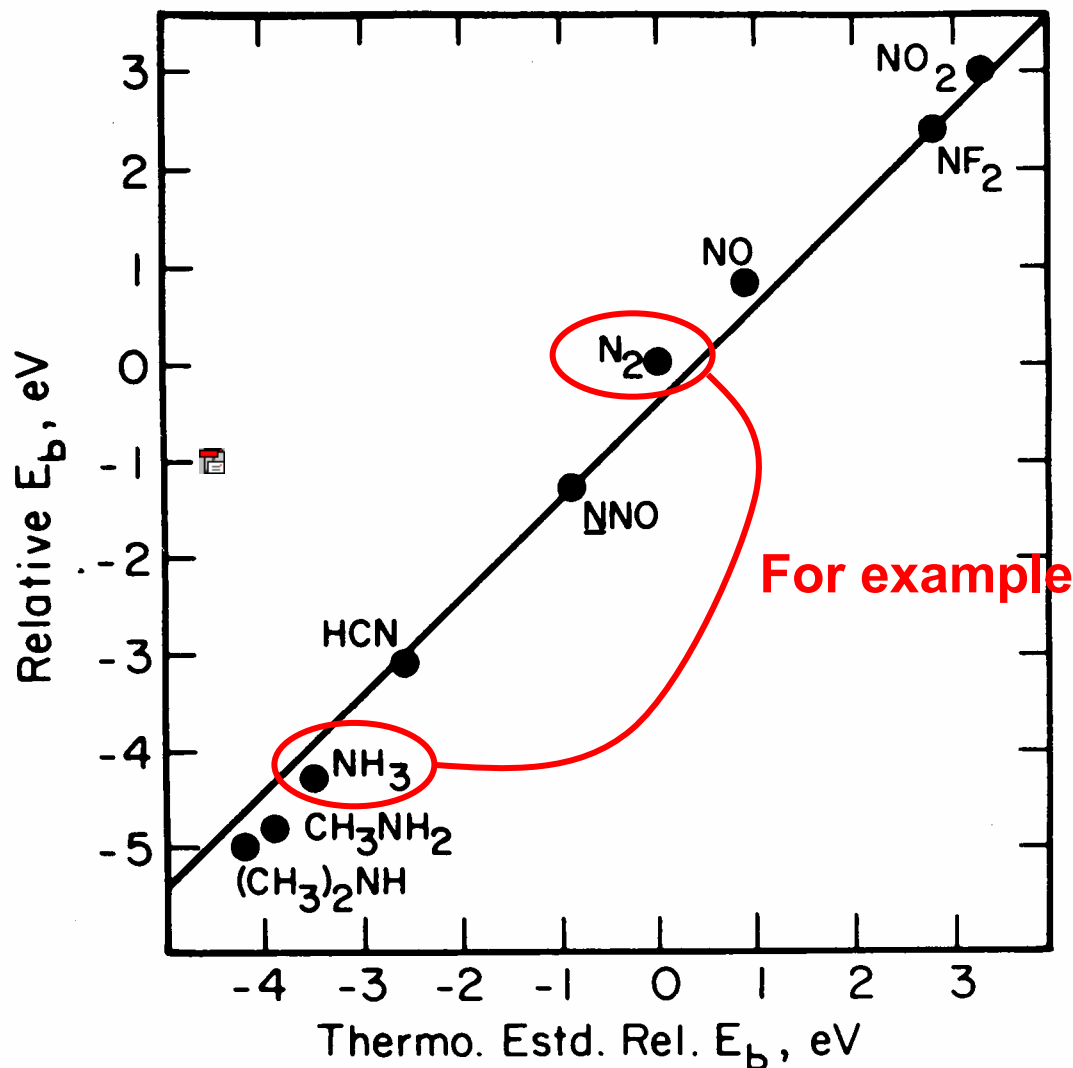
N core = N $1s^2$ = N⁵⁺



Assume:
N^{6+*} core with
1s hole = N^{6+*} =



Plus see pp. 92-93
in "Basic Concepts
of XPS"



Jolly et al.

Binding energies: * = N 1s core hole present



Adding and subtracting:

$$\begin{aligned} NH_3 + N_2^{+*} \rightarrow NH_3^{+*} + N_2 : \quad \Delta E &= \Delta E_1 - \Delta E_2 \\ &= E_b^V(N1s, NH_3) - E_b^V(N1s, N_2) \\ &= \Delta E_b^V(N1s, NH_3 - N_2) \end{aligned}$$

The chemical shift

Replacing real N 1s core with equivalent O 1s core:

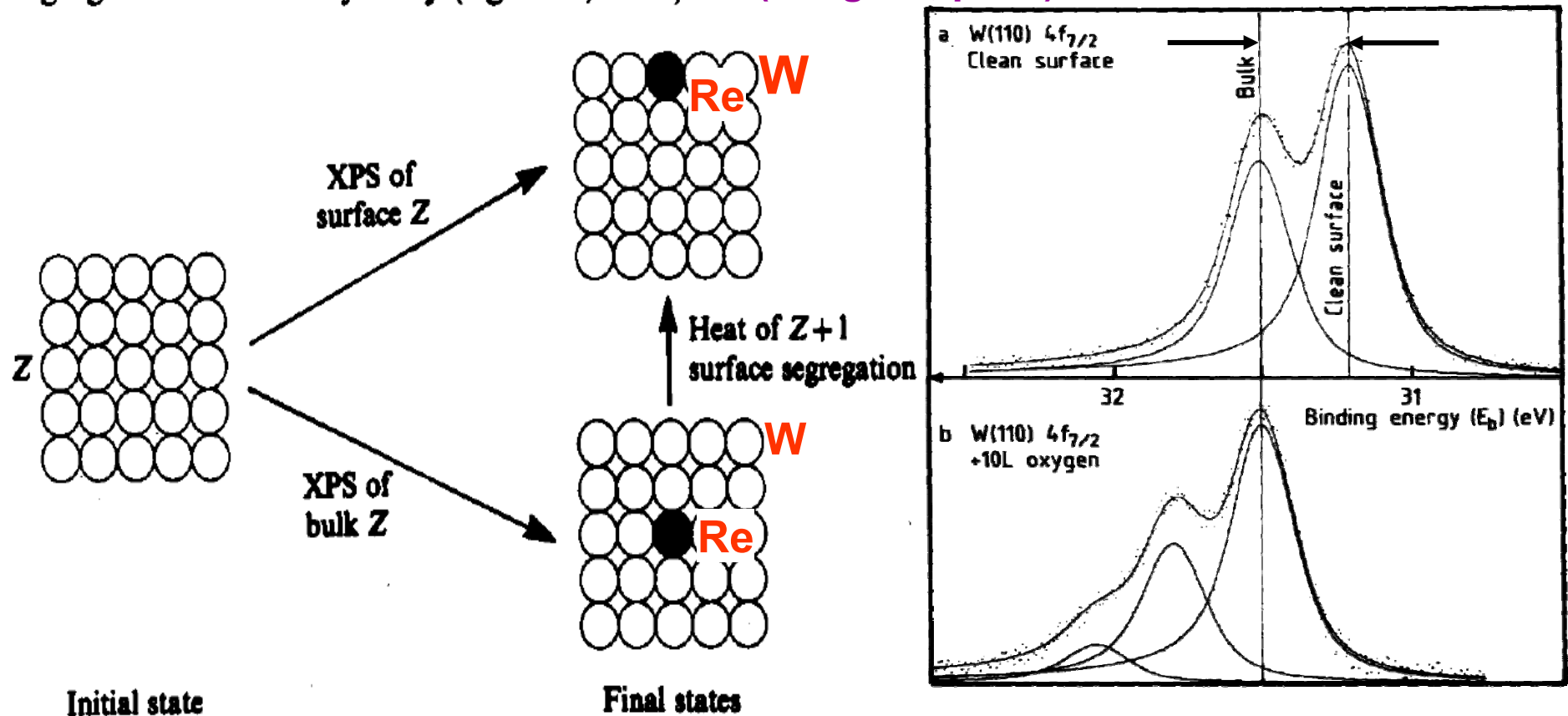
$$\begin{aligned} NH_3 + NO^+ \rightarrow OH_3^+ + N_2 : \quad \Delta E &= \Delta E_b^V(N1s, NH_3 - N_2) + \cancel{\Delta E_{ce}} - \cancel{\Delta E_{ce}} \\ &= \Delta E_b^V(N1s, NH_3 - N_2) \end{aligned}$$

A thermochemical energy

DERIVATION OF HEAT OF SURFACE SEGREGATION FROM SURFACE CORE-LEVEL CHEMICAL SHIFTS

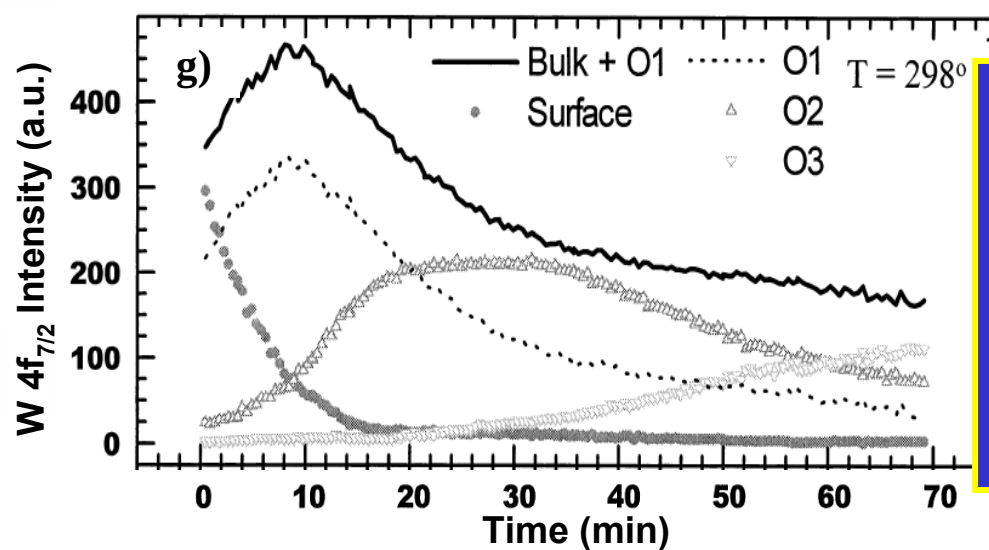
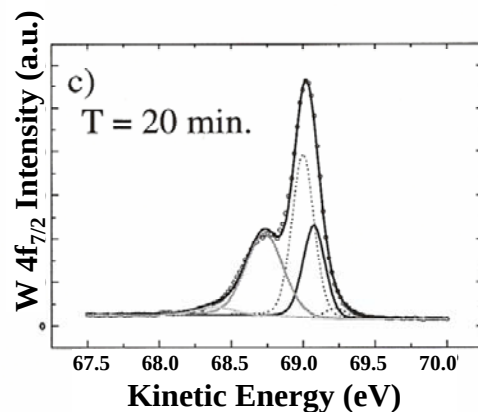
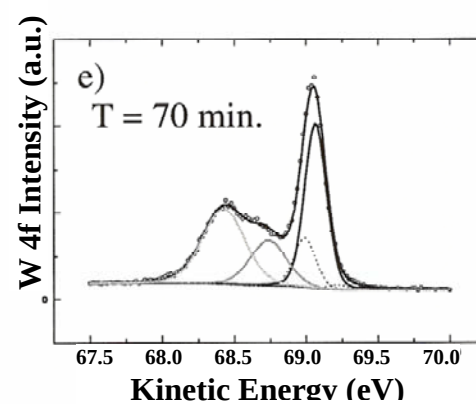
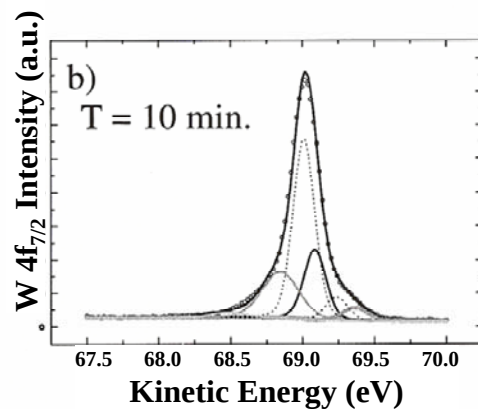
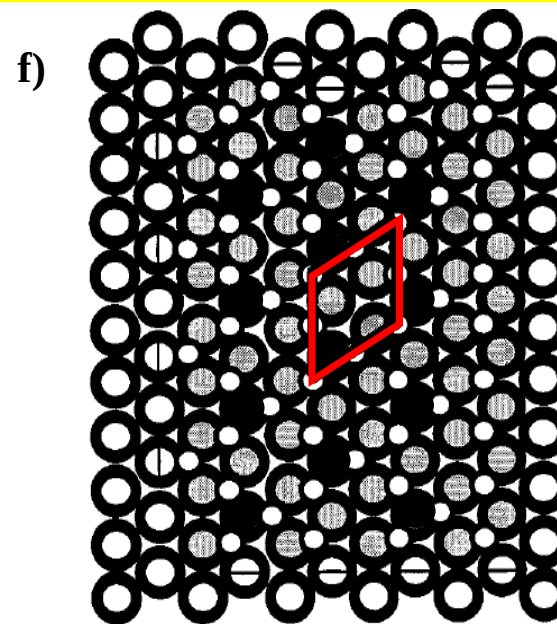
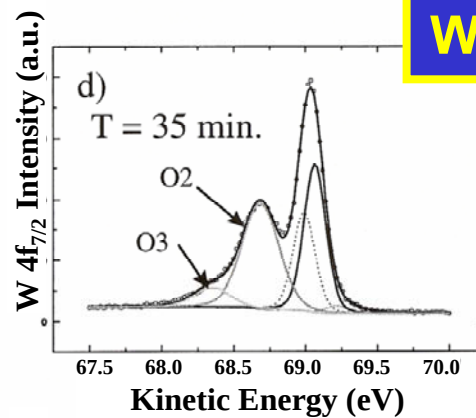
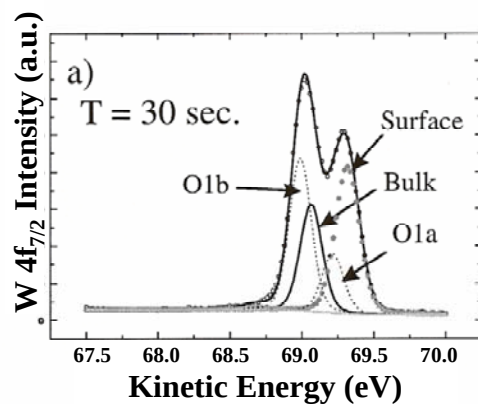
Fig. 4.30. The XPS surface core level shift approach to the heat of segregation of a binary alloy (Egelhoff, 1983). (Zangwill, p. 87)

$$\Delta E_b = \Delta H_{\text{surf. segregation}}$$



Spanjaard et al., Surf. Sci.
Repts. 5, 1 (1985)

W(110)/O—W 4f_{7/2} Chemical Shifts



An early
time-
resolved
reaction
study
(more
later)

Ynzunza et al.,
Surf. Sci. 459 (2000) 69

Outline

- Valence-band spectra: low-energy UPS limit and high-energy XPS limit

- Core-level chemical shifts: the potential model

- Core-level chemical shifts: equivalent-core ($Z+1$) and thermochemical energies



- Multiplet splittings and magnetism

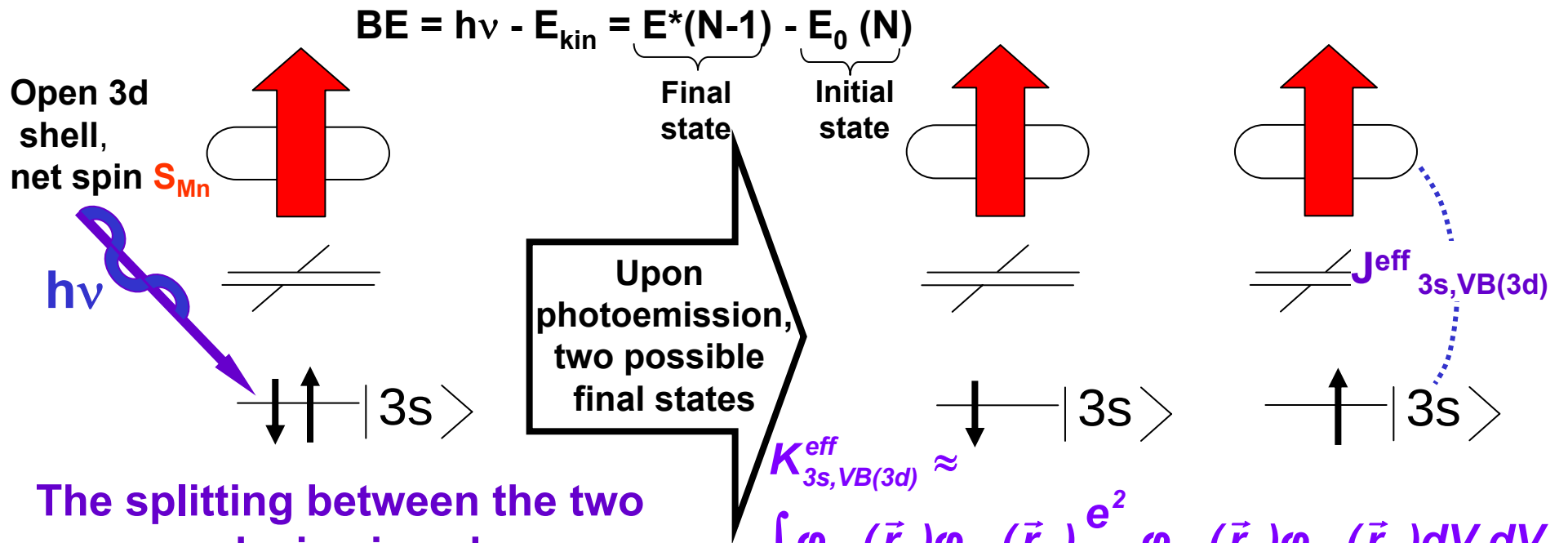
- Spin-orbit splitting, the Fano effect, and spin-polarized outgoing electrons

- Magnetic circular dichroism (MCD) in core-level emission

- Non-magnetic circular dichroism in core-level emission: a.k.a. circular dichroism in angular distributions (CDAD)

- Various other final state effects providing information in core-level spectra

Multiplet splitting in core levels of transition metal oxides

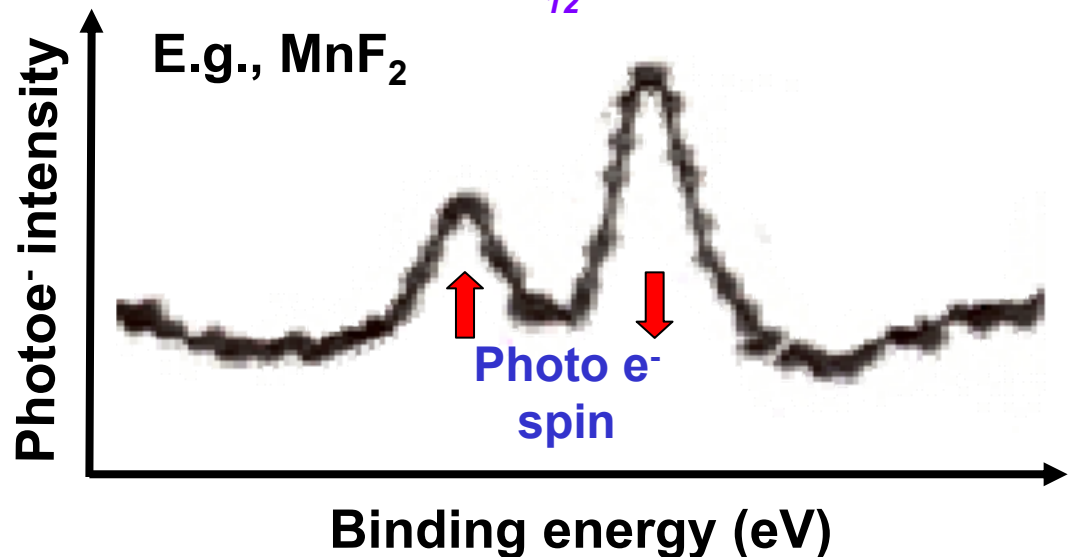


The splitting between the two peaks is given by

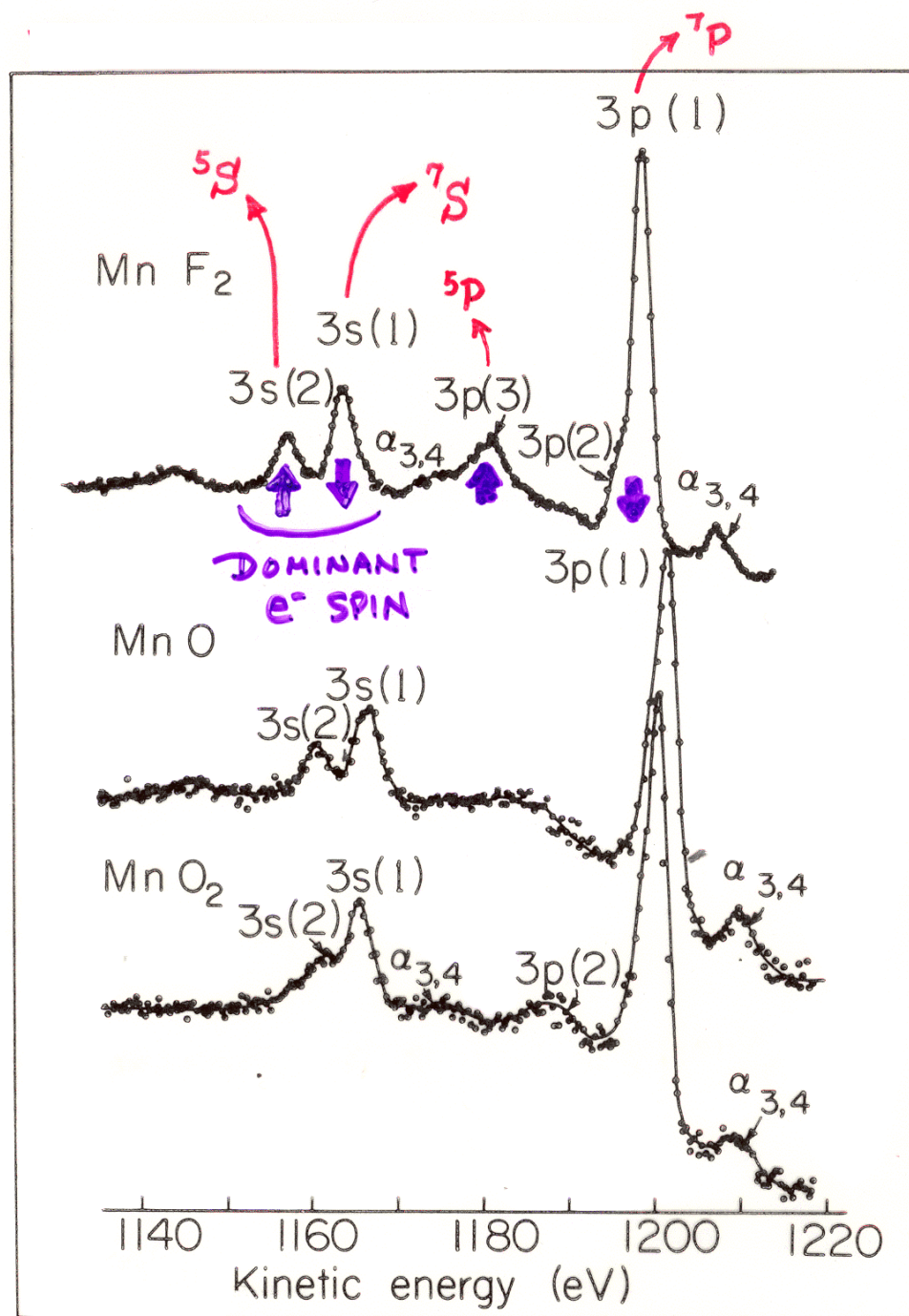
$$\Delta E_{3s} \approx (2S_{\text{Mn}} + 1) K_{3s, \text{VB}(3d)}^{\text{eff}} \quad (\text{Van Vleck Theorem})$$

For the cubic manganites in simplest doping model,

$$S_{\text{Mn}} = 1/2(4-x) \rightarrow \Delta E_{3s} \approx [5-x] K_{3s, \text{VB}(3d)}^{\text{eff}} \quad \text{with } K_{3s, \text{VB}}^{\text{eff}} \approx 1.1 \text{ eV}$$

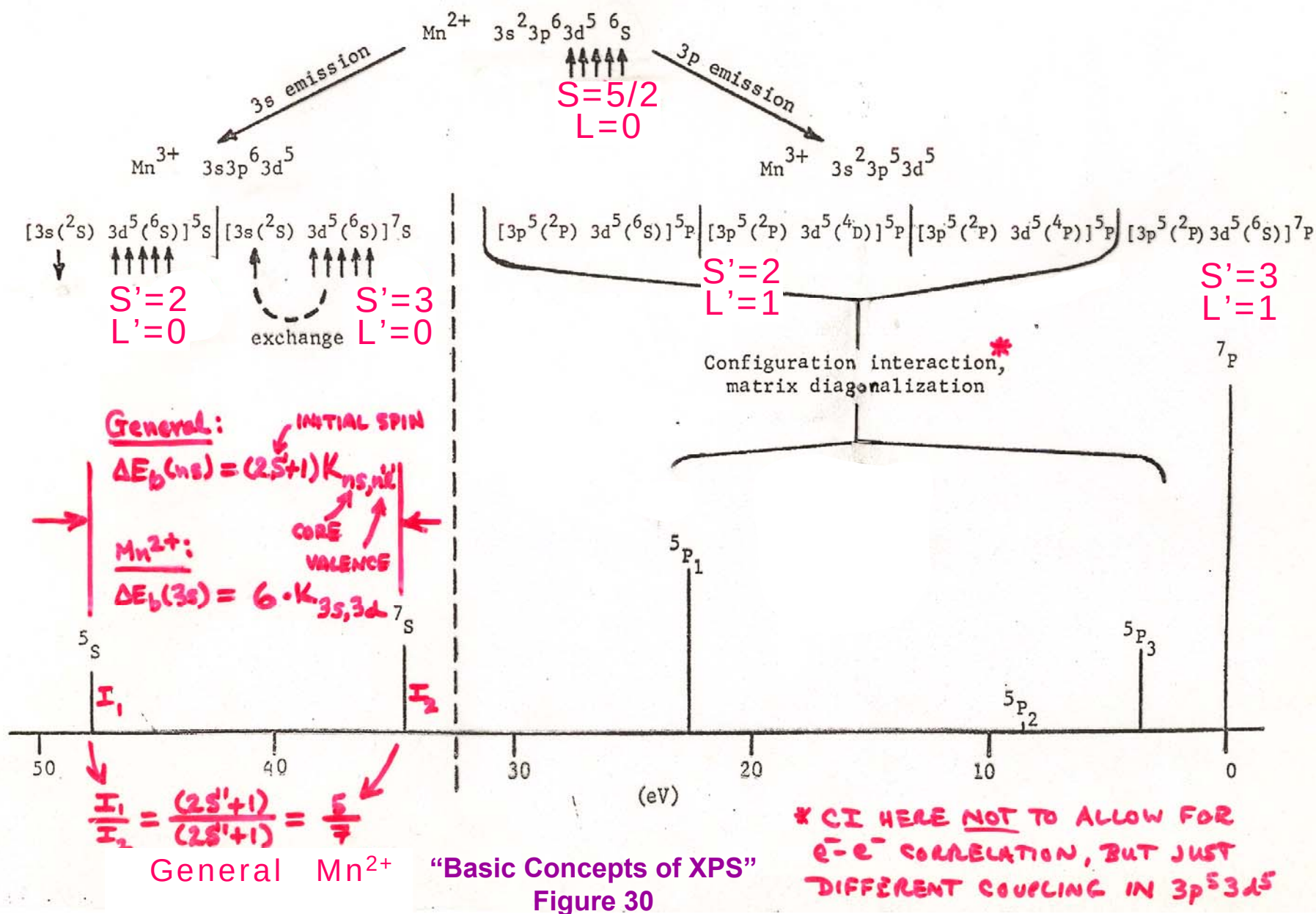


CORE-LEVEL MULTIPLY SPLITTINGS IN Mn COMPOUNDS



"Basic Concepts of XPS"
Figure 31

ORIGIN OF MULTIPLY SPLITTINGS IN Mn^{2+} : "ONE-ELECTRON" THEORY

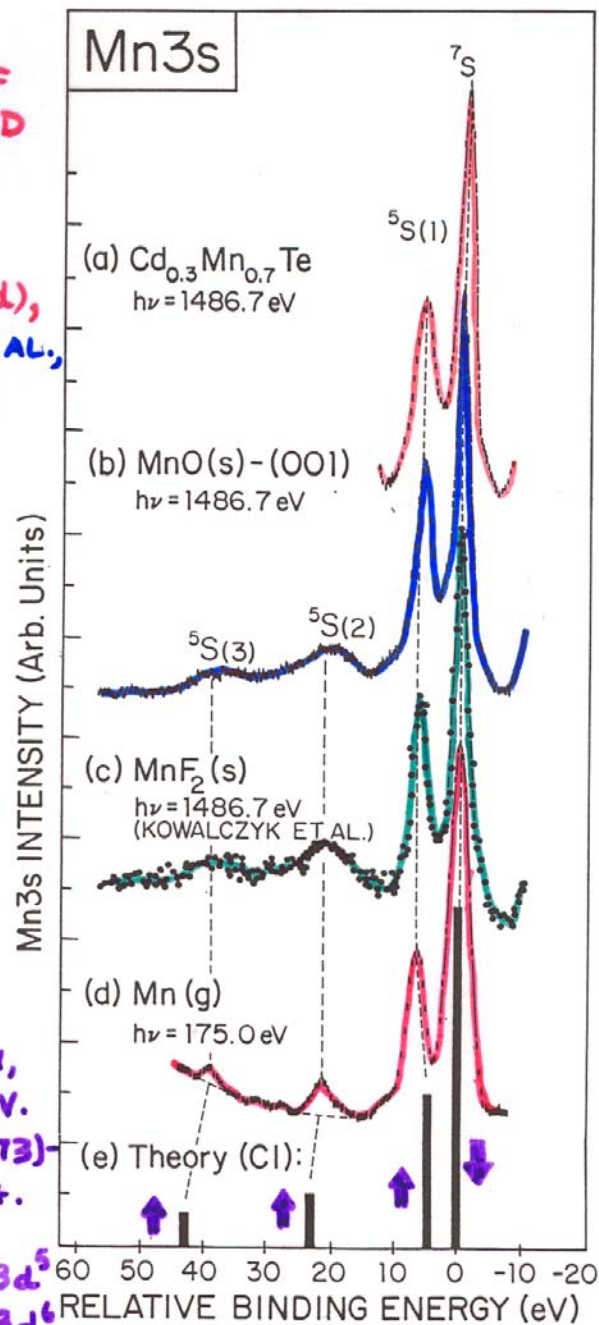


COMPARISON OF GAS-PHASE AND SOLID-STATE SPECTRA

EXPT. : (a), (b), (d),
HERMSMEIER ET AL.,
PHYS. REV. LETT.
61, 2592 (1988)
(OUR GROUP)

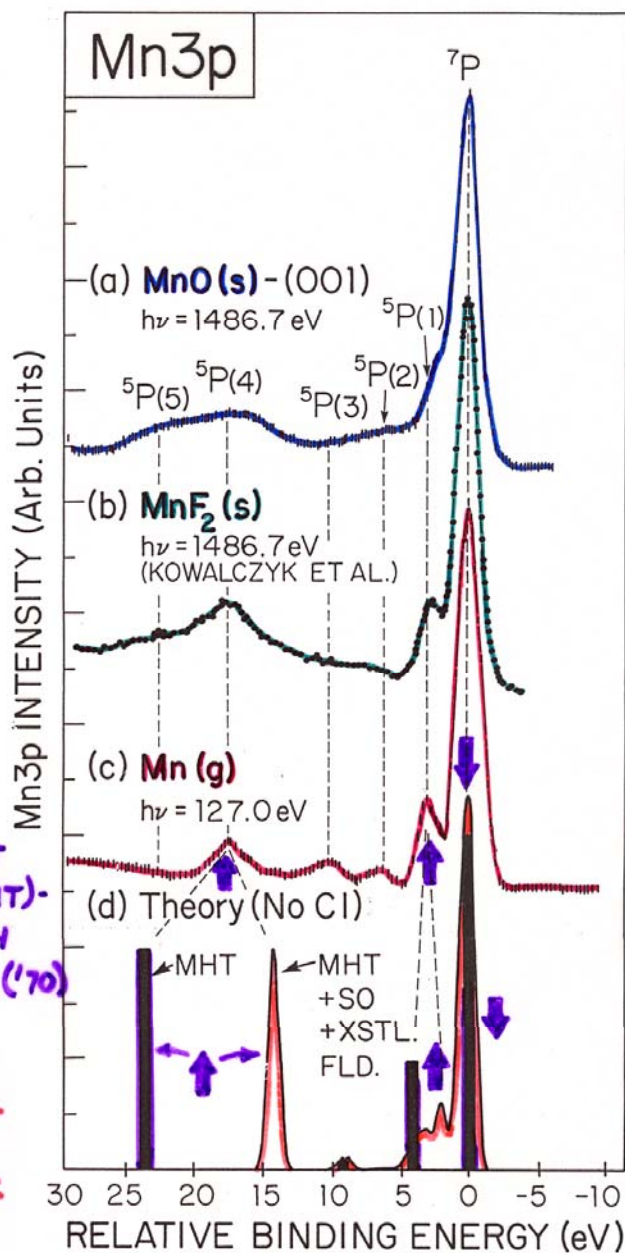
Correlation
CI effects:
anti-parallel
electrons

THEORY:
BAQUS, FREEMAN,
SASAKI, PHYS. REV.
LETT. 30, 850 (1973)
ATOMIC CONFIG.
INT. IN
 $Mn^{2+} \dots 3s^2 \dots 3d^5$
 $+ Mn^{3+} \dots 3s^2 3p^4 3d^6$



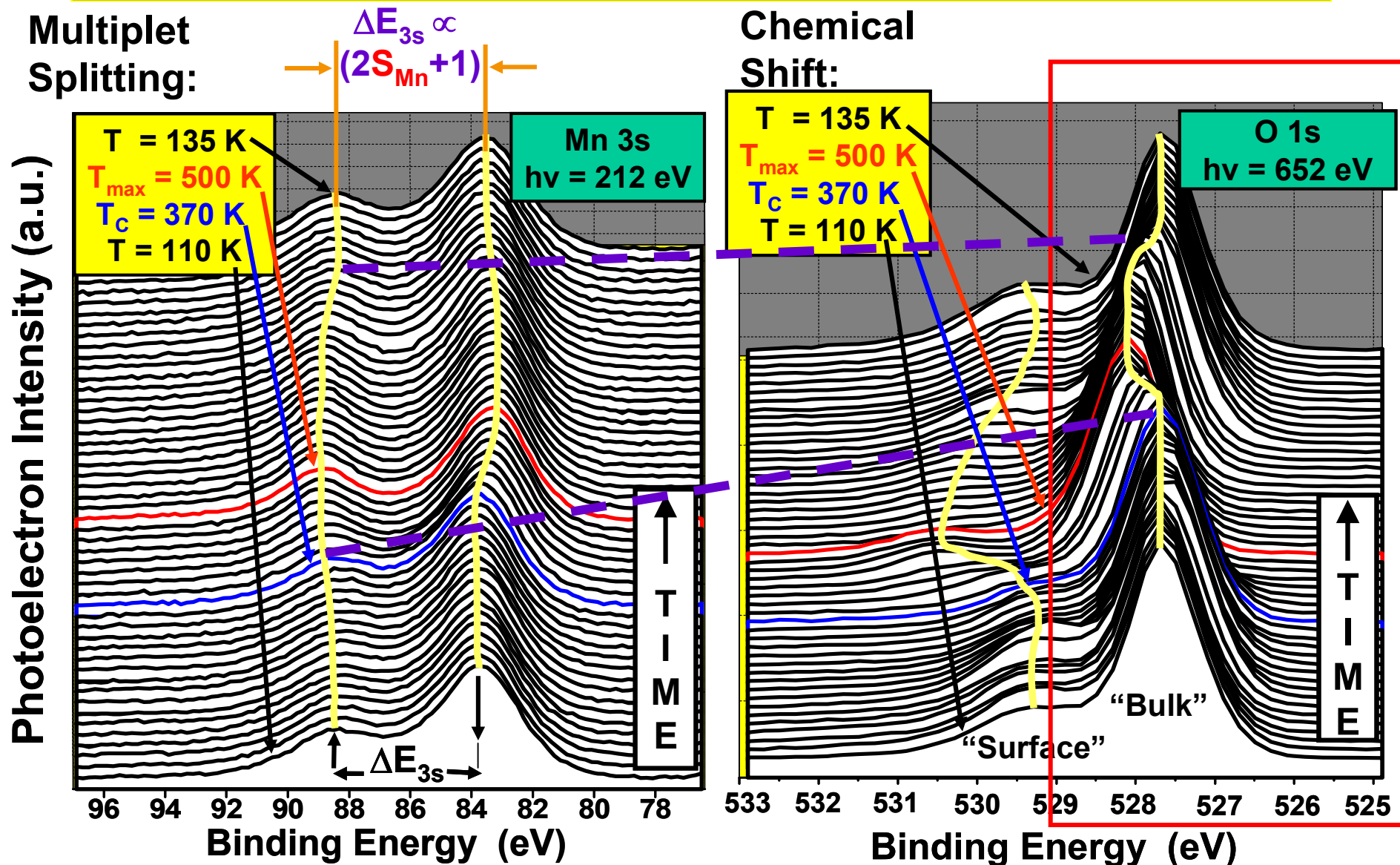
"Basic Concepts of XPS"
Figure 33

THEORY: NO CI
 SIMPLE MULTIPLET
 HOLE THEORY (MHT)-
 FADLEY, SHIRLEY
 PHYS. REV. A2, 1109 ('70)
 EMPIRICAL
 MHT WITH SPIN
 ORBIT & CRYSTAL
 FIELD - SUGANO
 ET AL., J. PHYS. C
 15, 2625 (1982)



HERMSMEIER
 ET AL.,
 P.R.L. 61, 2592 ('88)

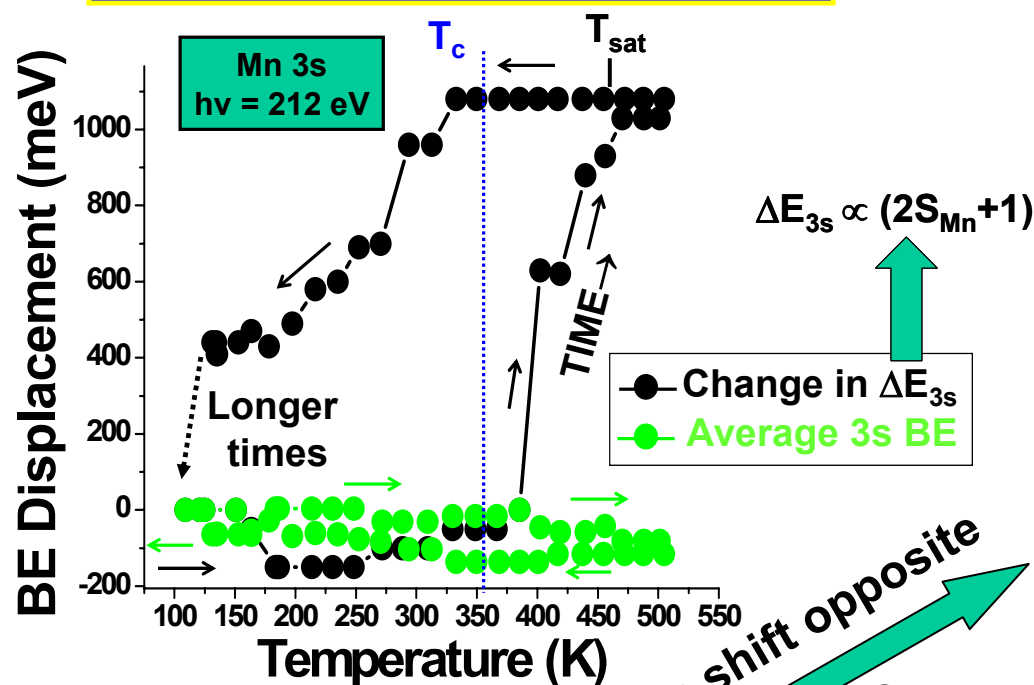
Temperature dependence of Mn3s and O1s spectra in a colossal magnetoresistive (CMR) oxide: $\text{La}_{0.7}\text{Sr}_{0.3}\text{MnO}_3$



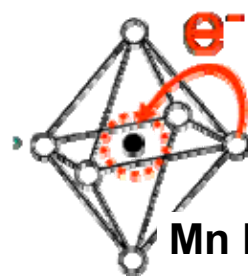
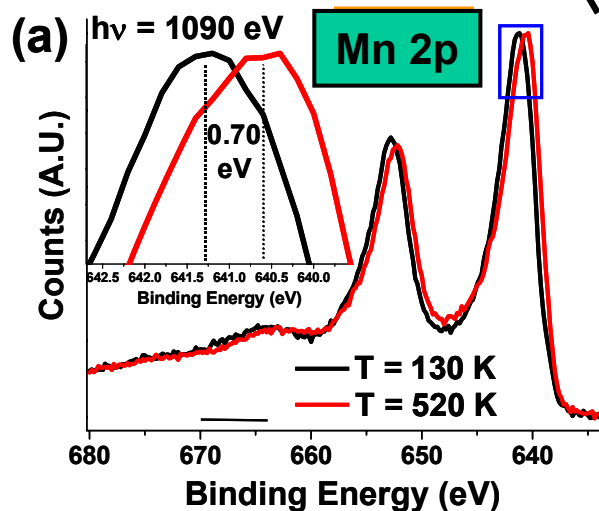
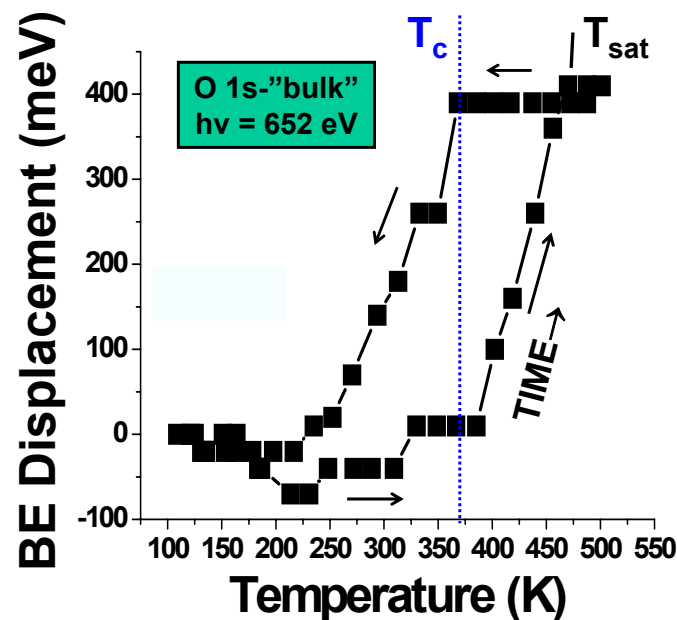
Increase of the Mn3s splitting-reversible

Increase of O1s BE-reversible

T dependence of Mn 3s binding energies



T dependence of bulk O 1s binding energy

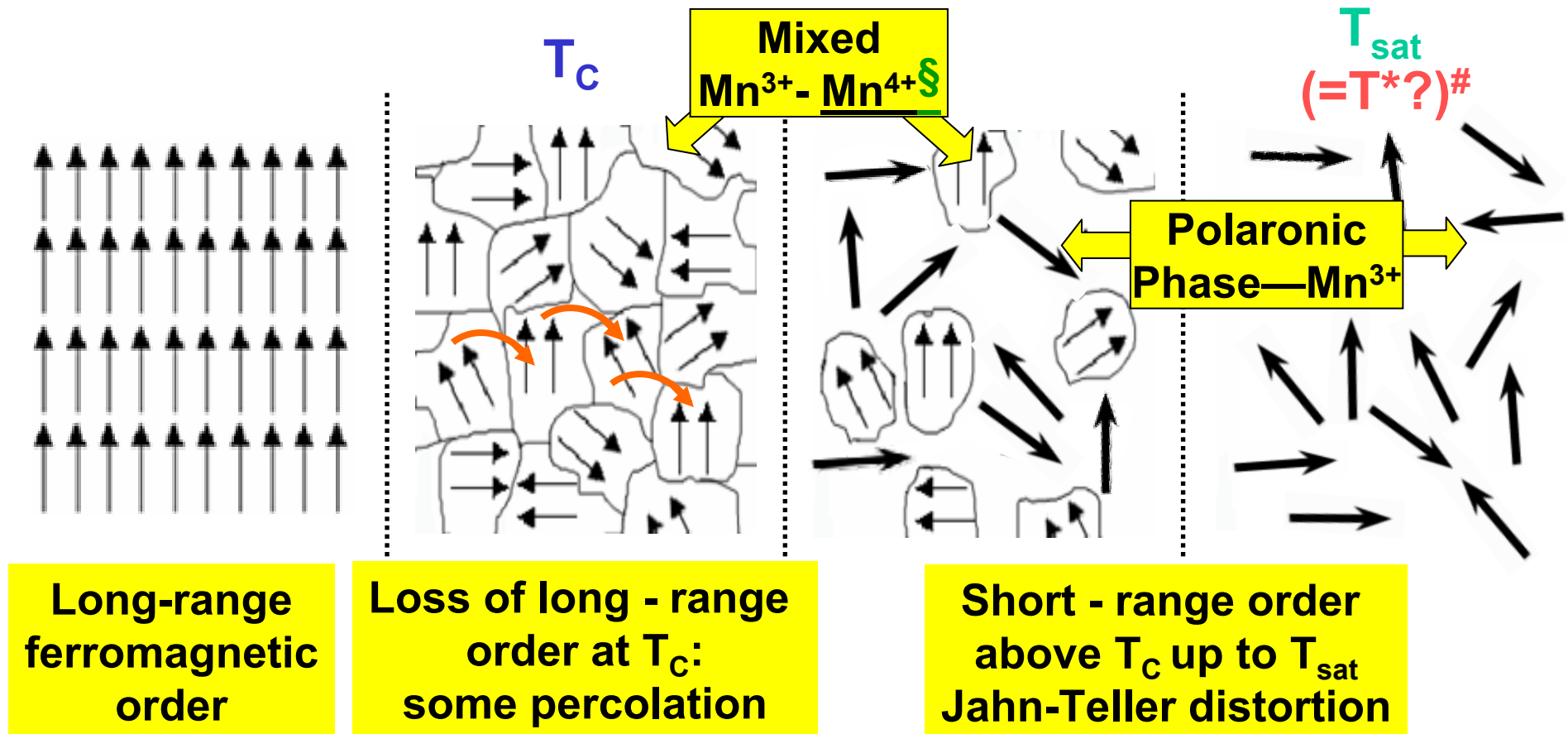


O more positive
Mn higher spin,
more negative

Theory: self-interaction corrected local spin density results for LSMO $\rightarrow$ elongating octahedron makes Mn^{3+} more stable than Mn^{4+}
Banach, Temmerman, PRB 69, 054427 ('04)

Mannella et al., Phys. Rev. Lett. 92, 166401 (2004)

Suggested scenario—LSMO, $x = 0.3, 0.4$



§ Self-interaction corrected local spin density calcs. suggest Mn^{4+} dominant, but conversion to Mn^{3+} with JT distortion
Banach, Temmerman, PRB 69, 054427 ('04)

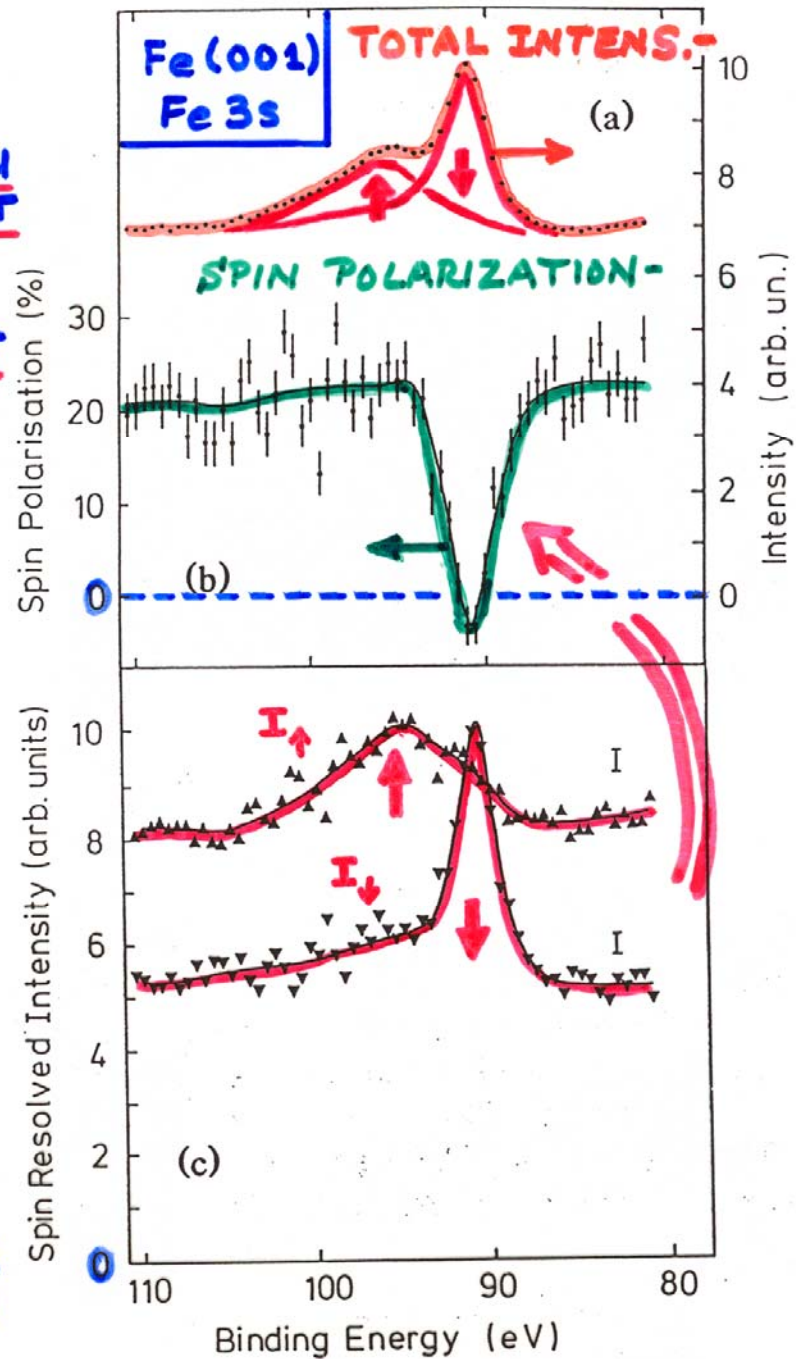
T^* = new T scale suggested from theory
Dagotto et al. PRL 87, 277202 (2001)

Mannella et al., PRL 92, 166401 ('04); PRB 70, 224433 ('04), and to be publ.

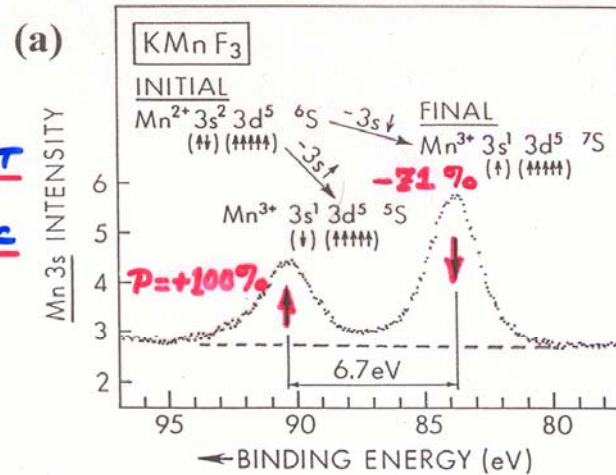
DIRECT
OBSERVATION
OF SPIN-SPLIT
CORE LEVELS
IN A
FERROMAGNET

$$\frac{I_{\uparrow} - I_{\downarrow}}{I_{\uparrow} + I_{\downarrow}}$$

HILLEBRECHT
ET AL.,
PHYS. REV. LETT.
65, 2450 (1990)



①
MULTIPLY
IN A
MAGNETIC
ATOM



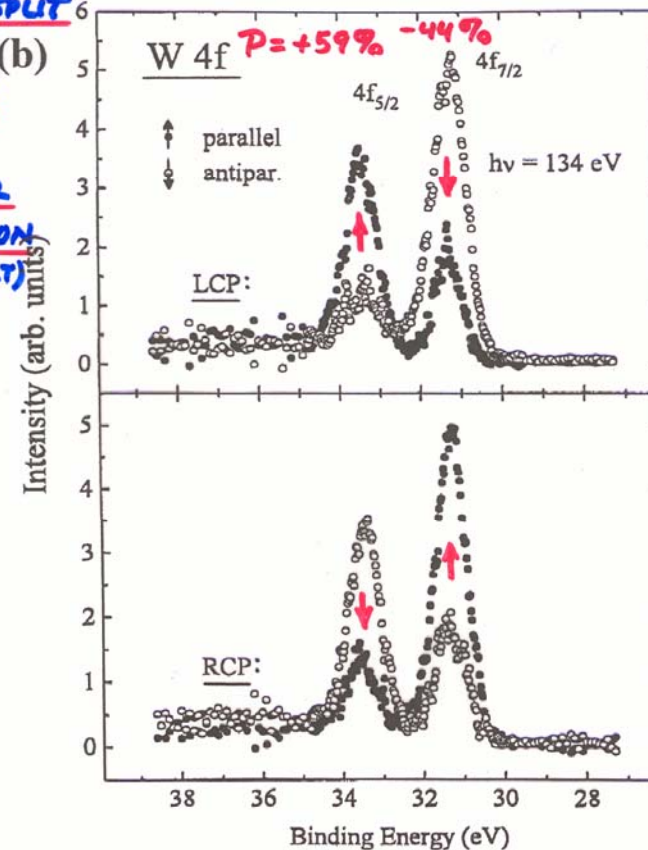
SPIN POLARIZATION
IN CORE SPECTRA

$$P = \frac{I_{\uparrow} - I_{\downarrow}}{I_{\uparrow} + I_{\downarrow}}$$

EXPT. - FINKOVIC
ET AL.
P.R.L. 55,
1227 (1985)

Spin
internally
referenced
to spin of
each ion

②
SPIN-ORBIT SPLIT
LEVEL
EXCITED
WITH
CIRCULAR
POLARIZATION
(FANO EFFECT)

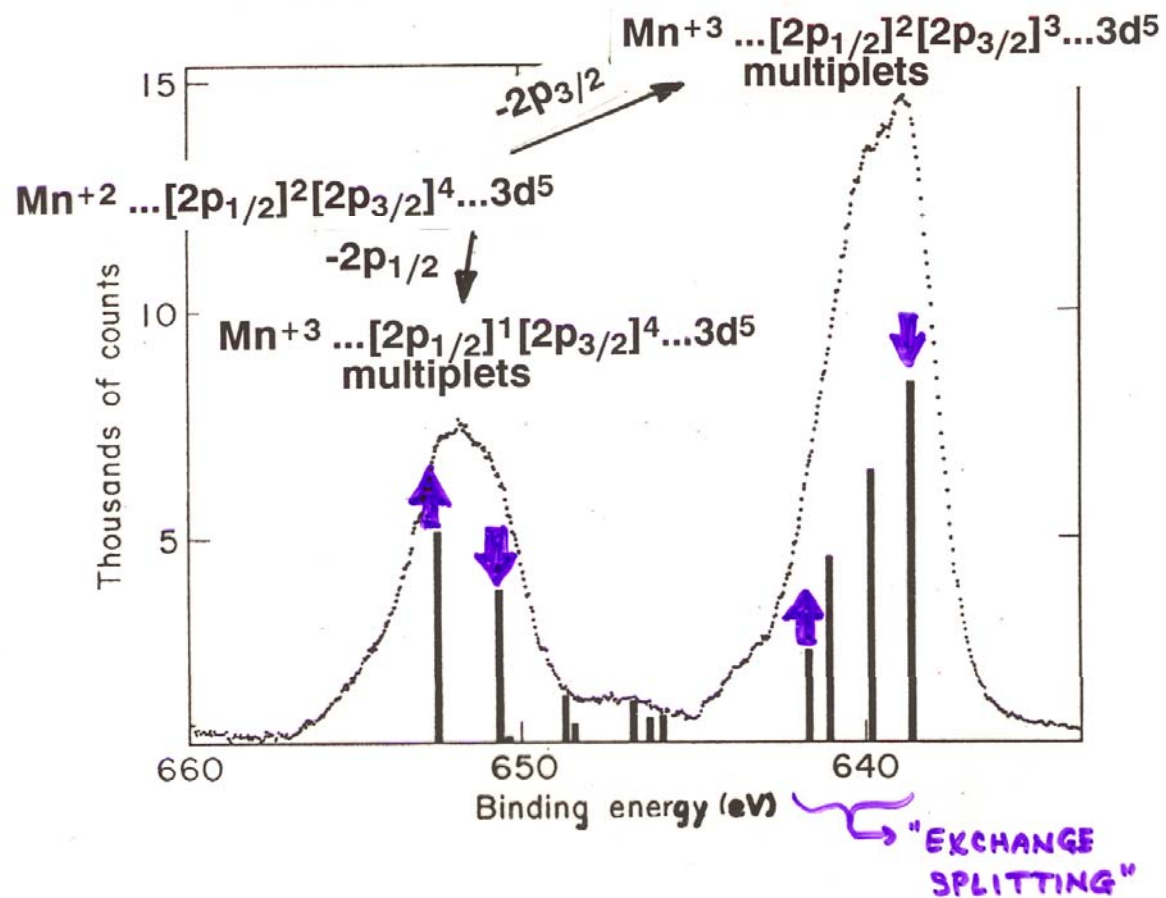


EXPT. - STARR ET
AL.
P.R.B. 53, 210544
(1996)

Spin
externally
referenced
to $\vec{k}_{h\nu}$ and $\vec{M}$
of sample

**+ MORE COMPLEX MULTIPLETS FOR $L > 0$
WITH SPIN-ORBIT COUPLING:**

Mn 2p emission from MnF_2 :

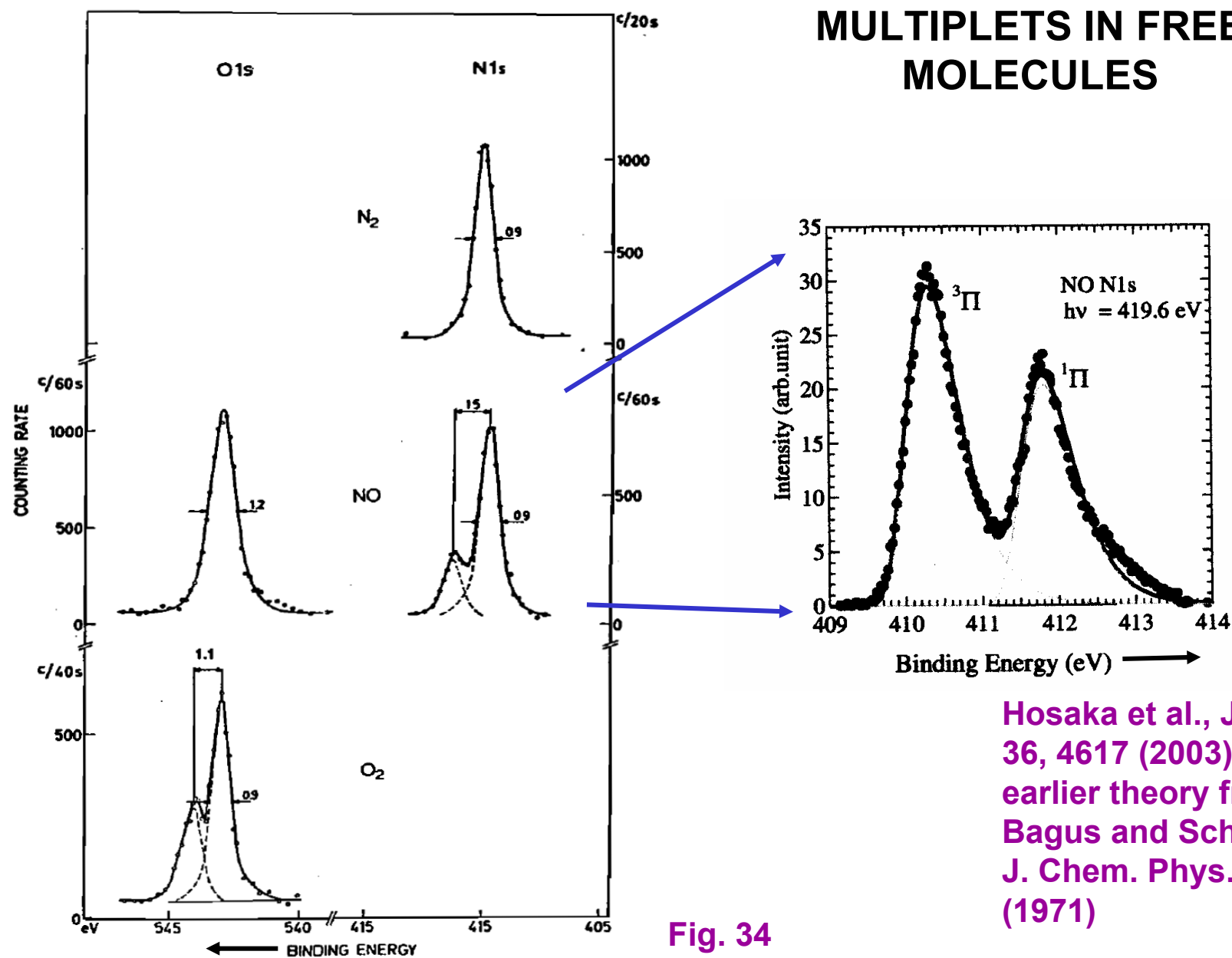


Expt.--Kowalczyk et al., Phys. Rev. B 11, 1721 (1975)

Theory--Gupta and Sen, Phys. Rev. B 10, 71 (1974)

Park et al., Phys. Rev. B 37, 10867 (1988)

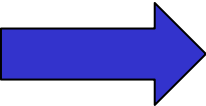
MULTIPLETS IN FREE MOLECULES



Hosaka et al., J. Phys. B
36, 4617 (2003), and
earlier theory from
Bagus and Schaefer,
J. Chem. Phys. 55, 1474
(1971)

Fig. 34
Basic Concepts of XPS

Outline

- Valence-band spectra: low-energy UPS limit and high-energy XPS limit
- Core-level chemical shifts: the potential model
- Core-level chemical shifts: equivalent-core ($Z+1$) and thermochemical energies
- Multiplet splittings
-  • Spin-orbit splitting, the Fano effect, and spin-polarized outgoing electrons
- Magnetic circular dichroism (MCD) in core-level emission
- Non-magnetic circular dichroism in core-level emission: a.k.a. circular dichroism in angular distributions (CDAD)
- Various other final state effects providing information in core-level spectra

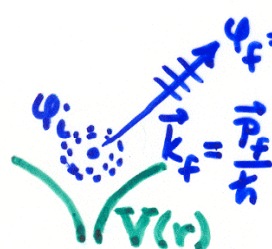
PHOTOELECTRON EMISSION-

BASIC MATRIX ELEMENTS + SELECTION RULES:

● ATOMIC-LIKE (LOCALIZED) STATES $\Rightarrow$ CORE:

PLUS SPIN:

$$\psi_i(\vec{r}) = \psi_{n_i l_i m_i}(\vec{r}, \theta, \phi) = R_{n_i l_i}(r) Y_{l_i m_i}(\theta, \phi) \begin{cases} \alpha(\sigma) = m_{s_i} = +1/2 = \uparrow \\ \beta(\sigma) = m_{s_i} = -1/2 = \downarrow \end{cases}$$



$$\psi_f(\vec{r}, \vec{k}_f) = \psi_{E_f}(\vec{r}, \vec{k}_f) \begin{cases} \alpha(\sigma) \\ \beta(\sigma) \end{cases}$$

$$= 4\pi \sum_{l_f, m_f} i^{l_f} e^{-i\delta_{l_f}} Y_{l_f m_f}^*(\theta, \phi) Y_{l_f m_f}(\theta, \phi) R_{E_f, l_f}(r) \begin{cases} \alpha(\sigma) \\ \beta(\sigma) \end{cases}$$

PHASE SHIFT OF l_f WAVE IN $V(r)$

DIPOLE: INT. $\propto |\langle \psi_f | \hat{E} \cdot \vec{r} | \psi_i \rangle|^2 = |\hat{E} \cdot \langle \psi_f | \vec{r} | \psi_i \rangle|^2 \Rightarrow \left\{ \begin{array}{l} \Delta l = l_f - l_i = \pm 1 \\ \text{TWO CHANNELS} \\ \Delta m = m_f - m_i = 0, \pm 1 \\ \text{LINEAR POLAR.} \\ \Delta m = \pm 1, \text{ CIRCULAR POLARIZATION} \end{array} \right\}$

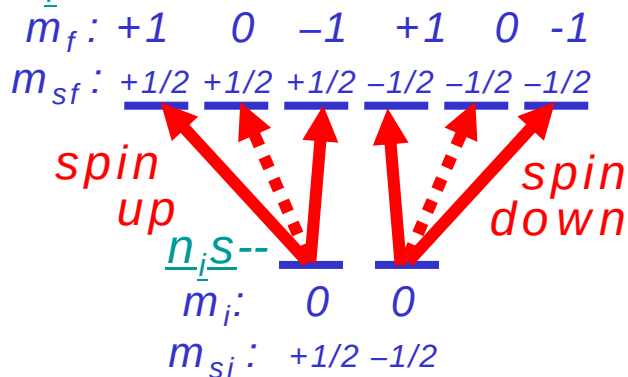
EQUIVALENT
WITHIN CONSTANT
FACTOR



$\Delta m = m_f - m_i = 0, \pm 1$
LINEAR POLAR.
 $\Delta m = \pm 1$, CIRCULAR POLARIZATION

$$\Delta m_s = m_{sf} - m_{si} = 0!$$

E_f p--

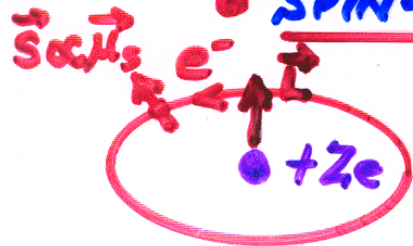


FOR A GIVEN $n_i / m_i m_{s_i}$: SUM OVER DEGENERATE INITIAL STATES $m_i m_{s_i}$ AND AVERAGE OVER FINAL STATES $E_f / m_f m_{s_f}$ ACCESSED FROM EACH m_i TO YIELD DIFFERENTIAL SUBSHELL PHOTOELECTRIC CROSS SECTION:

$$d\sigma_{n_i l_i} / d\Omega$$

$\propto$ PROBABILITY PER UNIT SOLID ANGLE OF EXCITING ONE ELECTRON FROM SUBSHELL n_i / l_i INTO THE DIRECTION k_f

• SPIN-ORBIT SPLITTING OF LEVELS:



⇒ EFFECTIVE $\vec{B}$ (NUCLEUS AROUND e^-) $\propto \vec{L}$

$$\hat{H}_{s.o} = \xi(r) \vec{L} \cdot \vec{S}$$

- SPLITS ALL nl LEVELS $2(2l+1)$
- $nl_j = l + 1/2 \rightarrow 2l+2$
 - $nl_j = l - 1/2 \rightarrow 2l$

• MIXES SPIN + ORBITAL ANGULAR MOM.:

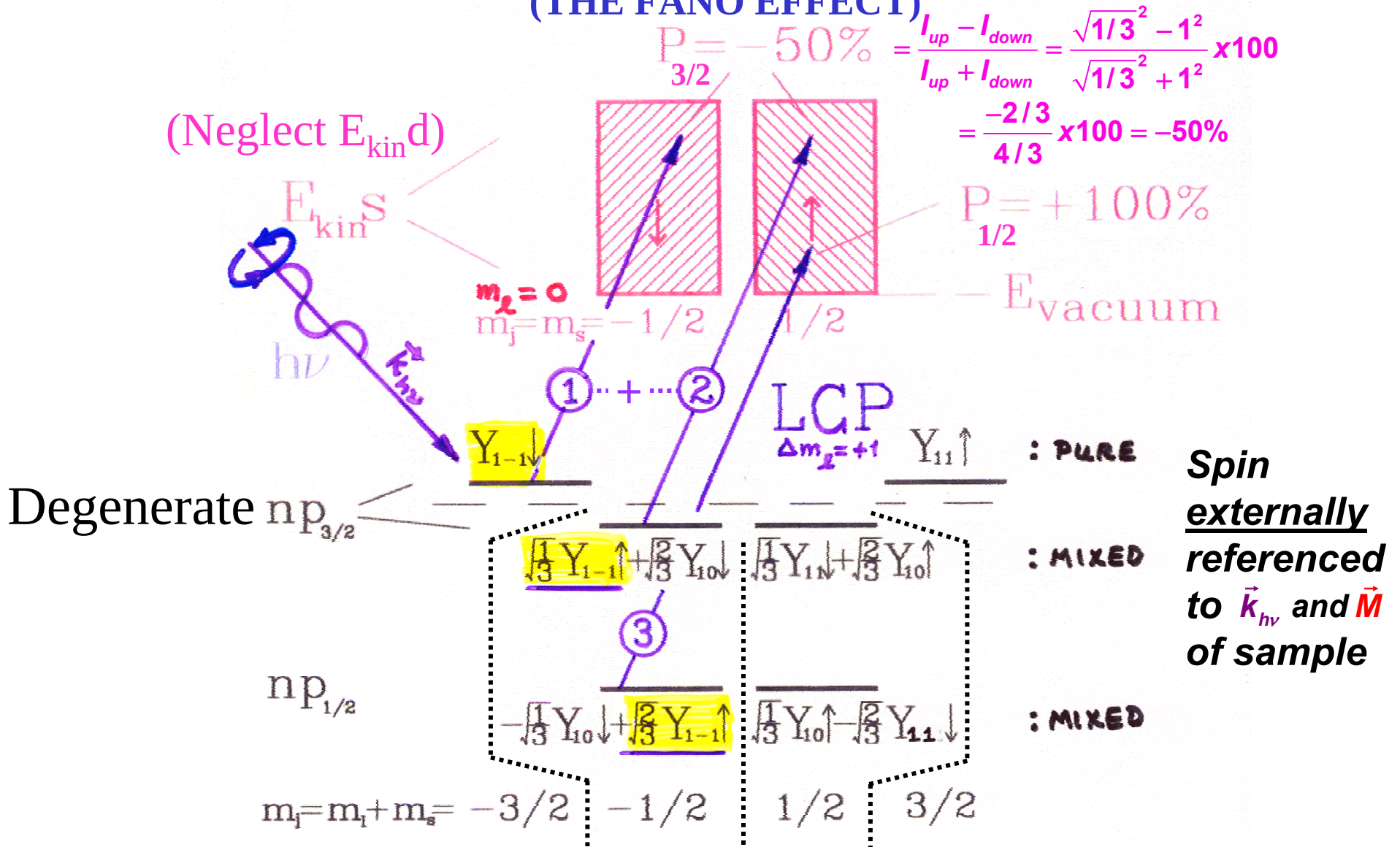
$$\psi_{nljm_j} = C_1 \psi_{nl, m_j - 1/2} \begin{pmatrix} 1 \\ 0 \end{pmatrix} + C_2 \psi_{nl, m_j + 1/2} \begin{pmatrix} 0 \\ 1 \end{pmatrix}$$

$\parallel$
 $m_s = +1/2$
 $\parallel$
 $\uparrow$

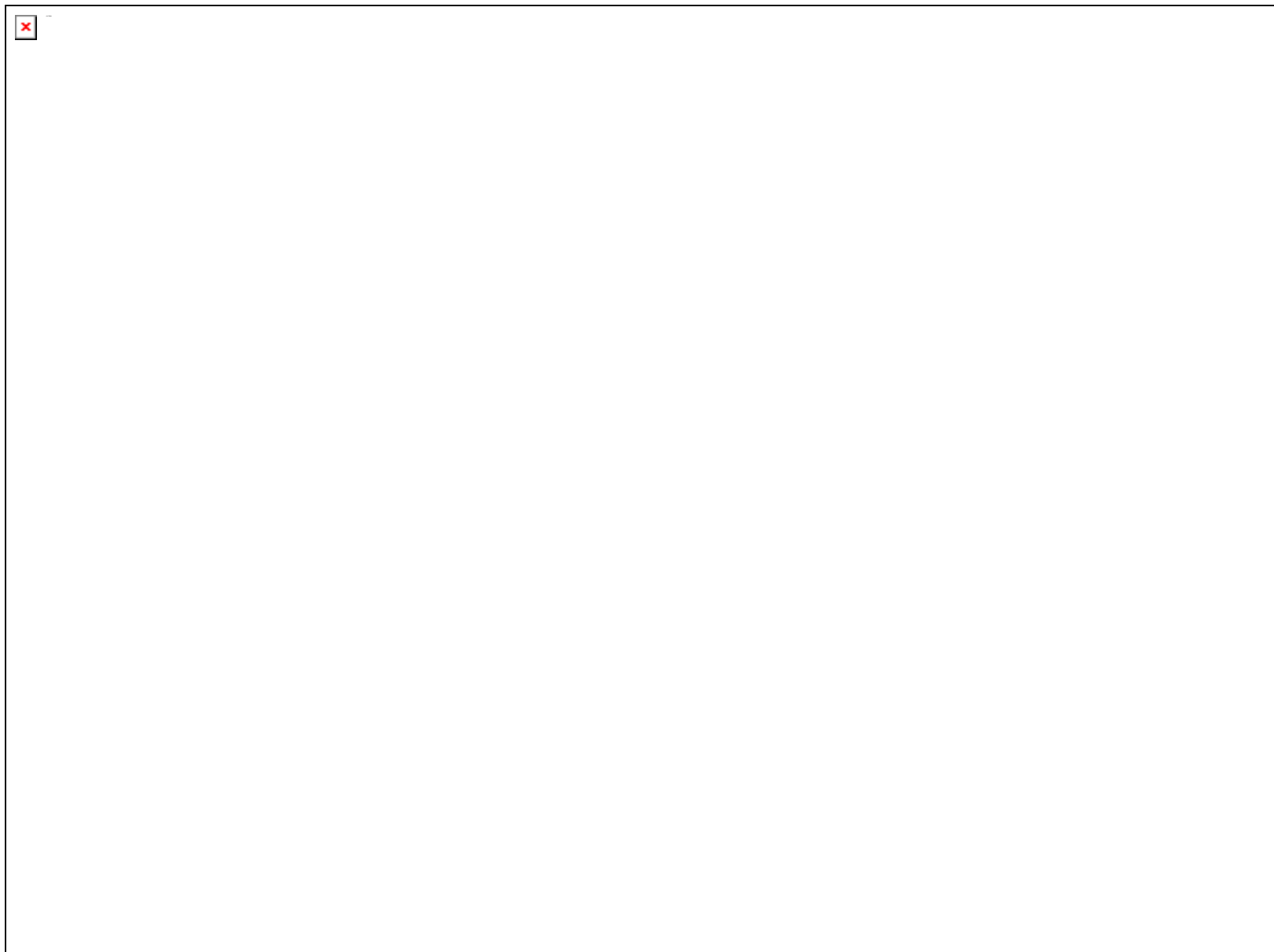
$\parallel$
 $m_s = -1/2$
 $\parallel$
 $\downarrow$

WITH C1 AND C2 TABULATED CLEBSCH-GORDAN OR WIGNER 3j SYMBOLS

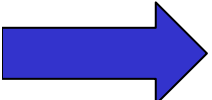
PHOTOELECTRON SPIN POLARIZATION FROM CIRCULAR POLARIZATION AND SPIN-ORBIT SPLITTING (THE FANO EFFECT)



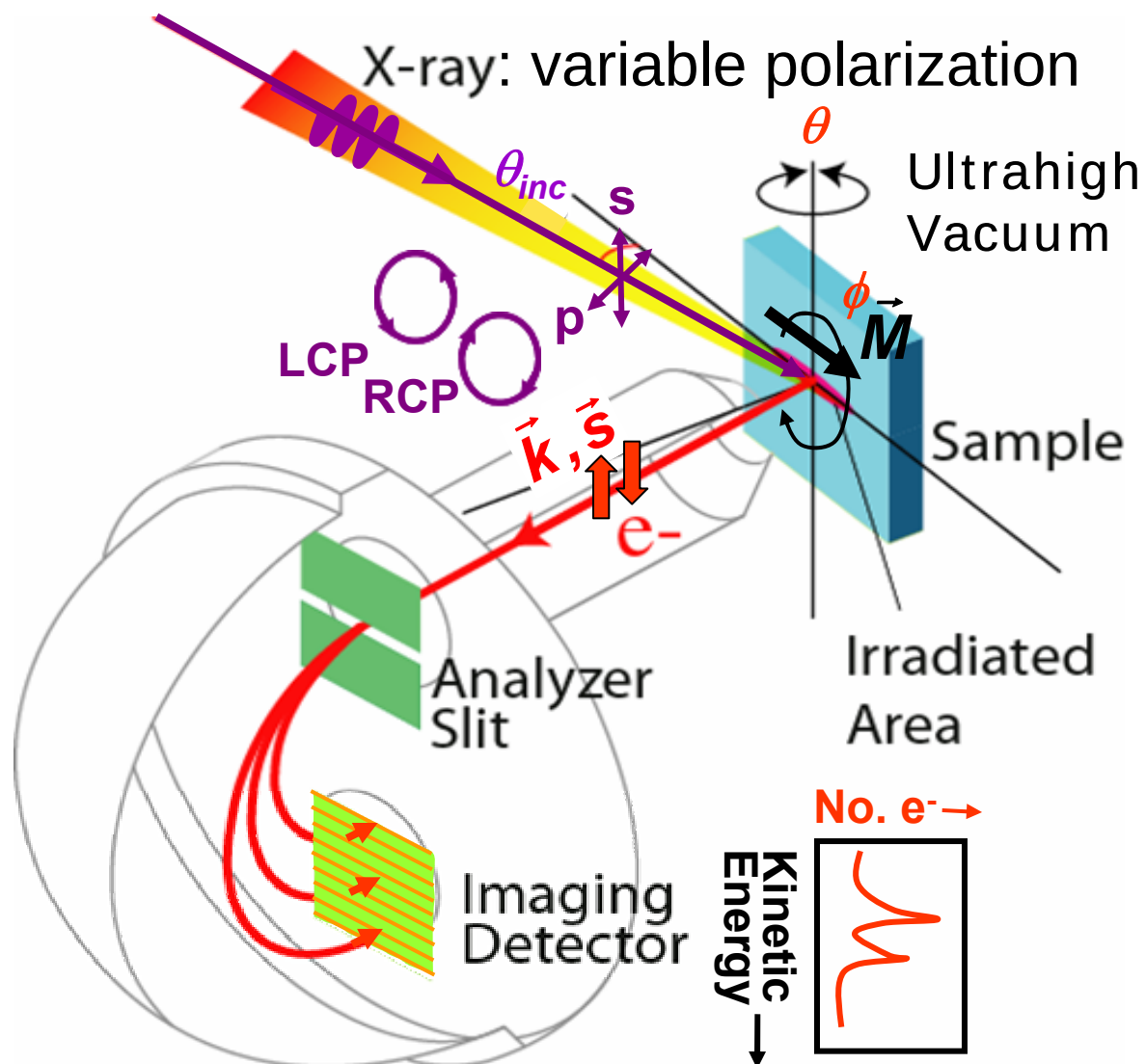
Spin polarization in core photoelectron spectra—expt.



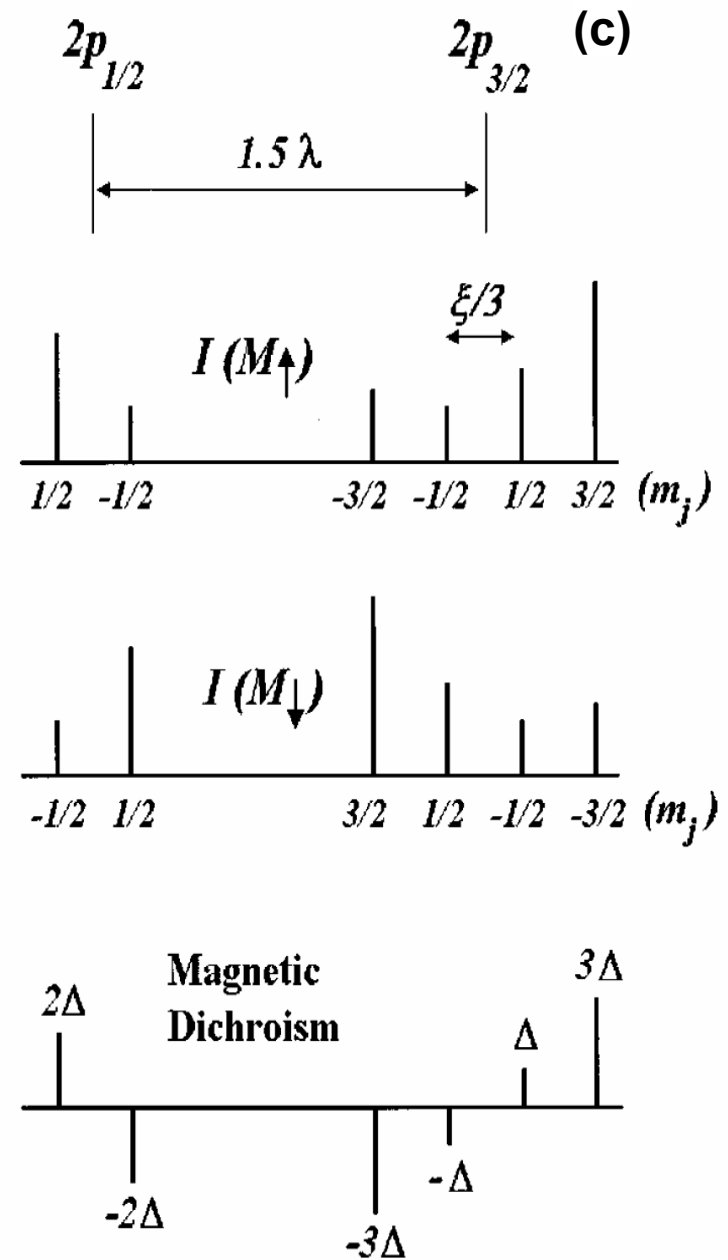
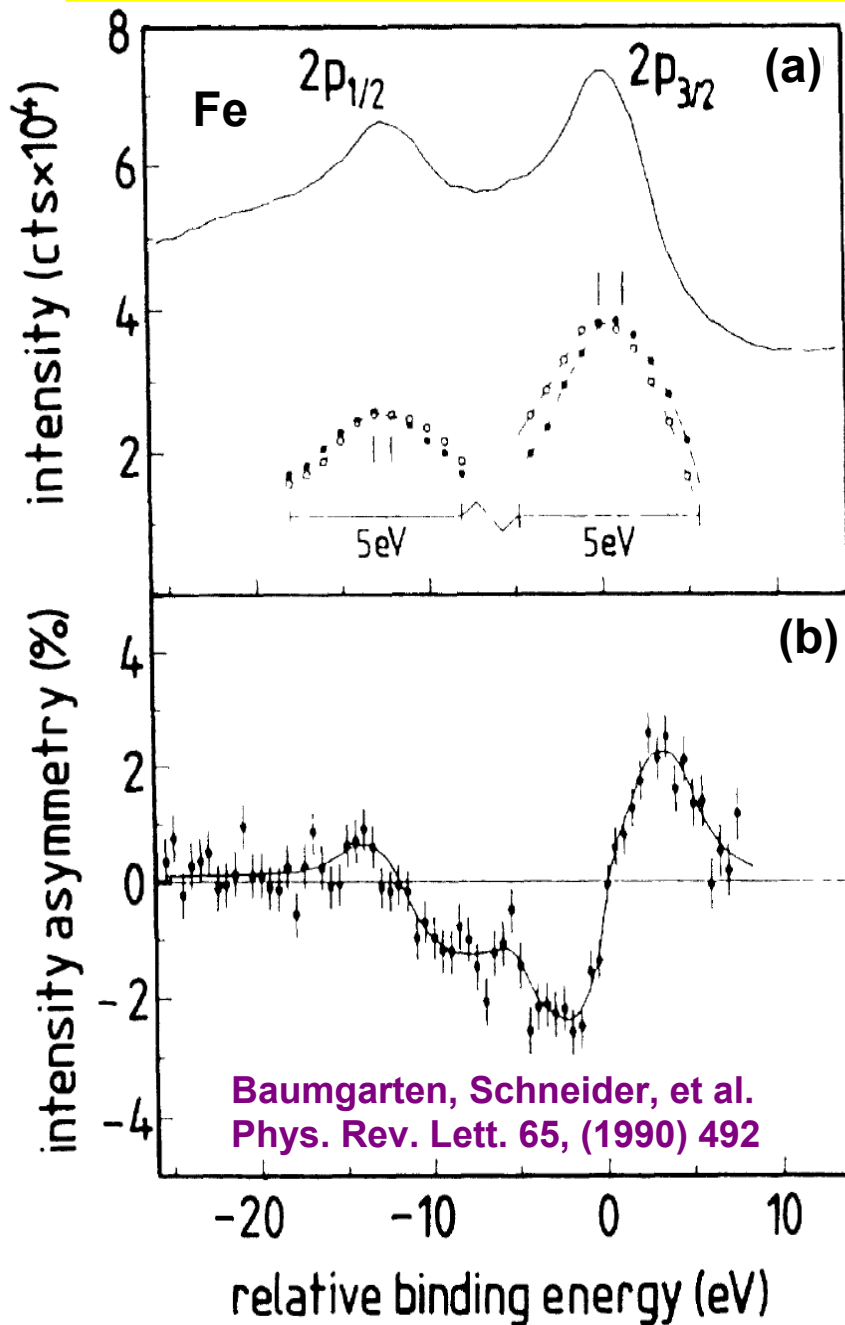
Outline

- Valence-band spectra: low-energy UPS limit and high-energy XPS limit
- Core-level chemical shifts: the potential model
- Core-level chemical shifts: equivalent-core ($Z+1$) and thermochemical energies
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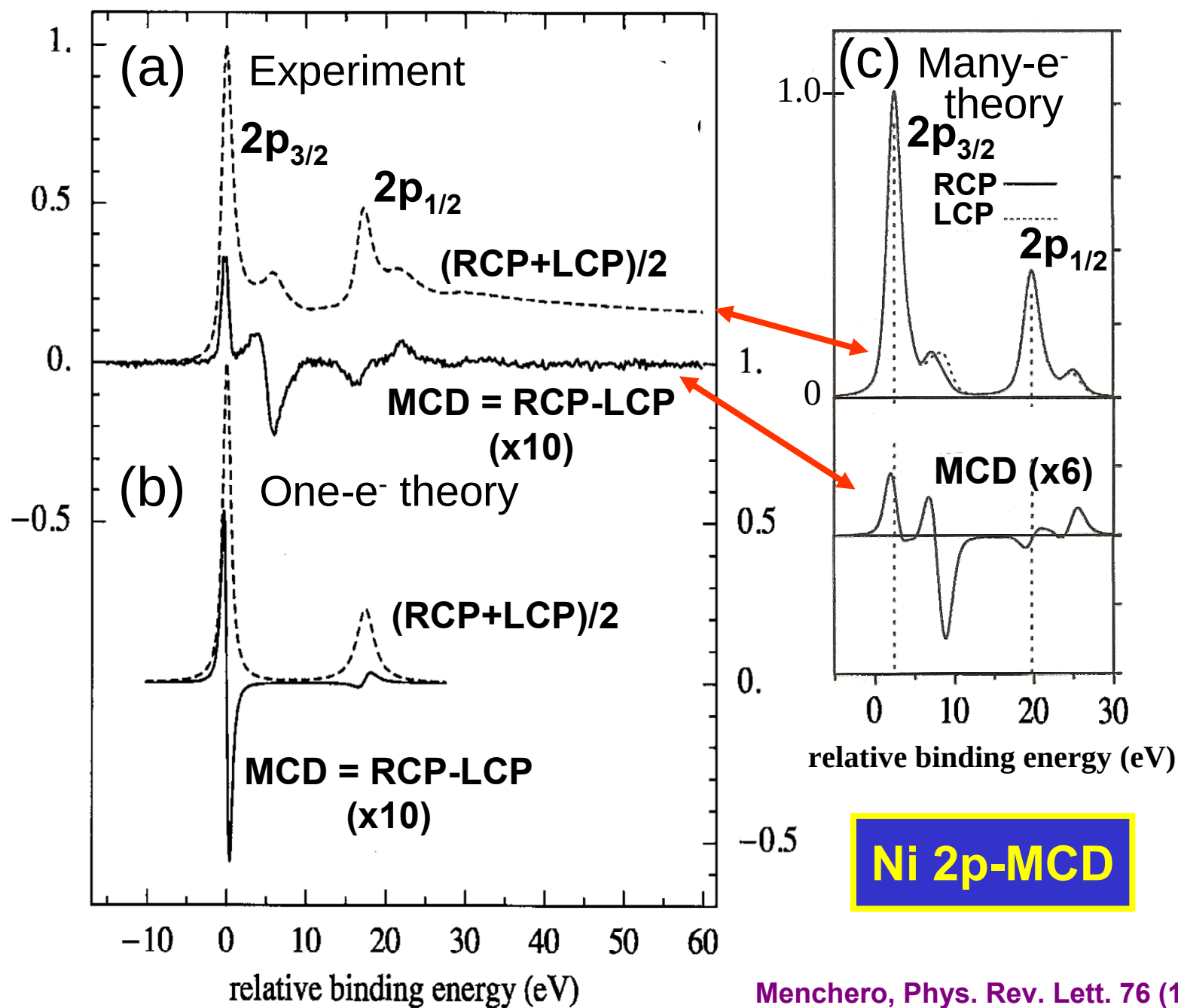
Typical experimental geometry for energy- and angle-resolved photoemission measurements



1st Expt. Magnetic Circular Dichroism 1-e- Theo.



Menchero, Phys. Rev. B 57 (1998) 993

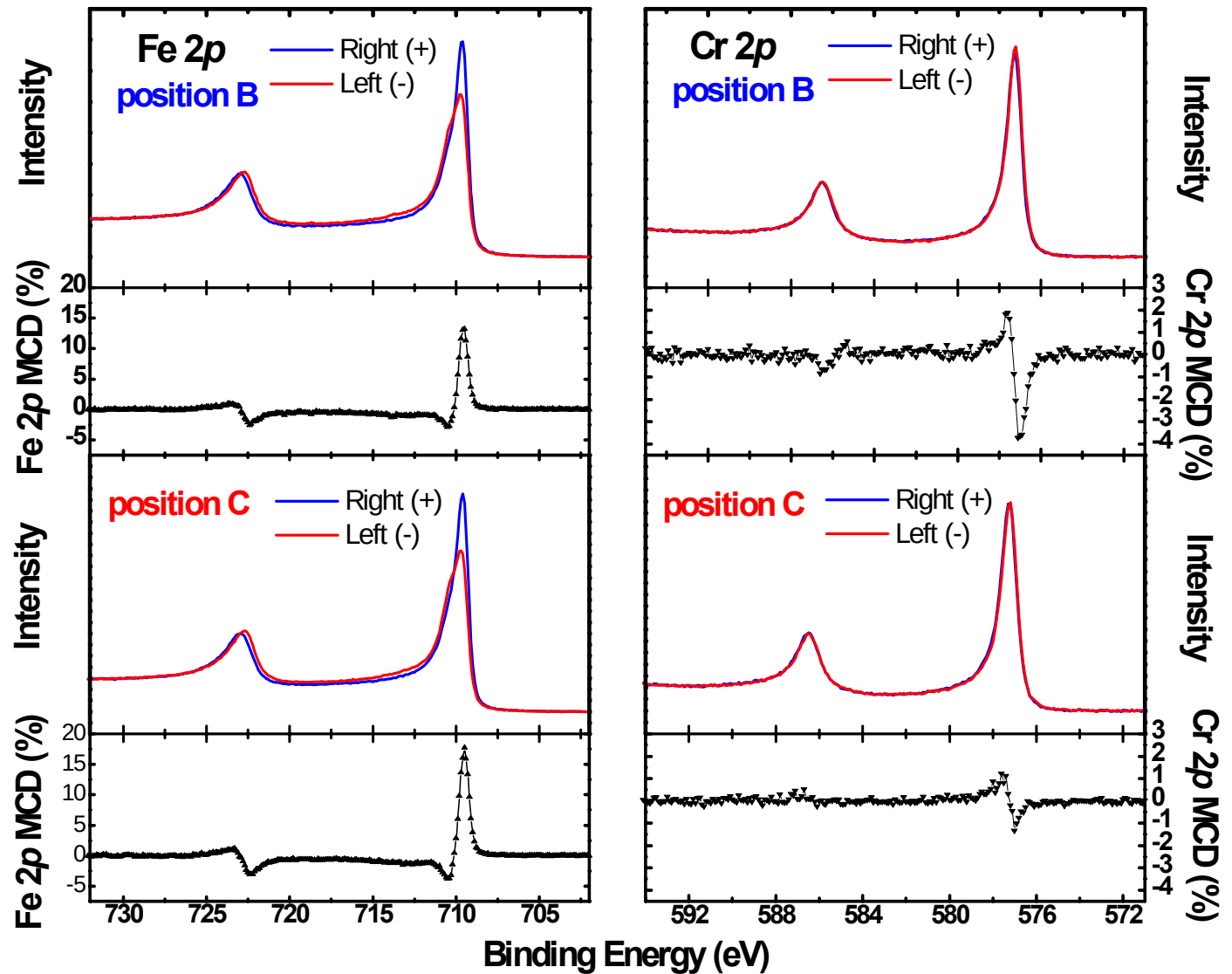
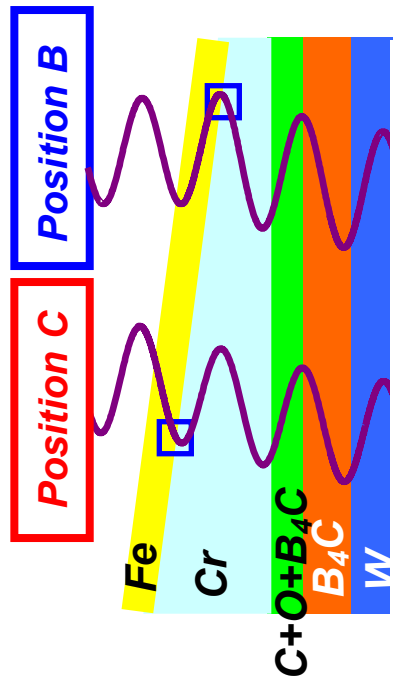
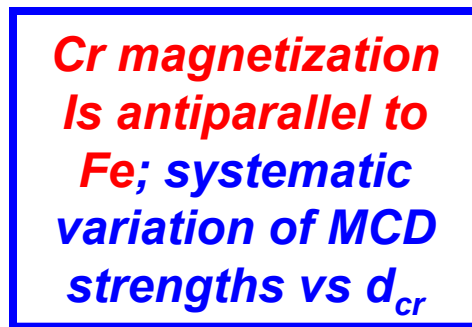


Menchero, Phys. Rev. Lett. 76 (1996) 3208

Van der Laan et al., J. Phys. Condens. Matter 12 (2000) L275

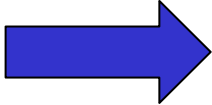
Application to a buried interface: with standing wave excitation

Fe & Cr 2p MCD Data from wedge (Fe/Cr)+SWG



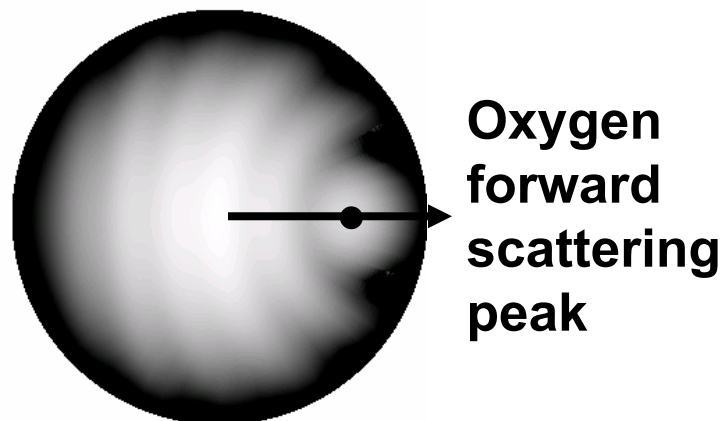
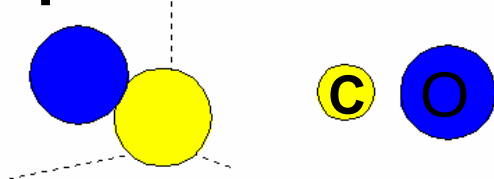
S.H. Yang , B.S. Mun et al., J. Phys. Cond. Matt. 14, L406 (2002)

Outline

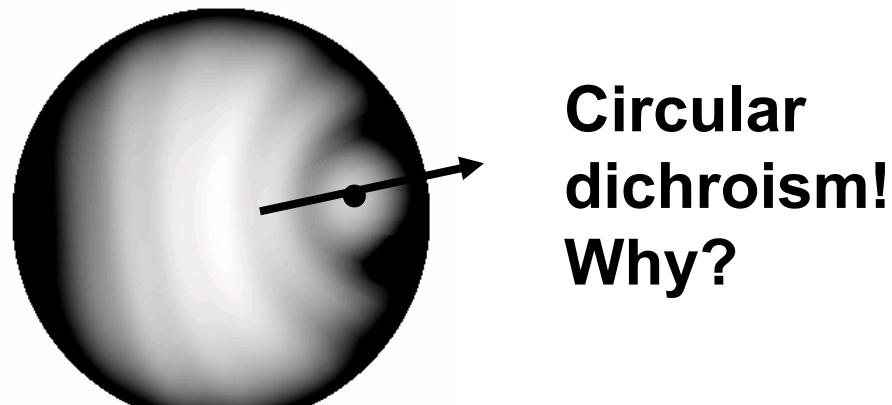
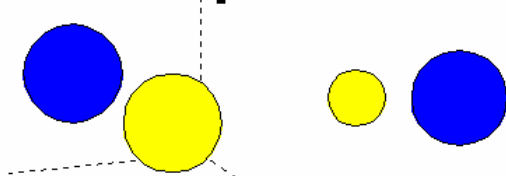
- Valence-band spectra: low-energy UPS limit and high-energy XPS limit
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Circular dichroism in angular distributions: C 1s emission from CO, $E_{\text{kin}} = 200$ eV

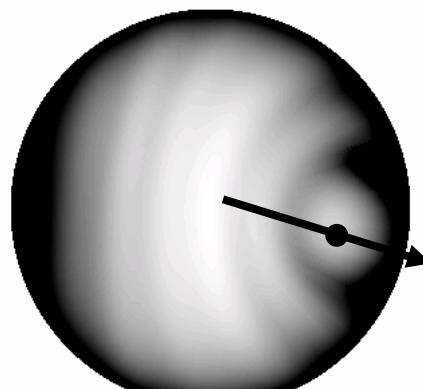
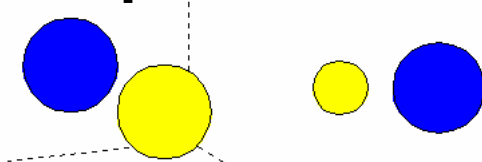
Linear p polarization:



Right circular polarization:

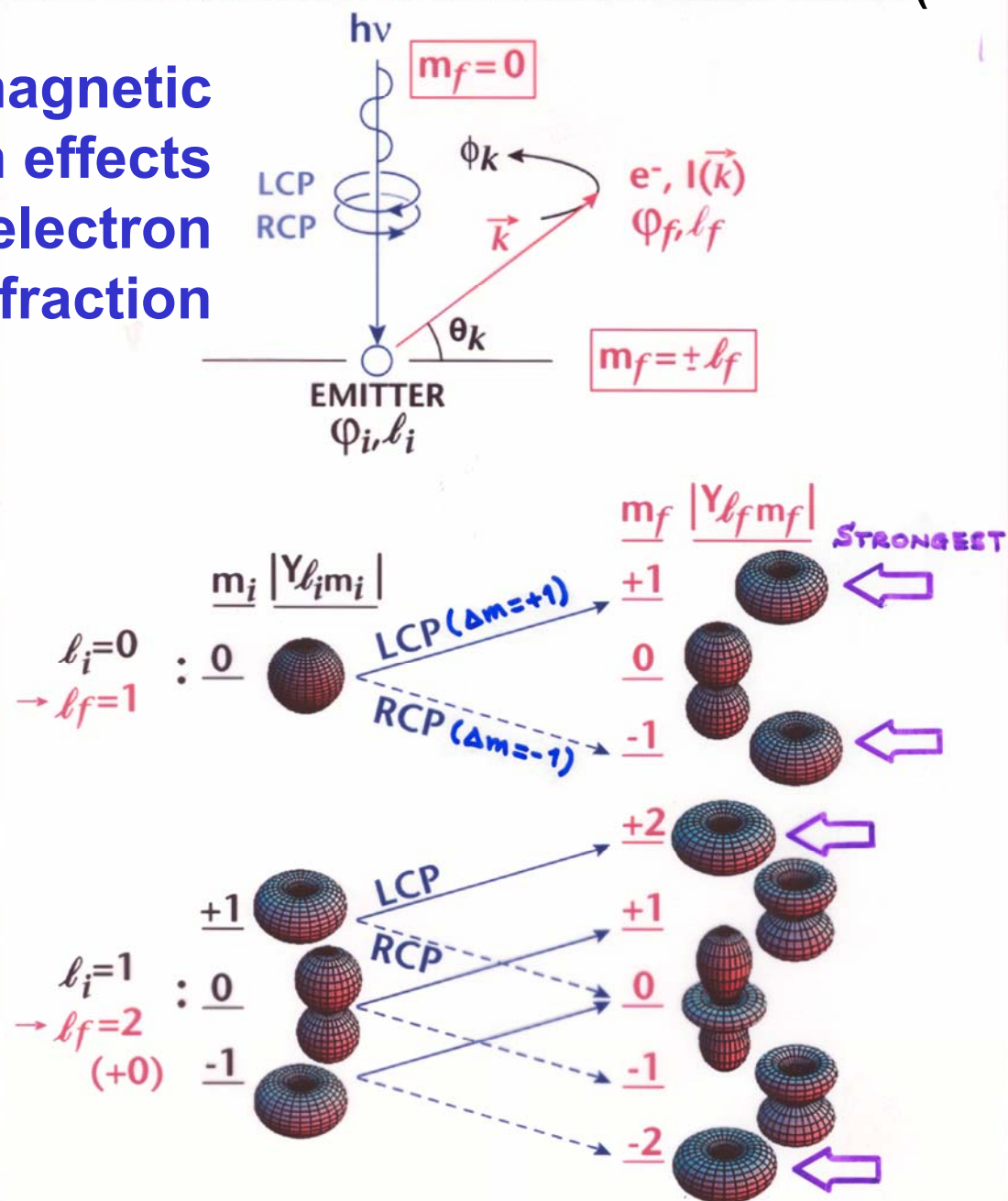


Left circular polarization:



CIRCULAR DICHROISM IN PHOTOELECTRON ANGULAR DISTRIBUTIONS (CDAD)

➡ Non-magnetic dichroism effects due to photoelectron diffraction

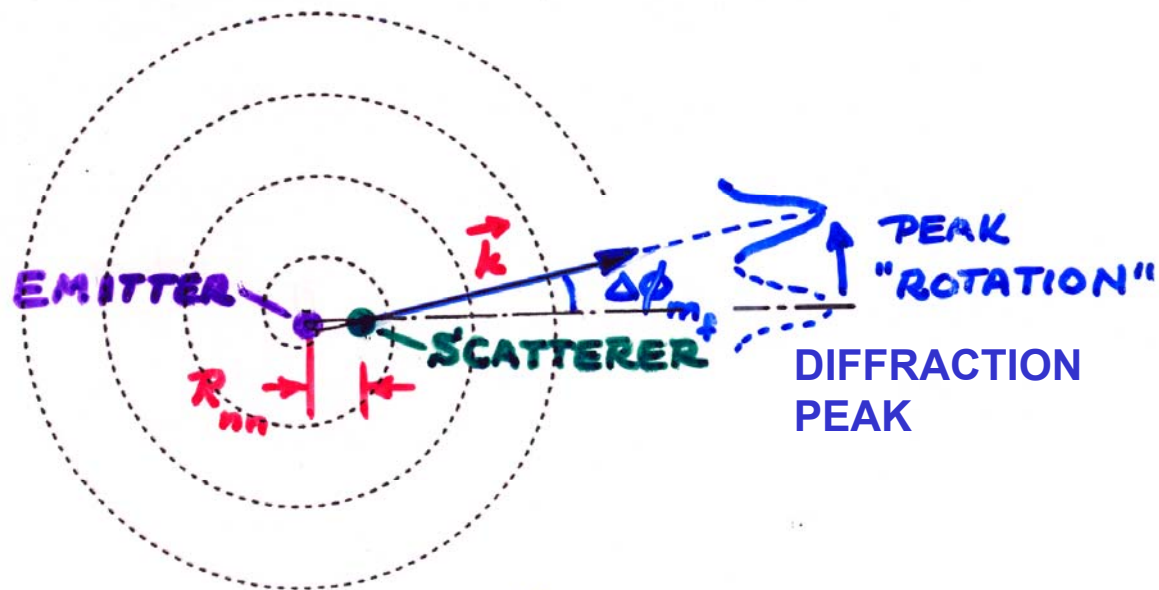


CIRCULAR DICHROISM IN PHOTOELECTRON DIFFRACTION

CONSTANT-PHASE SURFACE OF :

$$\psi_{\text{PHOTOE}^-}(r, \theta, \phi) \propto \frac{e^{ikr}}{r} \sum_{lm} e^{im_f \phi}$$

Z OUT
OF PLANE



$$\Delta\phi_{m_f} = \frac{m_f}{R_{nn, || k_{||}}}$$

$$\overline{m}_f \approx m_{f, \max}$$

DAIMON ET AL.
JPN. J. APPL. PHYS.
32, L1480 ('93)

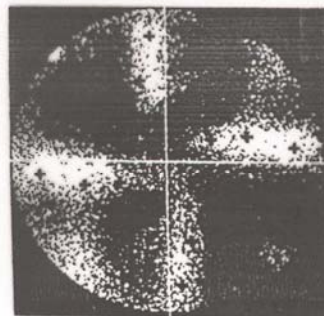
CIRCULAR DICHOISM - NON-MAGNETIC SYSTEMS

Si2p -- 250eV = E_{kin}

EXPERIMENT



(a) LCP



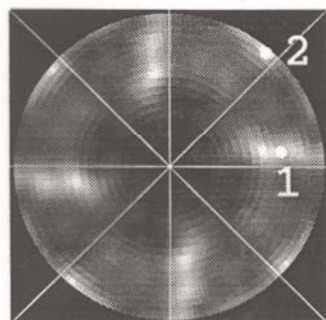
(b) RCP



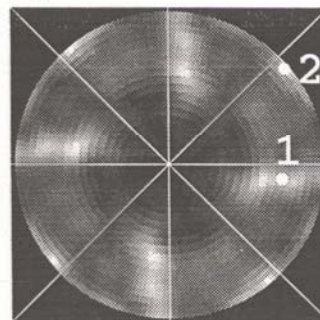
DAIMON ET AL.
JPN. J. APPL. PHYS.
32, L1480 ('93)

THEORY

(c) LCP

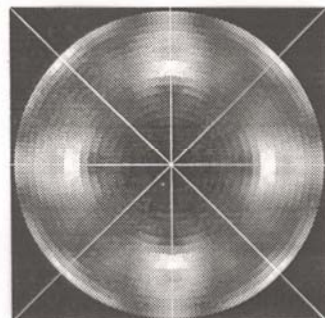


(d) RCP



IKADUWELA ET AL.
P. R. B 50, 6203 ('94)

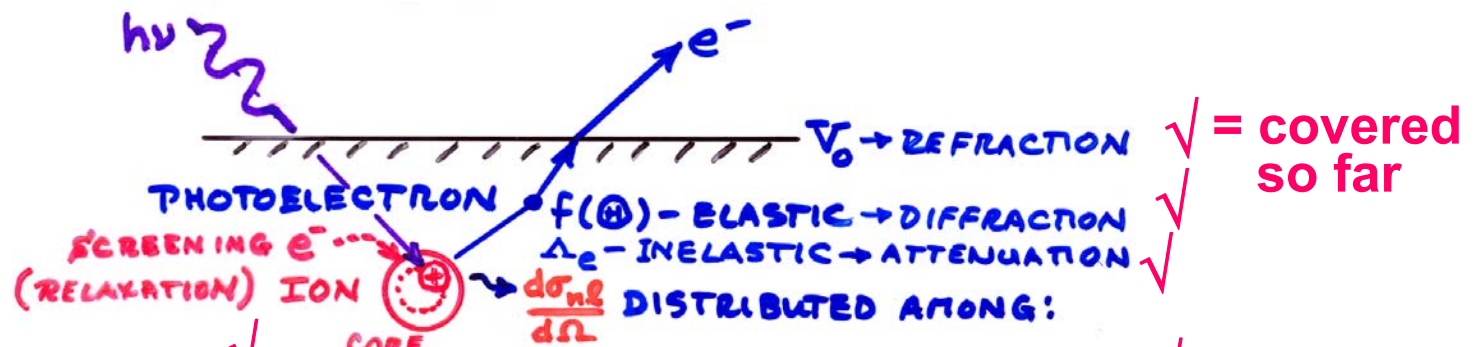
(e) UNPOLARIZED



Outline

- Valence-band spectra: low-energy UPS limit and high-energy XPS limit
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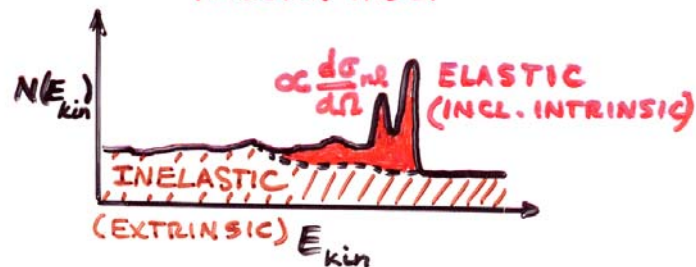


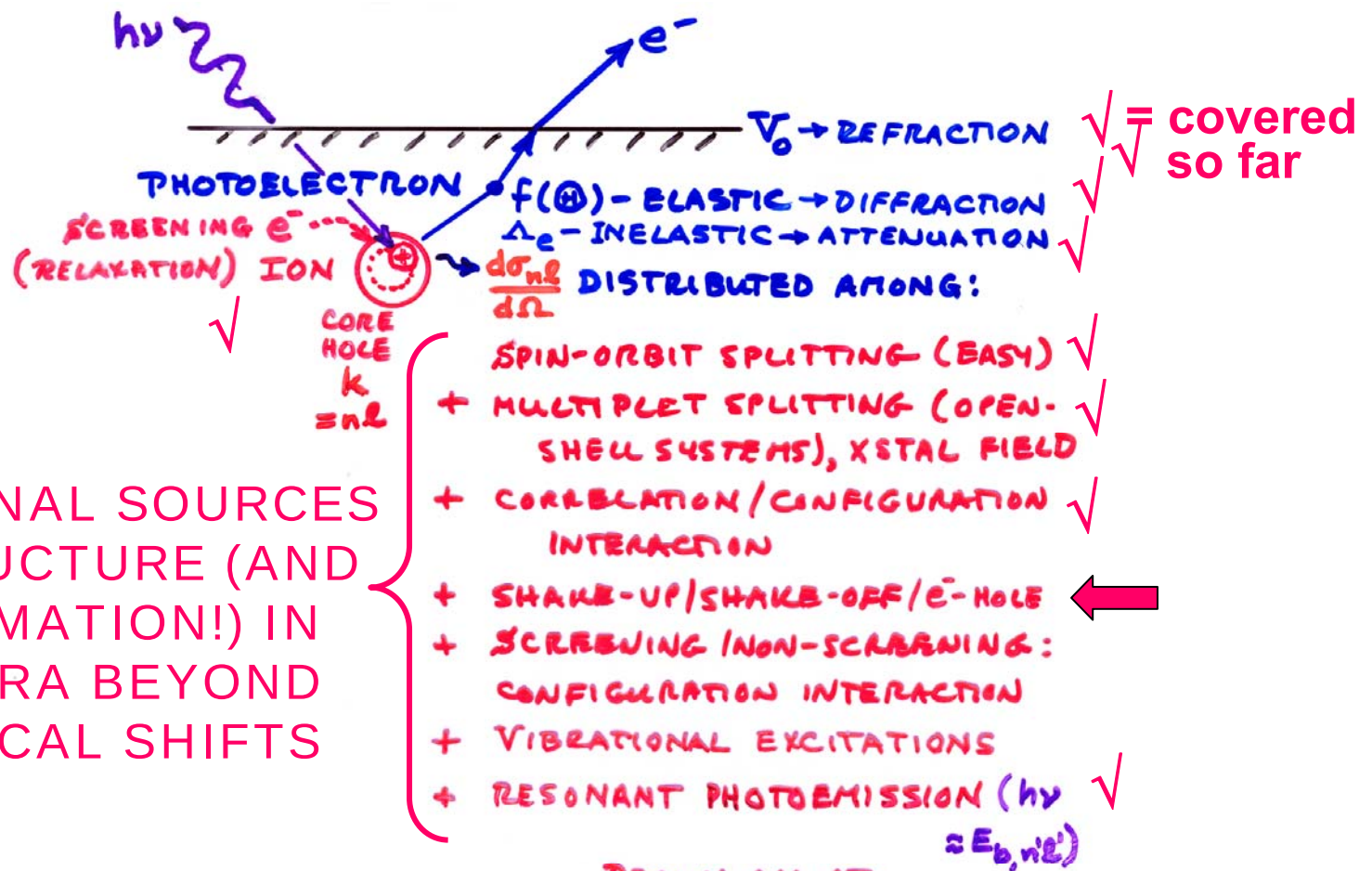


ADDITIONAL SOURCES
OF STRUCTURE (AND
INFORMATION!) IN
SPECTRA BEYOND
CHEMICAL SHIFTS

- + SPIN-ORBIT SPLITTING (EASY) $\checkmark$
- + MULTIPLY SPLITTING (OPEN-SHELL SYSTEMS), XSTAL FIELD $\checkmark$
- + CORRELATION/CONFIGURATION INTERACTION $\checkmark$
- + SHAKE-UP/SHAKE-OFF/ e^- -HOLE
- + SCREENING/NON-SCREENING: CONFIGURATION INTERACTION
- + VIBRATIONAL EXCITATIONS $\checkmark$
- + RESONANT PHOTOEMISSION ($h\nu \approx E_{b,n\ell}$) $\checkmark$

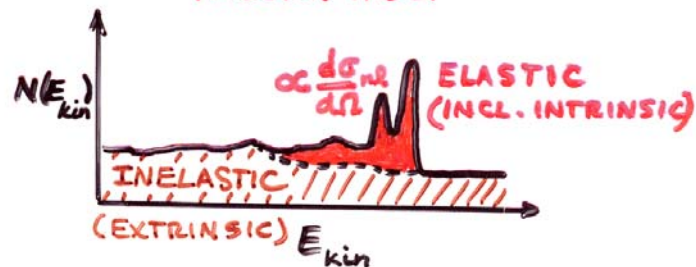
REALLY ALL AT
ONCE, BUT
SUM RULES +
THEORY HELP

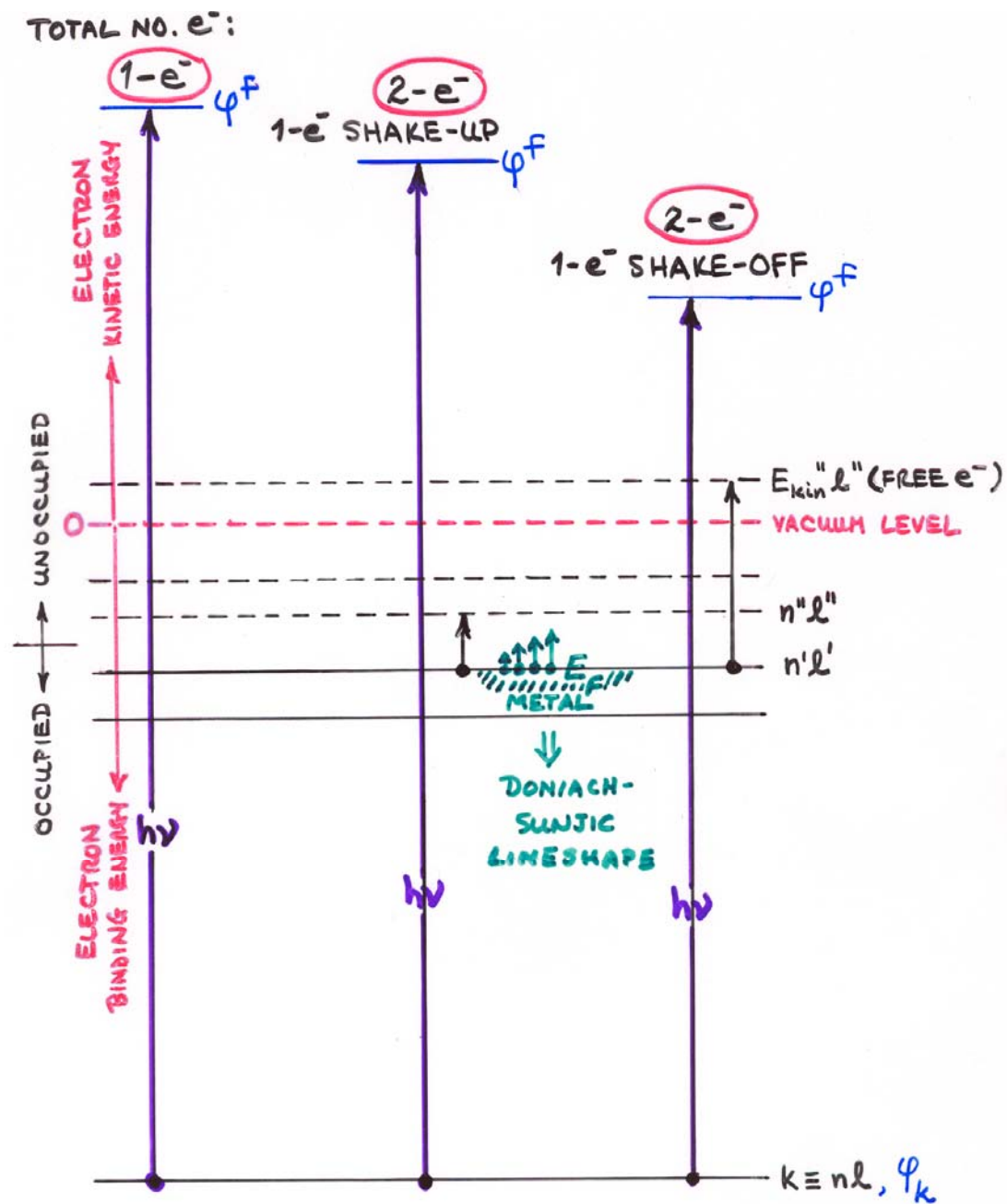




ADDITIONAL SOURCES OF STRUCTURE (AND INFORMATION!) IN SPECTRA BEYOND CHEMICAL SHIFTS

REALLY ALL AT ONCE, BUT SUM RULES + THEORY HELP





MULTIELECTRON EFFECTS IN CORE EMISSION

INTENSITIES IN PHOTOELECTRON SPECTRA:

- GENERAL: FINAL STATE K (k -SUBSHELL + ALL OTHER DESIG.)

$$\text{INT.}_K \propto |\hat{e} \cdot \langle \Psi_{\text{tot}}^f(N, K) | \sum_{i=1}^N \vec{r}_i | \Psi_e^i(N) \rangle|^2 \quad (\text{DIPOLE APPROX.})$$

- BORN-OPPENHEIMER: e^- 's FAST, VIBRATIONS SLOW

$$\text{INT.}_K \propto |\langle \Psi_{\text{vib}, v}^f | \Psi_{\text{vib}, v}^i \rangle|^2 |\hat{e} \cdot \langle \Psi_e^f(N, K) | \sum_{i=1}^N \vec{r}_i | \Psi_e^i(N) \rangle|^2$$

FRANCK-CONDON FACTOR

- SUDDEN APPROXIMATION: $\Psi_K \rightarrow \Psi_f = \text{PHOTO}^-$ (FAST)



$$\text{INT.}_K \propto |\langle \Psi_{\text{vib}, v}^f | \Psi_{\text{vib}, v}^i \rangle|^2 |\langle \Psi_e^f(N-1, K) | \Psi_e^i(N-1, K) \rangle|^2$$

$$|\hat{e} \cdot \langle \Psi_f | \vec{r} | \Psi_K \rangle|^2$$

SAME SUBSHELL COUPLING +
TOTAL $L, S \rightarrow$ "MONOPOLE"

$$\rightarrow \text{NORMAL } \frac{d\sigma_K}{d\Omega}$$

- SLATER DETS. FOR $\Psi_e^f = \det(\psi'_1, \psi'_2, \dots, \psi'_{k-1}, \psi'_{k+1}, \dots, \psi'_N)$

$$\Psi_K = \det(\psi_1, \psi_2, \dots, \psi_{k-1}, \psi_{k+1}, \dots, \psi_N)$$

$$\text{INT.}_K \propto |\langle \Psi_{\text{vib}, v}^f | \Psi_{\text{vib}, v}^i \rangle|^2 |\langle \psi'_1 | \psi_1 \rangle|^2 |\langle \psi'_2 | \psi_2 \rangle|^2 \dots$$

$$|\langle \psi'_{k-1} | \psi_{k-1} \rangle|^2 |\langle \psi'_{k+1} | \psi_{k+1} \rangle|^2 \dots |\langle \psi'_N | \psi_N \rangle|^2$$

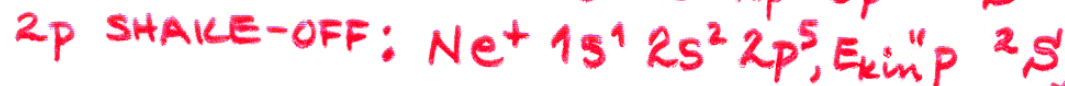
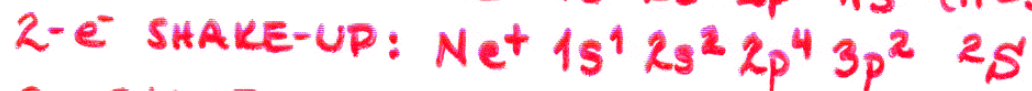
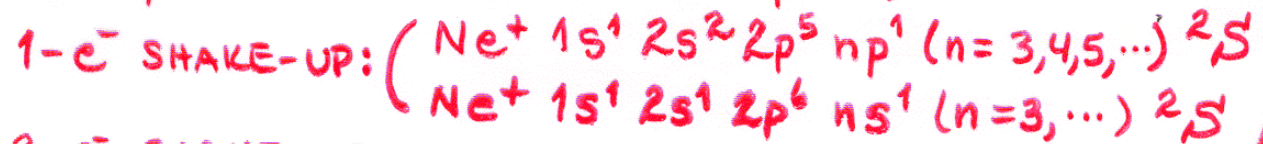
$$|\hat{e} \cdot \langle \Psi_f | \vec{r} | \Psi_K \rangle|^2$$

$1e^-$ DIPOLE $\rightarrow d\sigma/d\Omega$

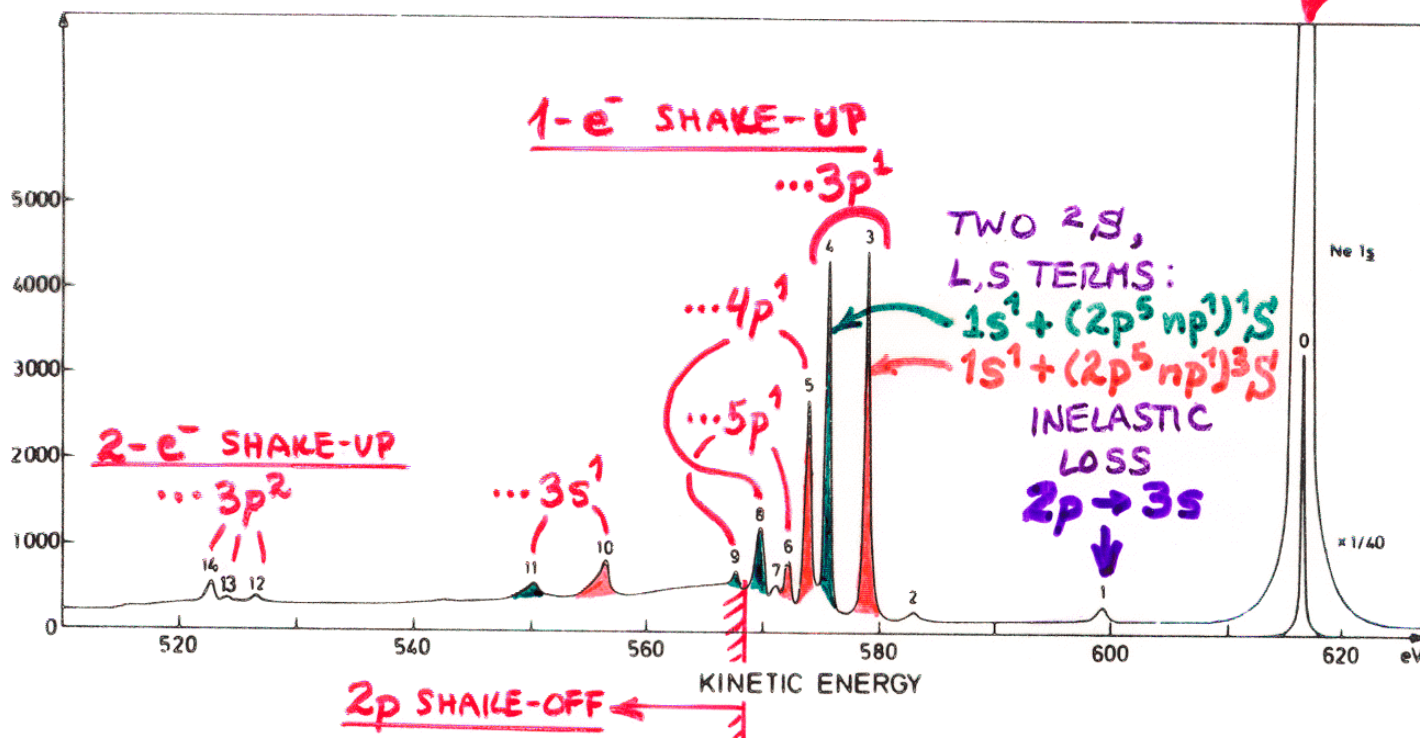
($N-1$) e^- SHAKE-UP/
SHAKE-OFF $\rightarrow$
"MONOPOLE"

- PLUS DIFFRACTION EFFECTS IN Ψ_f ESCAPE

NEON 1S SHAKE-UP / SHAKE-OFF:

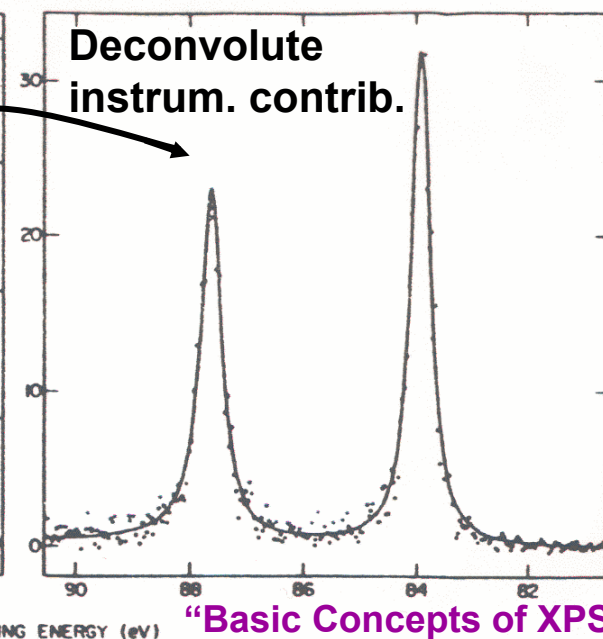
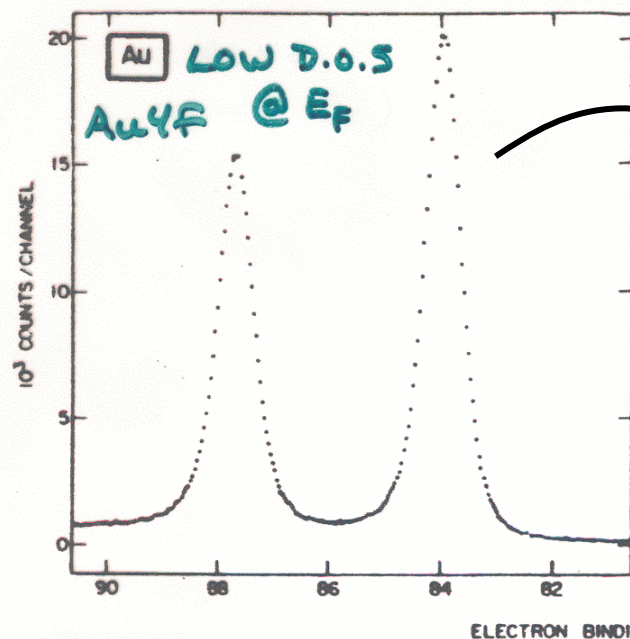
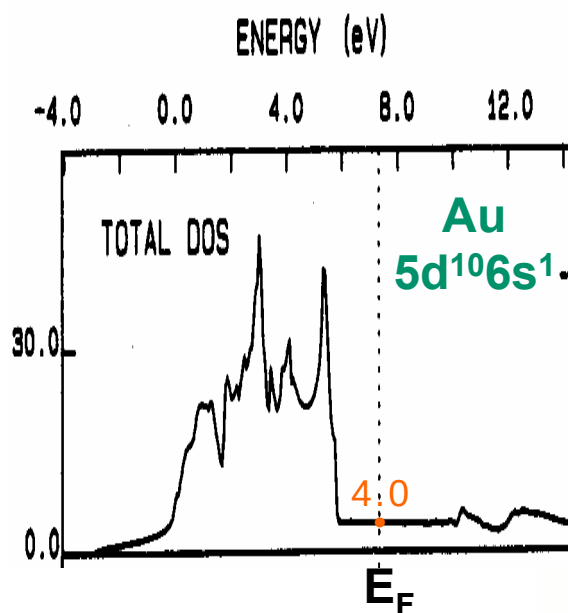


"Basic Concepts of XPS"
Figure 36

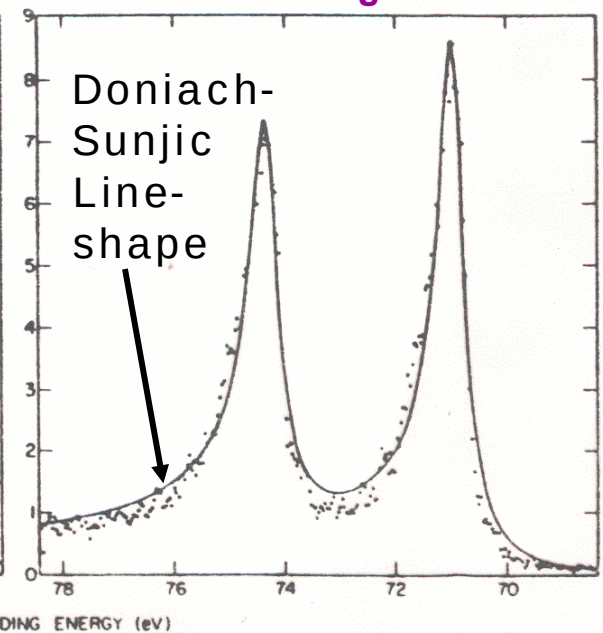
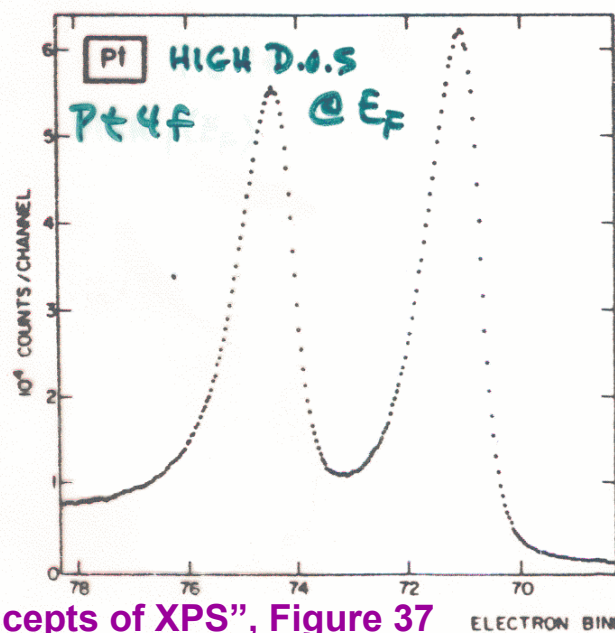
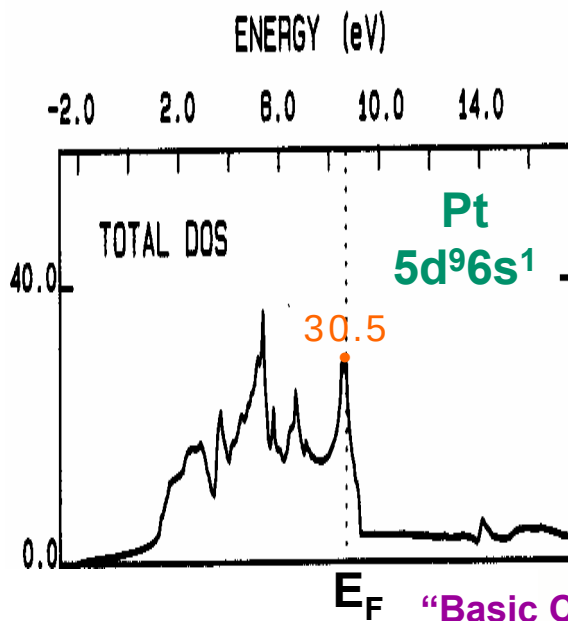


OVERALL: ~12% SHAKE-UP + 16% SHAKE-OFF ≈ 28% OF EVENTS

BAND THEORY—D.O.S:



“Basic Concepts of XPS”
Figure 10



“Basic Concepts of XPS”, Figure 37

TWO SUDDEN-APPROXIMATION

SUM RULES:

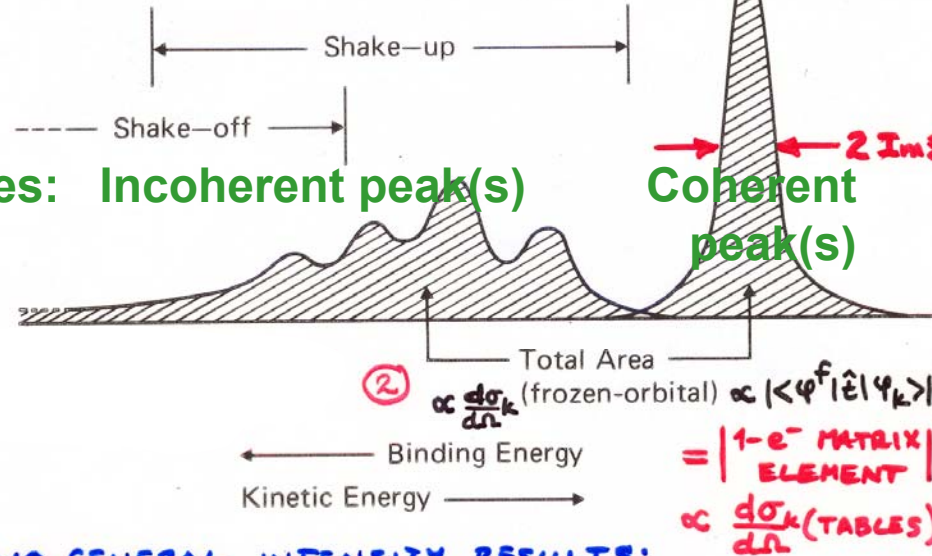
$$\textcircled{1} \left\{ \begin{array}{l} \text{AVERAGE} \\ \text{BINDING} \\ \text{ENERGY} \end{array} \right\} = \frac{\sum_{j=1}^{\text{ALL}} I_j E_b^V(k)_j}{\sum_{j=1}^{\text{ALL}} I_j} = \text{KOOPMANS' } -\epsilon_k$$

Ground-
State
of Ion =
Adiabatic
peak

$E_b^V(k)_1$

$\approx \delta E_{\text{relax}}$
 $= \text{Re } \Sigma$

$\Sigma = \text{many-body}$
"self energy"
 $= \text{Re } \Sigma + i \text{Im } \Sigma$



In valence-band studies: Incoherent peak(s)

Coherent
peak(s)

$$\Delta E \tau_{\text{lifetime}} \approx \hbar/2$$

TWO GENERAL INTENSITY RESULTS:

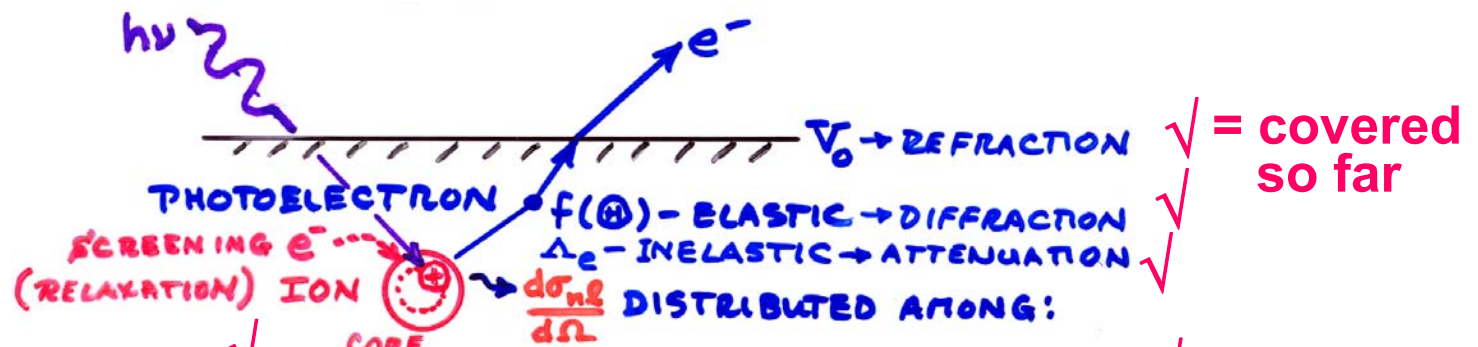
$$\textcircled{1} I_j \propto |\langle \varphi^f(1) | \hat{e} | \varphi_k(1) \rangle|^2 |\langle \Psi^f(N-1, j) | \Psi_R(N-1) \rangle|^2$$

$K e^- \text{ MISSING}$

Figure 8 -- Schematic illustration of a photoelectron spectrum involving shake-up and shake-off satellites. The weighted average of all binding energies yields the Koopmans' Theorem binding energy $-\epsilon_k$ (sum rule (77)), and the sum of all intensities is proportional to a frozen-orbital cross section σ_k (sum rule (78)). The adiabatic peak corresponds to formation of the ground-state of the ion ($E_b(k)_1 \equiv E_b(K=1)$).

$$\textcircled{2} \left(\begin{array}{l} \text{TOTAL SHAKE-UP} \\ + \text{SHAKE-OFF} \end{array} \right) = 1 - |\langle \Psi^f(N-1, 1) | \Psi_R(N-1) \rangle|^2$$

$\approx 15-25\% \text{ FOR ATOMS/MOLEC.}$

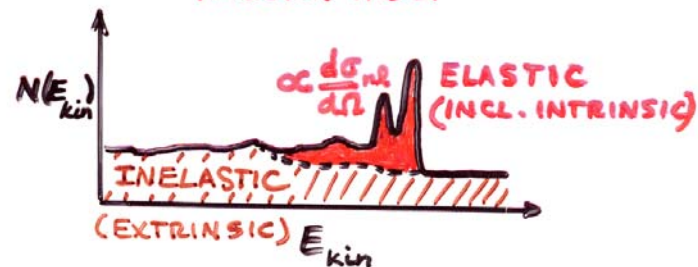


= covered so far

ADDITIONAL SOURCES OF STRUCTURE (AND INFORMATION!) IN SPECTRA BEYOND CHEMICAL SHIFTS

- + SPIN-ORBIT SPLITTING (EASY)
- + MULTIPLY SPLITTING (OPEN-SHELL SYSTEMS), XSTAL FIELD
- + CORRELATION / CONFIGURATION INTERACTION
- + SHAKE-UP / SHAKE-OFF / e^- -HOLE
- + SCREENING / NON-SCREENING: CONFIGURATION INTERACTION
- + VIBRATIONAL EXCITATIONS
- + RESONANT PHOTOEMISSION ($h\nu \approx E_{b,n\ell}$)

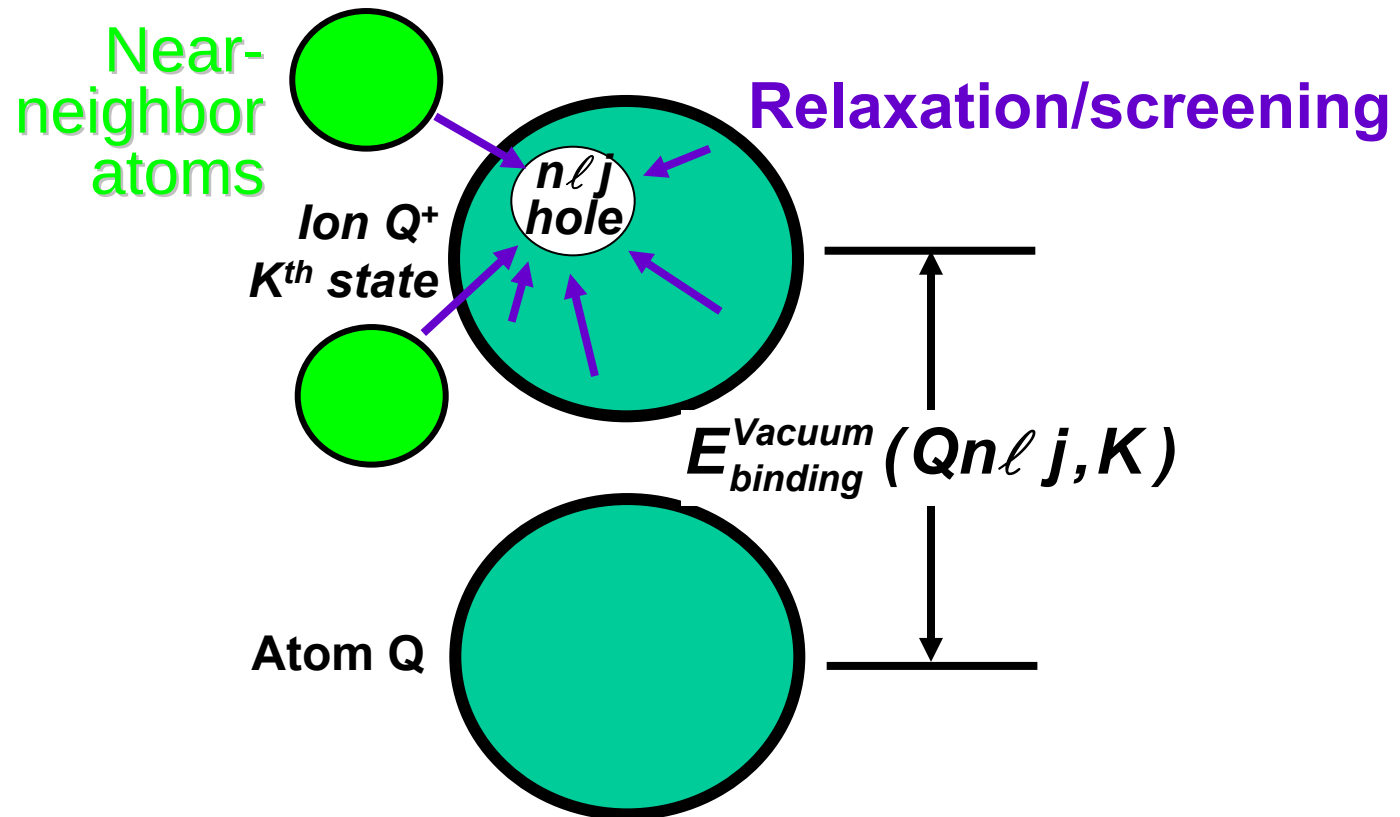
REALLY ALL AT ONCE, BUT SUM RULES + THEORY HELP

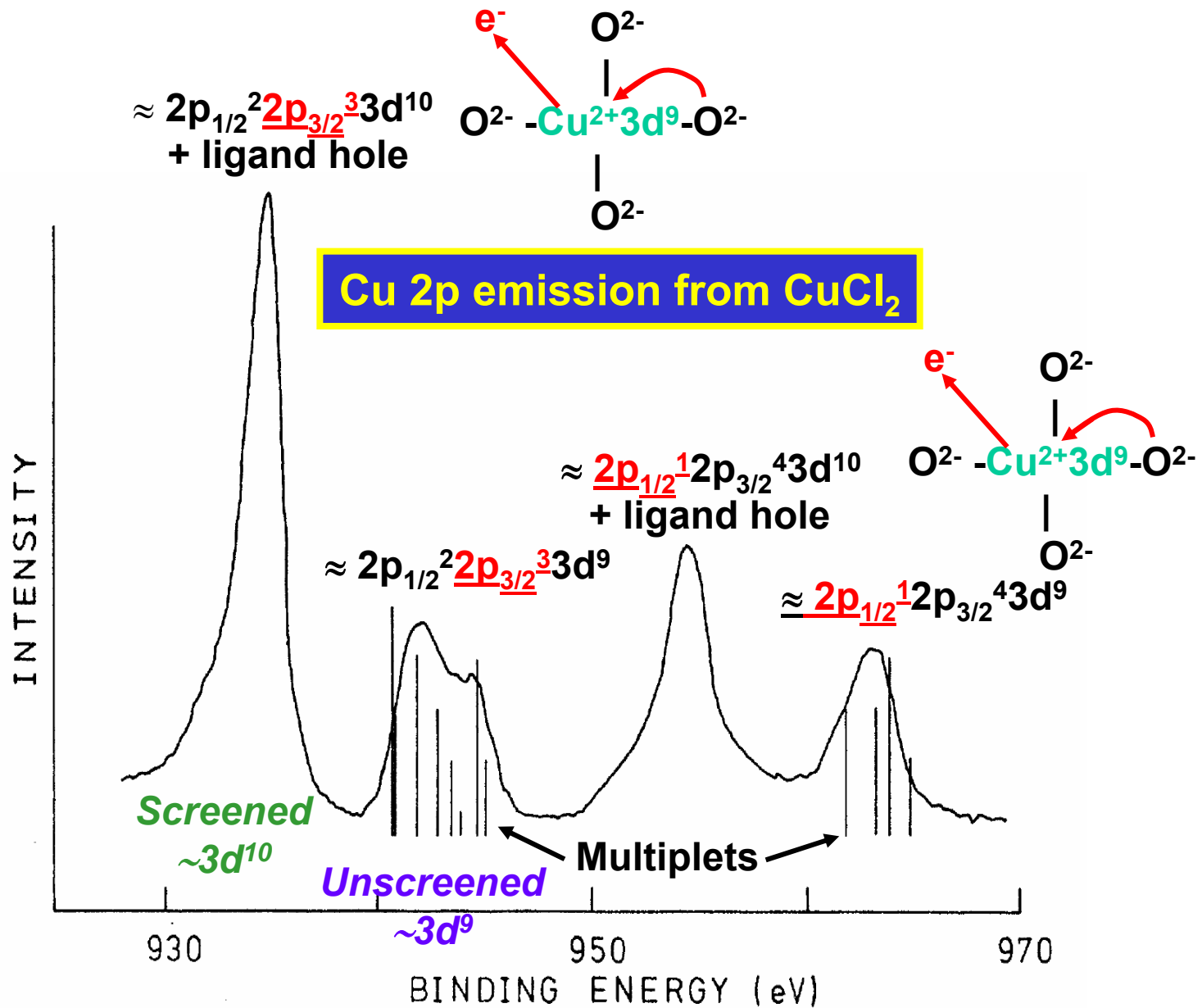


Basic energetics—Many e⁻ & many atom picture

$$h\nu = E_{\text{binding}}^{\text{Vacuum}} + E_{\text{kinetic}} = E_{\text{binding}}^{\text{Fermi}} + \varphi_{\text{spectrometer}} + E_{\text{kinetic}}$$

$$E_{\text{binding}}^{\text{Vacuum}}(Qn\ell j, K) = E_{\text{final}}(N-1, Qn\ell j \text{ hole}, K) - E_{\text{initial}}(N)$$

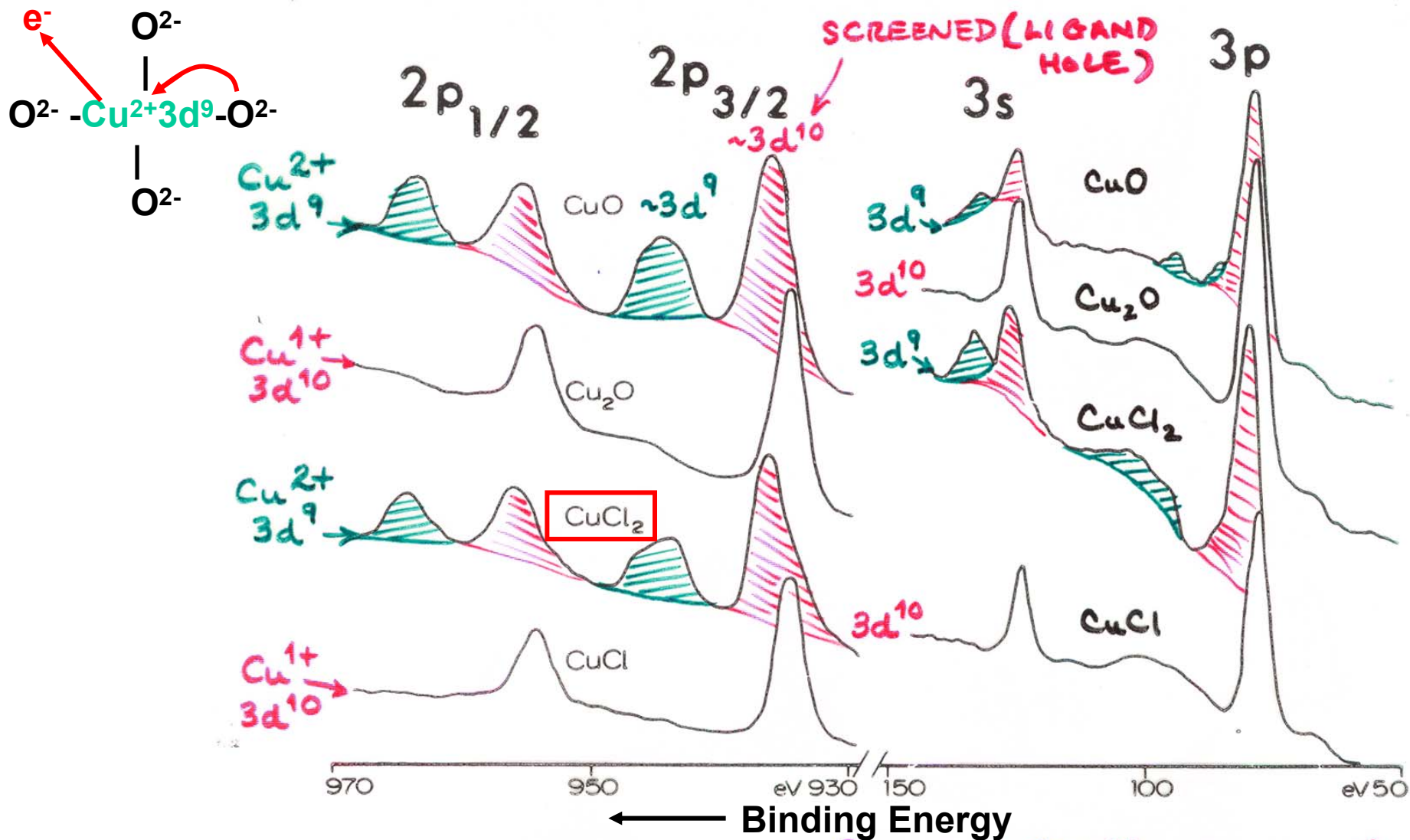




$$\psi_{final,K}(N-1) = C_{1,K}(2p_{1/2}^2 \underline{2p_{3/2}}^3 3d^{10} + C_{\ell} \text{ hole}) + C_{2,K}(2p_{1/2}^2 \underline{2p_{3/2}}^3 3d^9)$$

Van der Laan et al., Phys. Rev. B 23 (1981) 4369

SATELLITES & CHARGE-TRANSFER SCREENING



"Basic Concepts of XPS" **ACTUAL FINAL STATE** $\Psi \approx C_1 \phi_1 (3d^{10} - \text{SCREENED}) + C_2 \phi_2 (3d^9 - \text{UNSCREENED})$

Figure 38

Screening depends on ionicity/covalency → satellite intensities can be used to measure interaction parameters

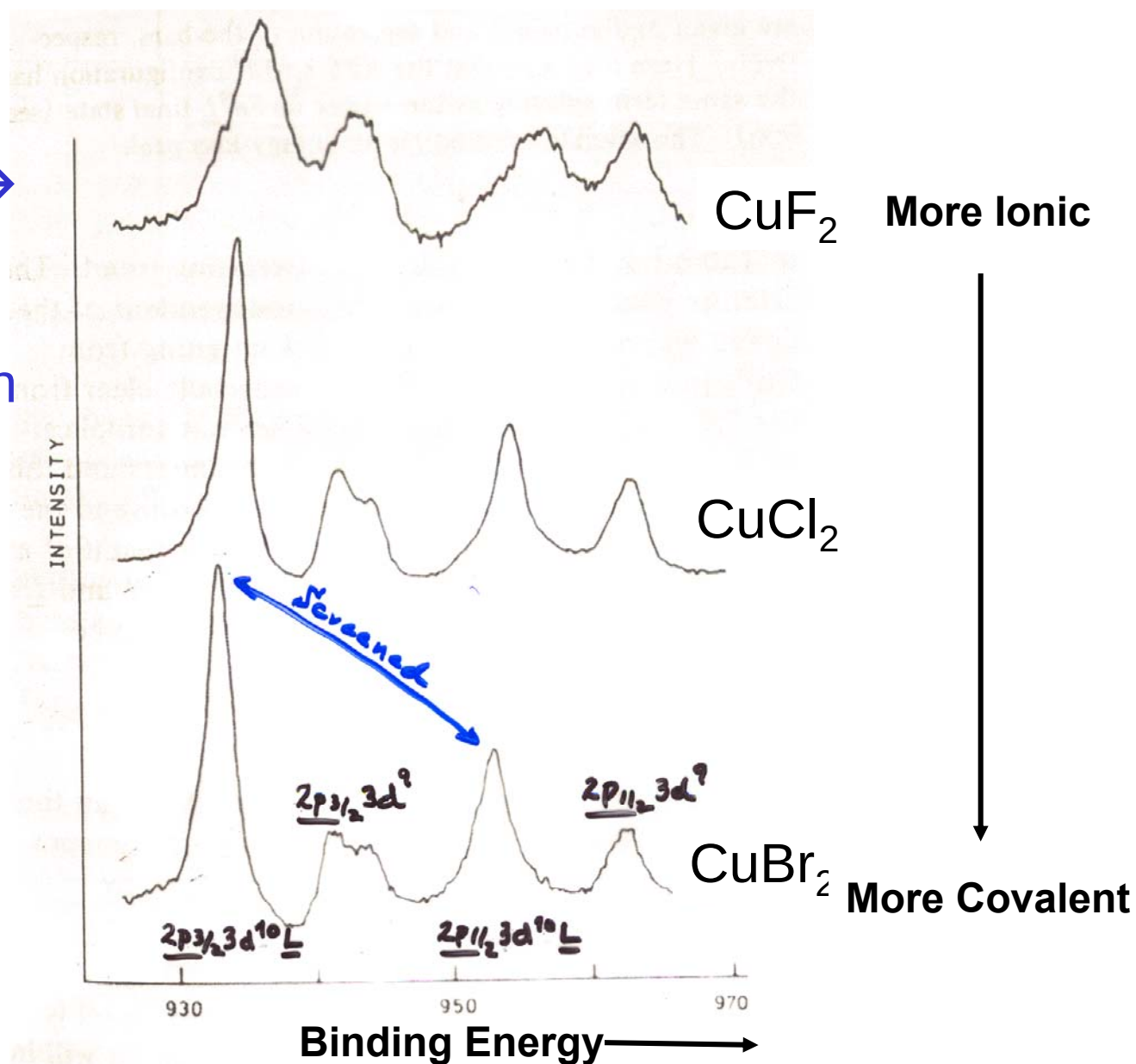


FIG. 1. Cu 2p photoelectron spectra of Cu dihalides. The lines leading to a final state with a ligand hole ($\underline{L}$) show a chemical shift.

Screening
depends on
Ionicity/covalency →
satellite intensities
can be used to
measure interaction
parameters

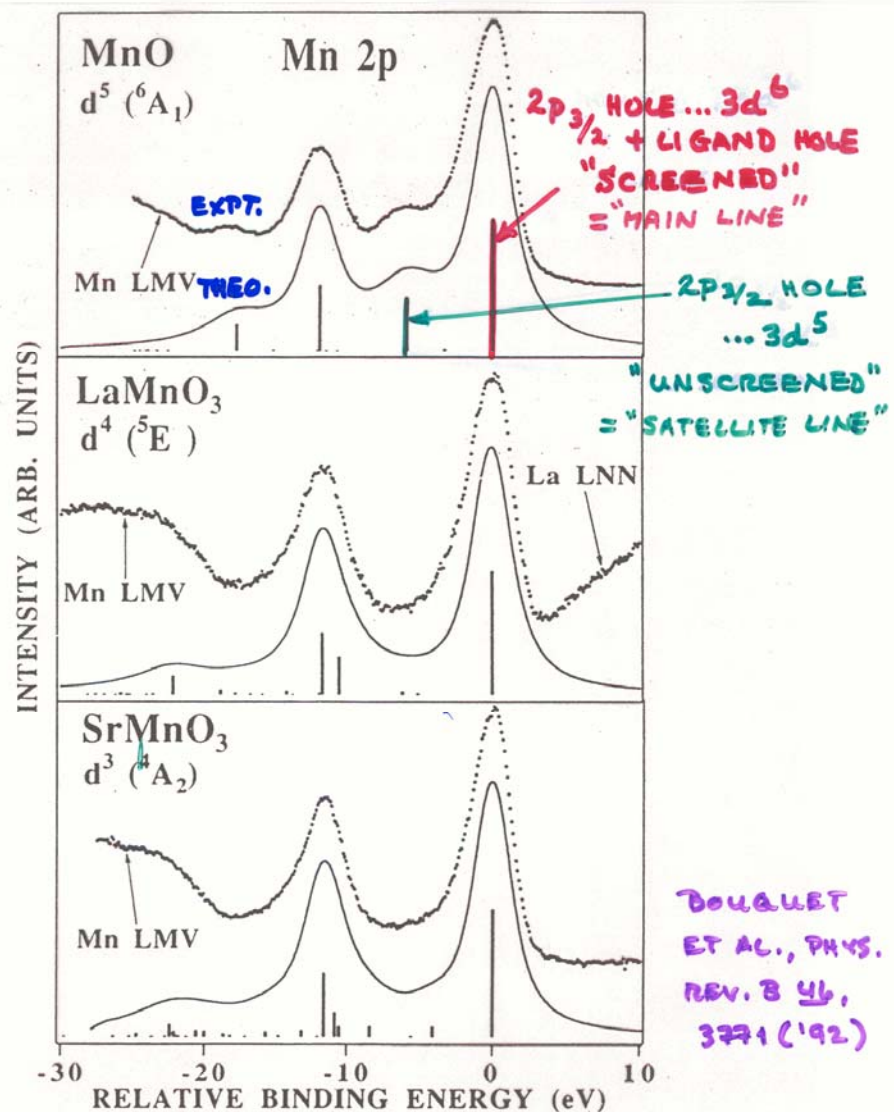
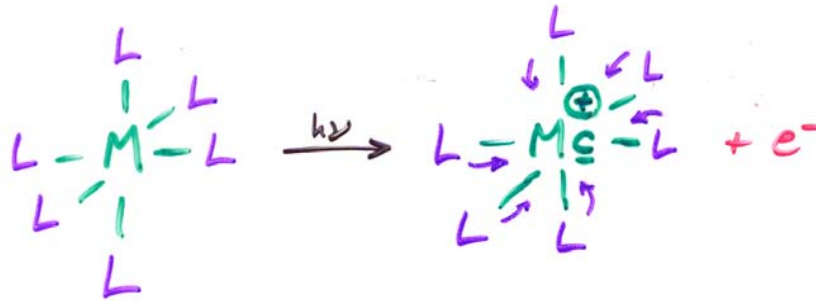


FIG. 1. Theoretical $2p$ core-level XPS spectra (solid line) compared with experimental data (dots) after background subtraction for Mn cations with varying valence. Emission due to the Mn LMV Auger peak is observed on the high-binding-energy side of the $2p_{1/2}$ spin-orbit peak, partially obscuring the $2p_{1/2}$ satellite structure.

CONFIGURATION INTERACTION APPROACH TO SCREENING IN TRANSITION-METAL (RARE-EARTH) COMPOUNDS:

(SUGANO, LARSSON → SAWATZKY, VANDER LAAN, FUTIMORI, OH, ET AL.)



$\underline{c}$ = CORE HOLE ON METAL

$\underline{l}$ = VALENCE (?) HOLE ON LIGAND

$$\psi_i = a_0 |d^n\rangle + \sum_m a_m |d^{(n+m)} \underline{l}^m\rangle$$

$$\psi_f = b_0 |\underline{c} d^n\rangle + \sum_m b_m |\underline{c} d^{(n+m)} \underline{l}^m\rangle$$

WITH INTERACTIONS OF:

$10Dq$ = CRYSTAL FIELD (OFTEN NEGLECTED)

Δ = LIGAND-TO-METAL CHARGE TRANSF. ENERGY
 $= E(d^{n+1} \underline{l}) - E(d^n)$

U = d-d COULOMB REPULSION ENERGY
 $= E(d^{n-1}) + E(d^{n+1}) - 2E(d^n)$

T = LIGAND p-TO-METAL d HYBRIDIZATION
 $= \langle d_\alpha | \hat{H} | p_\alpha \rangle$ (α = SAME SYMMETRY)

Q = CORE-HOLE-TO-d INTERACTION: $\langle \underline{c} | \hat{H} | d \rangle \approx J_{cd}$

WITH INTENSITIES FROM SUDDEN APPROX.

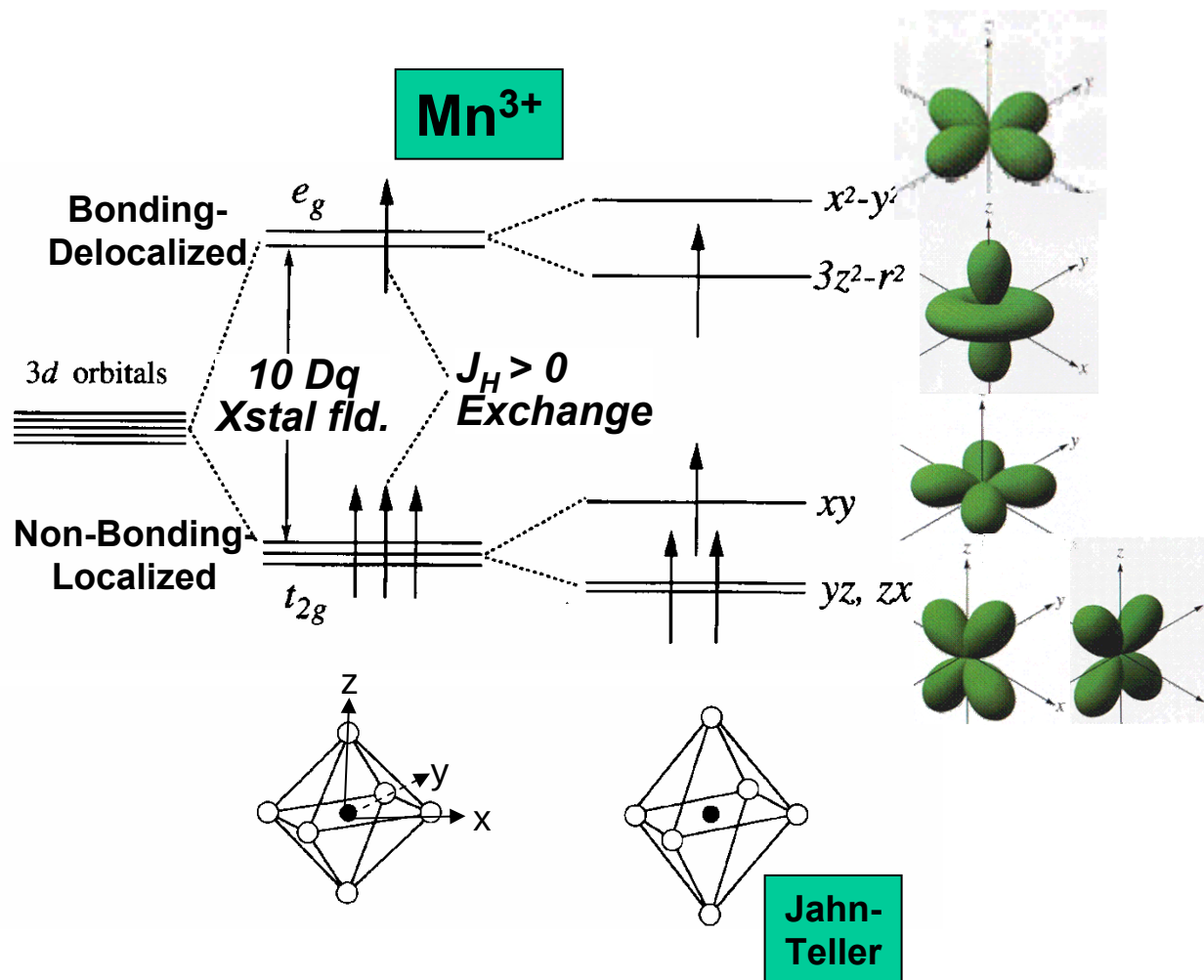
AS:

$$I(E_{kin}) \propto \sum_{f,k} |\langle \psi_f(N-1, k) | \psi_i(N-1, k) \rangle|^2 \delta(h\nu - E_f - E_{kin})$$

$\underline{f} = \underline{c}$ = CORE HOLE

WHERE: $\psi_R(N-1, k) = \psi_i(N \text{ WITH } k \text{ HOLE} = \underline{c})$

E.g.—Crystal field in Mn^{3+} with negative octahedral ligands



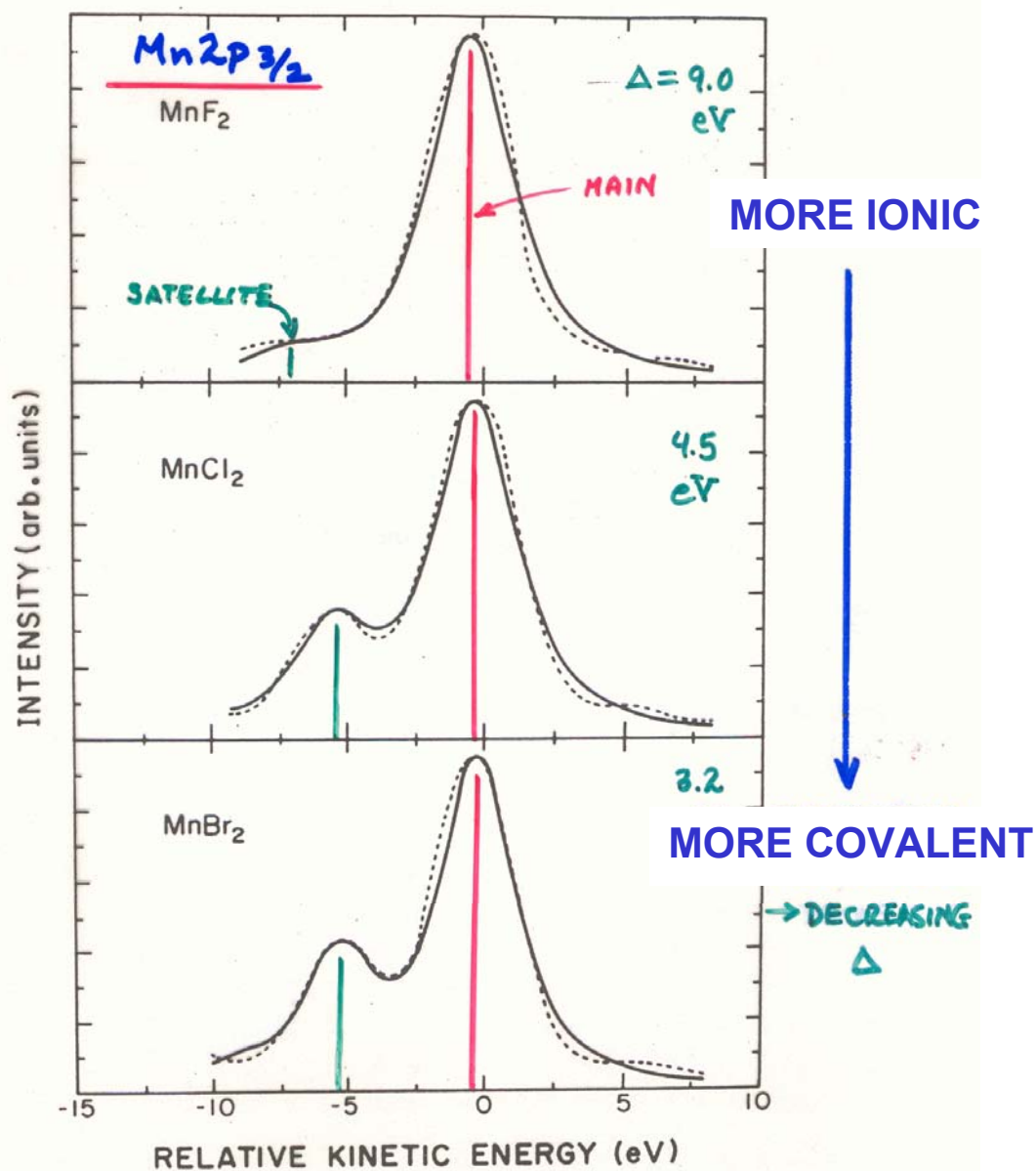
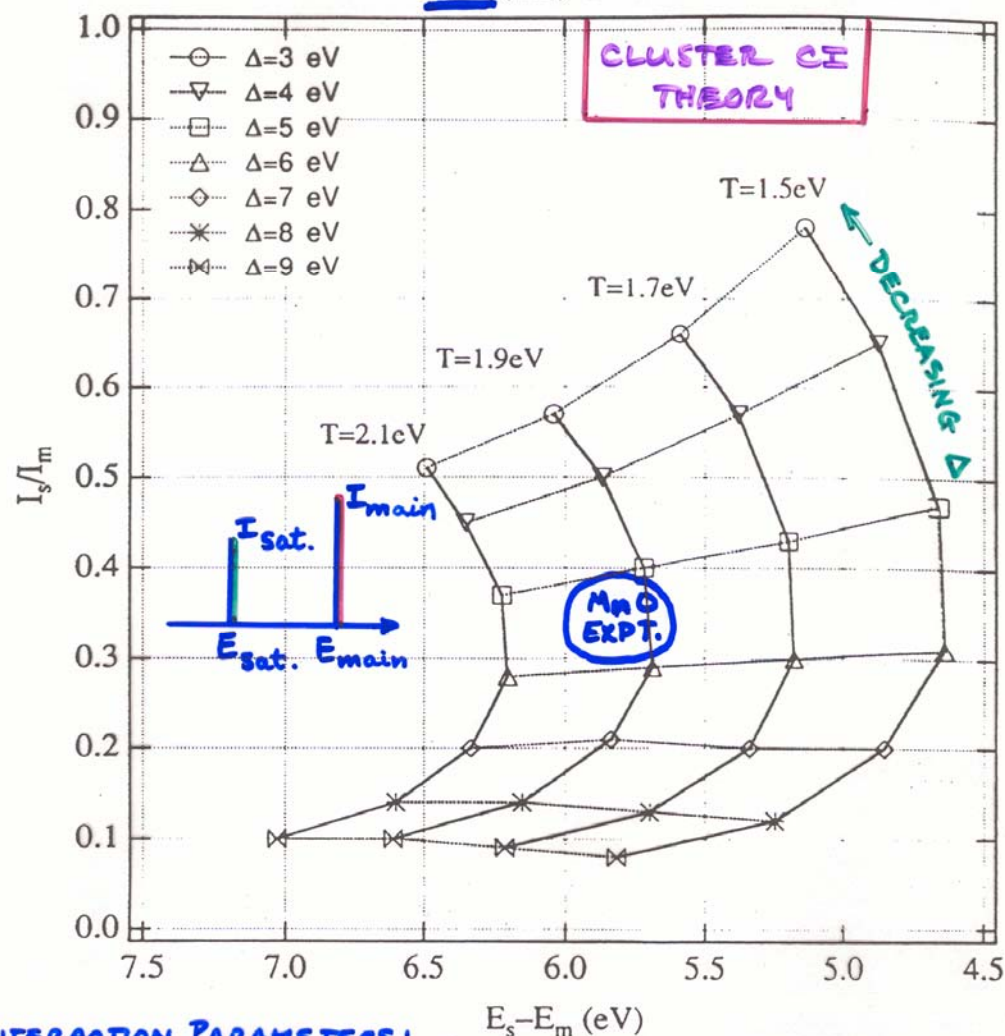


FIG. 6. Fits of the cluster model results with the experimental $2p_{3/2}$ spectra of the manganese dihalides. The parameters used are listed in Table II. A Lorentzian broadening is 2.6–3.0 eV, and a Gaussian broadening of 1.2 eV (FWHM) was used.

ANALYSIS VIA ANDERSON IMPURITY MODEL

$\text{Mn}^{2+}(\text{HS})$ $U=6.0 \text{ eV}$



INTERACTION PARAMETERS:

U = 3d-3d COULOMB REPULSION ENERGY

Δ = LIGAND-TO-METAL CHARGE TRANSFER ENERGY

T = LIGAND p - METAL 3d HYBRIDIZATION ENERGY

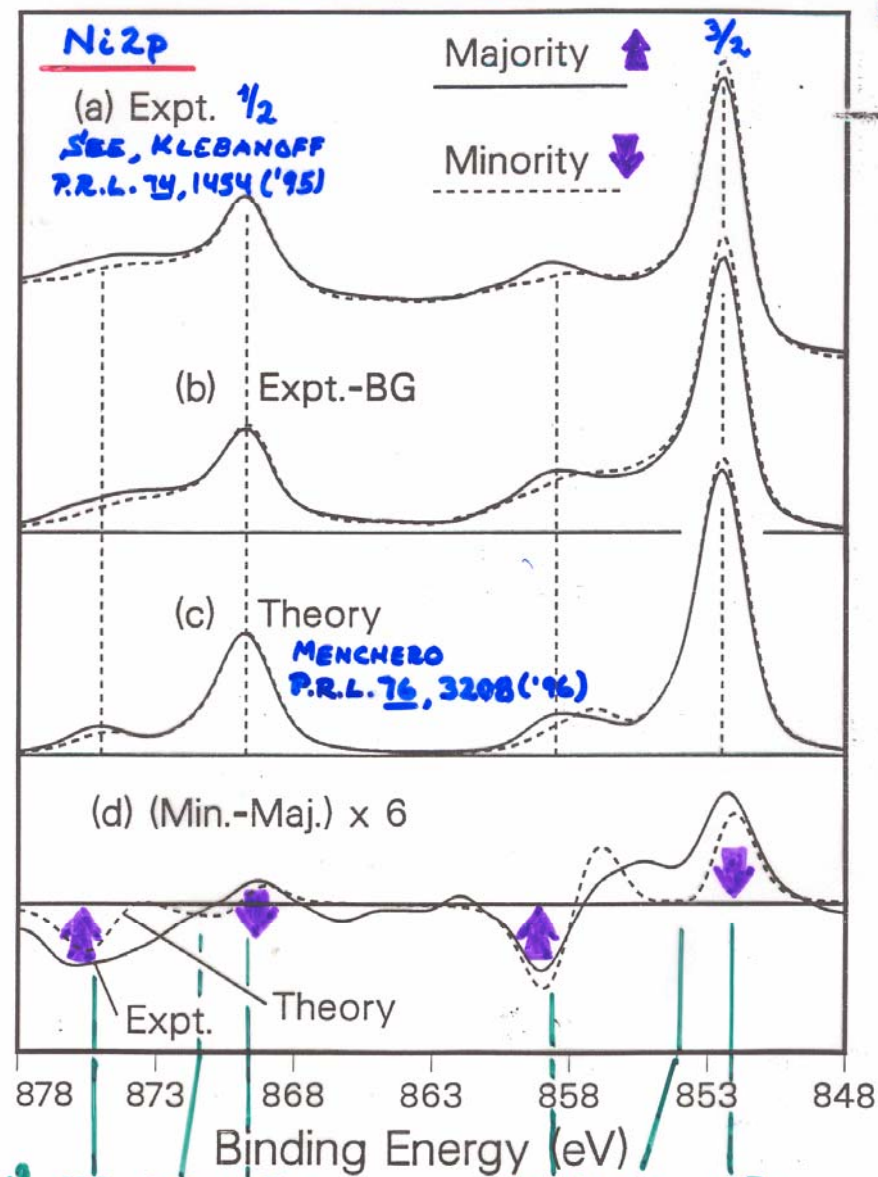
Q = CORE HOLE-3d COULOMB

BOUQUET ET AL.,
J. EL. SP. 82, 87 (196)

SPIN-ORBIT SPLITTING + MULTIPLETS +
SCREENING IN A METAL : Ni - INITIAL CONFIG. :

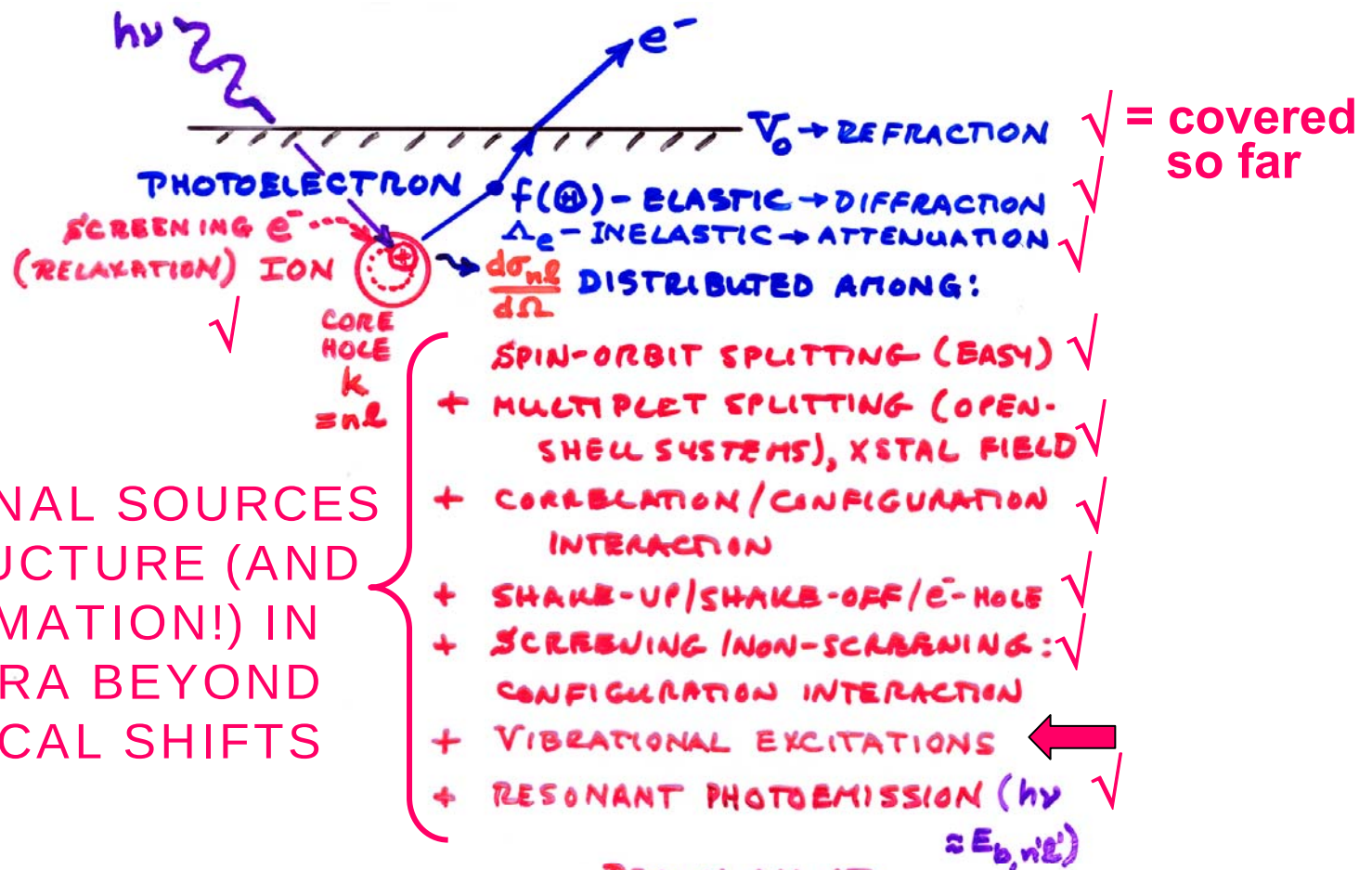
$\sim 15\% 3d^8$
 $43\% 3d^9$
 $42\% 3d^{10}$

Counts (Arb. Units)



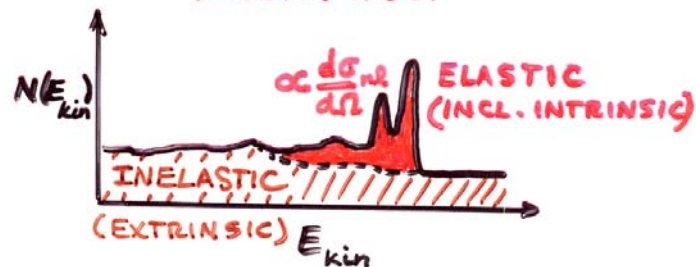
FINAL CONFIG.:

$3d^9$	65%	15%	10%	"UNSCREENED"	65%	15%	10%	"SCREENED"
$3d^{10}$	35%	85%	90%		35%	85%	90%	



ADDITIONAL SOURCES OF STRUCTURE (AND INFORMATION!) IN SPECTRA BEYOND CHEMICAL SHIFTS

REALLY ALL AT ONCE, BUT SUM RULES + THEORY HELP



INTENSITIES IN PHOTOELECTRON SPECTRA:

- GENERAL: FINAL STATE K (k -SUBSHELL + ALL OTHER DESIG.)

$$\text{INT.}_K \propto |\hat{e} \cdot \langle \Psi_{\text{tot}}^f(N, K) | \sum_{i=1}^N \vec{r}_i | \Psi_e^i(N) \rangle|^2 \quad (\text{DIPOLE APPROX.})$$

- BORN-OPPENHEIMER: e^- 's FAST, VIBRATIONS SLOW

$$\text{INT.}_K \propto \underbrace{|\langle \Psi_{\text{vib}, v}^f | \Psi_{\text{vib}, v}^i \rangle|^2}_{\text{FRANCK-CONDON FACTOR}} |\hat{e} \cdot \langle \Psi_e^f(N, K) | \sum_{i=1}^N \vec{r}_i | \Psi_e^i(N) \rangle|^2$$

- SUDDEN APPROXIMATION: $\Psi_K \rightarrow \Psi_f = \text{PHOTO}^-$ (FAST)



$$\text{INT.}_K \propto |\langle \Psi_{\text{vib}, v}^f | \Psi_{\text{vib}, v}^i \rangle|^2 |\langle \Psi_e^f(N-1, K) | \Psi_e^i(N-1, K) \rangle|^2$$

$$|\hat{e} \cdot \langle \psi_f | \vec{r} | \psi_K \rangle|^2$$

SAME SUBSHELL COUPLING +
TOTAL $L, S \rightarrow$ "MONOPOLE"

$$\rightarrow \text{NORMAL } \frac{d\sigma_K}{d\Omega}$$

- SLATER DETS. FOR $\Psi_e^f = \det(\psi'_1, \psi'_2, \dots, \psi'_{k-1}, \psi'_{k+1}, \dots, \psi'_N)$

$$\Psi_e^i = \det(\psi_1, \psi_2, \dots, \psi_{k-1}, \psi_{k+1}, \dots, \psi_N)$$

$$\text{INT.}_K \propto |\langle \Psi_{\text{vib}, v}^f | \Psi_{\text{vib}, v}^i \rangle|^2 |\langle \psi'_1 | \psi_1 \rangle|^2 |\langle \psi'_2 | \psi_2 \rangle|^2 \dots$$

$$|\langle \psi'_{k-1} | \psi_{k-1} \rangle|^2 |\langle \psi'_{k+1} | \psi_{k+1} \rangle|^2 \dots |\langle \psi'_N | \psi_N \rangle|^2$$

$$|\hat{e} \cdot \langle \psi_f | \vec{r} | \psi_K \rangle|^2$$

1e- DIPOLE $\rightarrow d\sigma/d\Omega$

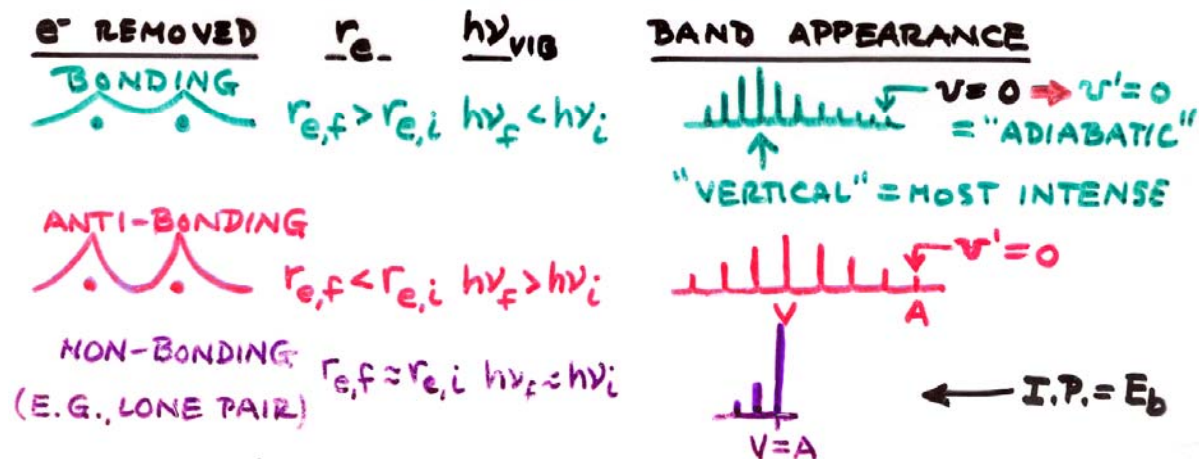
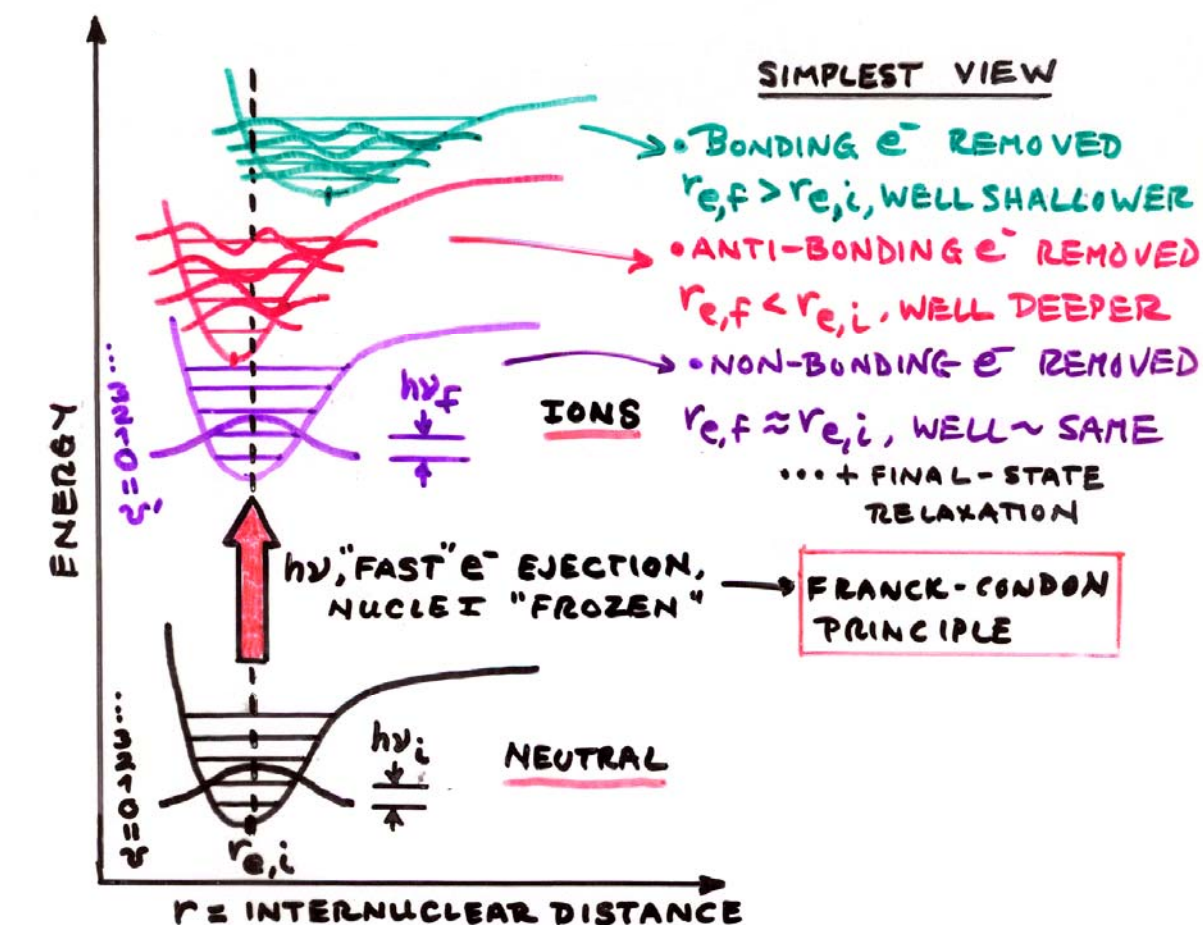
(N-1)e- SHAKE-UP/
SHAKE-OFF $\rightarrow$
"MONOPOLE"

- PLUS DIFFRACTION EFFECTS IN Ψ_f ESCAPE

VIBRATIONAL STRUCTURE IN VALENCE-LEVEL (MO) SPECTRA

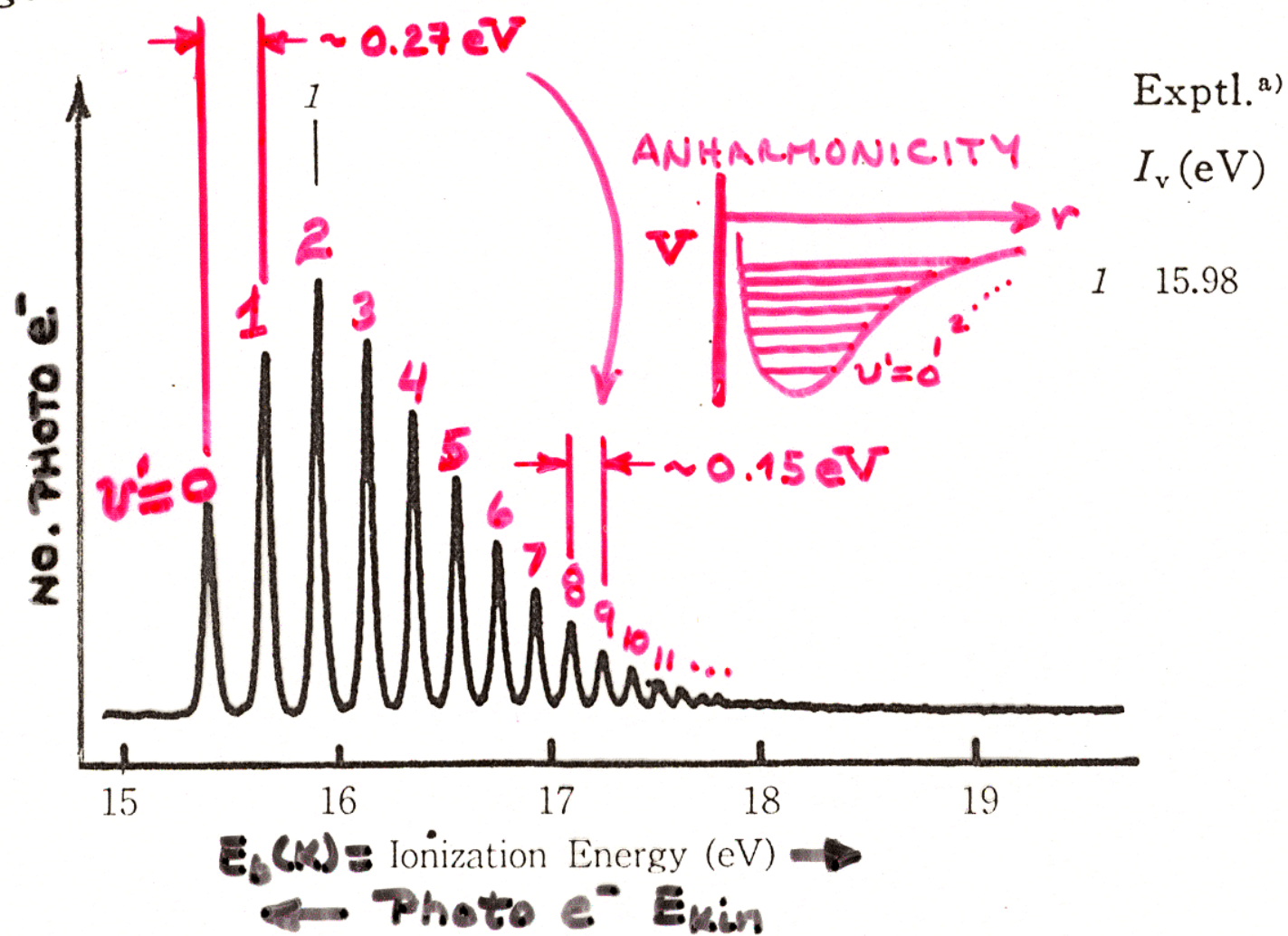
Diatomic A-B example

(Also applies to core-level emission if equilibrium distance changes on forming core hole)



VIBRATIONAL STRUCTURE IN VALENCE-LEVEL (MO) SPECTRA

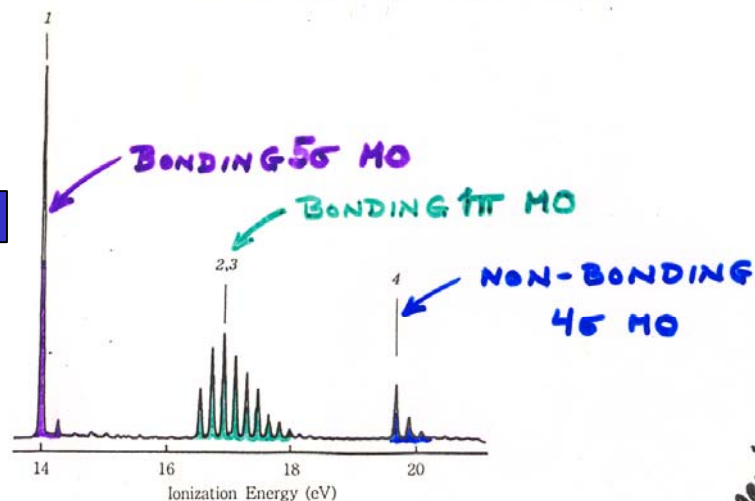
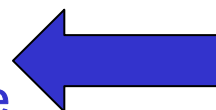
H₂ Hydrogen



(9) CO Carbon Monoxide

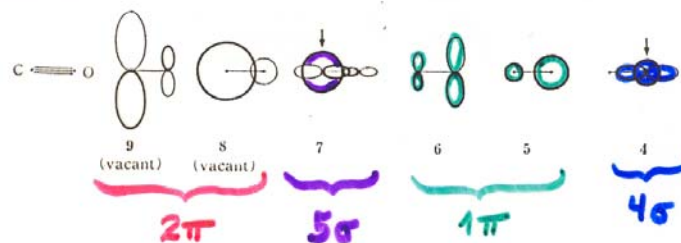
UV PHOTOELECTRON SPECTRUM OF CO

Vibrational fine structure



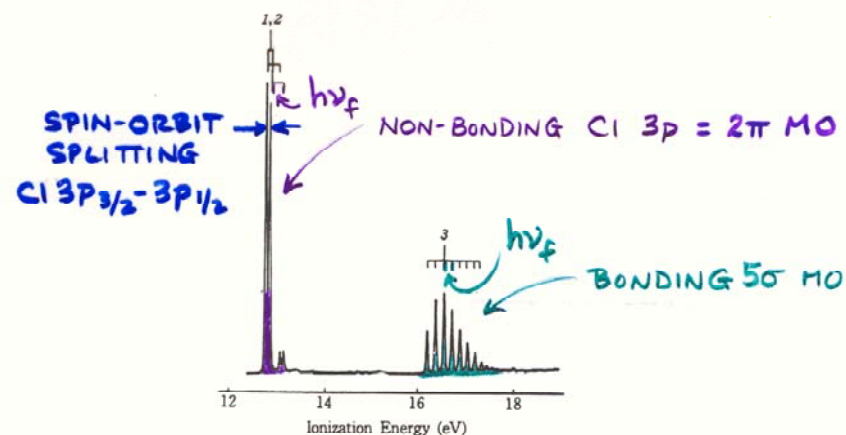
KOOPMANS'				CI FINAL STATE			PRIMARY HOLES
Exptl. ^{a)}	SCF MO [6-31 G] ^{b)}			CI (Ionic State) [6-31 G] ^{c)}			
I_v (eV)	$-\epsilon$ (eV)	MO	Character	E (eV)	State	Configuration	
1 14.01	14.99	5σ (7)	σ _{CO}	13.11	1 ² Σ ⁺	0.93(7 ⁻¹) -0.15(6 ⁻¹ , 7 ⁻¹ , 9 ¹) _a -0.15(5 ⁻¹ , 7 ⁻¹ , 8 ¹) _a	RELAX. + CORREL.
2 16.91	17.48	1π (6, 5)	π _{bond}	16.69	1 ² Π	0.95(6 ⁻¹) ; 0.95(5 ⁻¹)	
3 16.91	17.48						
4 19.72	21.69	4σ (4)	n _O	19.29	2 ² Σ ⁺	0.92(4 ⁻¹) +0.16(6 ⁻¹ , 7 ⁻¹ , 9 ¹) _a +0.16(5 ⁻¹ , 7 ⁻¹ , 8 ¹) _a	

- a) The spectrum: this work. The I_v 's: Turner *et al.* (215). See also other works: Turner and May (215 a); Carlson and Jonas (54); Gardner and Samson (104); Edqvist *et al.* (90); Potts and Williams (182 a); and Natalis *et al.* (165).
- b) We used the bond length reported (A 3); symmetry $C_{\infty h}$. $E_{SCF} = -112.6672$ hartree. In 4-31 G calculations, $E_{SCF} = -112.5524$ hartree and $-\epsilon$ (eV) = 14.93, 17.41, 17.41, and 21.60.
- c) CI-II. (9, 8) = 1π. |N> = 0.98 (SCF). The results obtained in other CI levels are given in Appendix B.



Kimura et al.,
 "Handbook of Hel
 Photoelectron Spectra"

THE UV PHOTOELECTRON SPECTRUM OF HCl



		KOPPHANS' THEO.		CONFIG. INT. ON FINAL STATE		
Exptl. ^{a)}		SCF MO [4-31 G] ^{b)}		~Cl (Ionic State) [4-31 G] ^{c)}		
I_v (eV)	$-\epsilon$ (eV)	MO	Character	E (eV)	State	Configuration
1	12.75	12.77	2π (9, 8) n_{Cl}	11.97	1 ² Π	0.98 (9 ⁻¹); 0.98 (9 ⁻¹)
2	12.85	12.77				
3	16.28	16.50	5σ (7) σ_{HCl}	16.10	1 ² Σ ⁺	0.98 (7 ⁻¹)

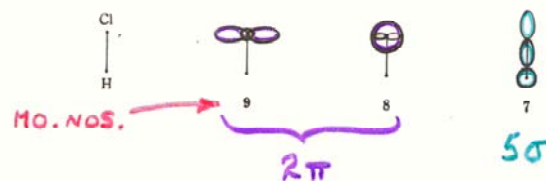
a) The spectrum: this work. The I_v 's: Frost *et al.* (102). See also other works: Lempka *et al.* (150); Turner *et al.* (215); and Weiss *et al.* (224).

b) We used the bond length reported in Ref. (A 5); symmetry $C_{\infty h}$. $E_{SCF} = -459.5631$ hartree.

c) CI-V: $|N\rangle = 0.99$ (SCF).

CI-V': E (eV) = 12.01 and 16.11.

CI-III: E (eV) = 12.60 and 16.79.

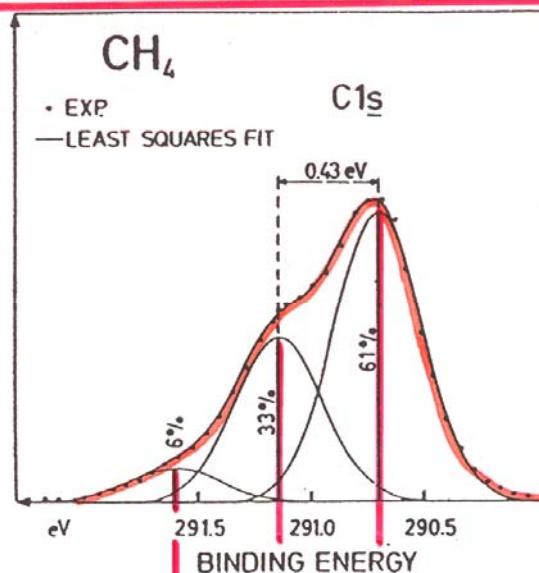


(FROM KIMURA ET AL., "HANDBOOK OF He I PHOTO-ELECTRON SPECTRA OF FUND. ORGANIC MOLECULES")

VIBRATIONAL FINE STRUCTURE IN CORE SPECTRA

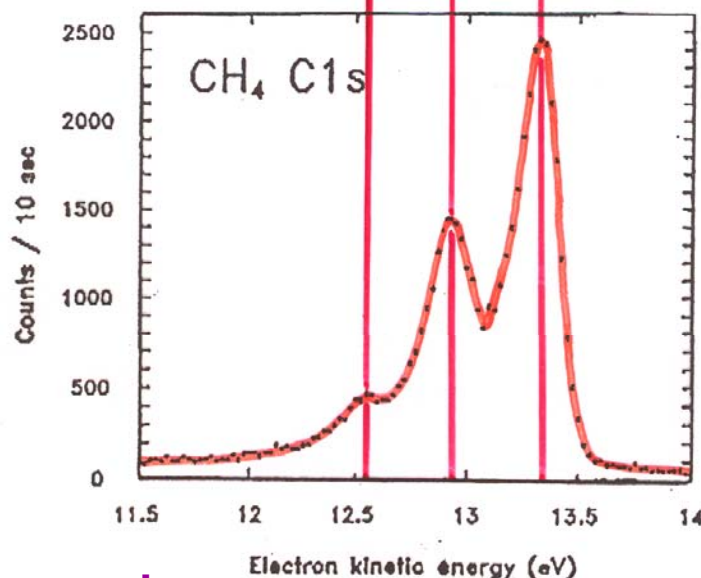
MONOCHROMATIZED
LABORATORY
X-RAY SOURCE

"Basic Concepts of XPS"
Figure 40



1970
SIEGBAHN,
GELWIS,
ET AL.

NEW SR
BEAMLINE
AT
BROOK-
HAVEN
(~5-10X
FASTER
@ALS)

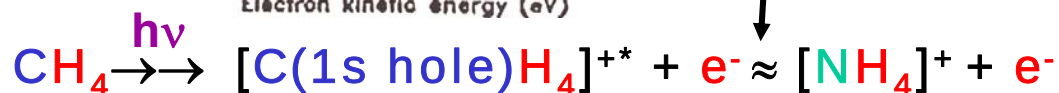


1991
BRADSHAW,
KAINDL,
ET AL.

CH distance = 1.10 Å

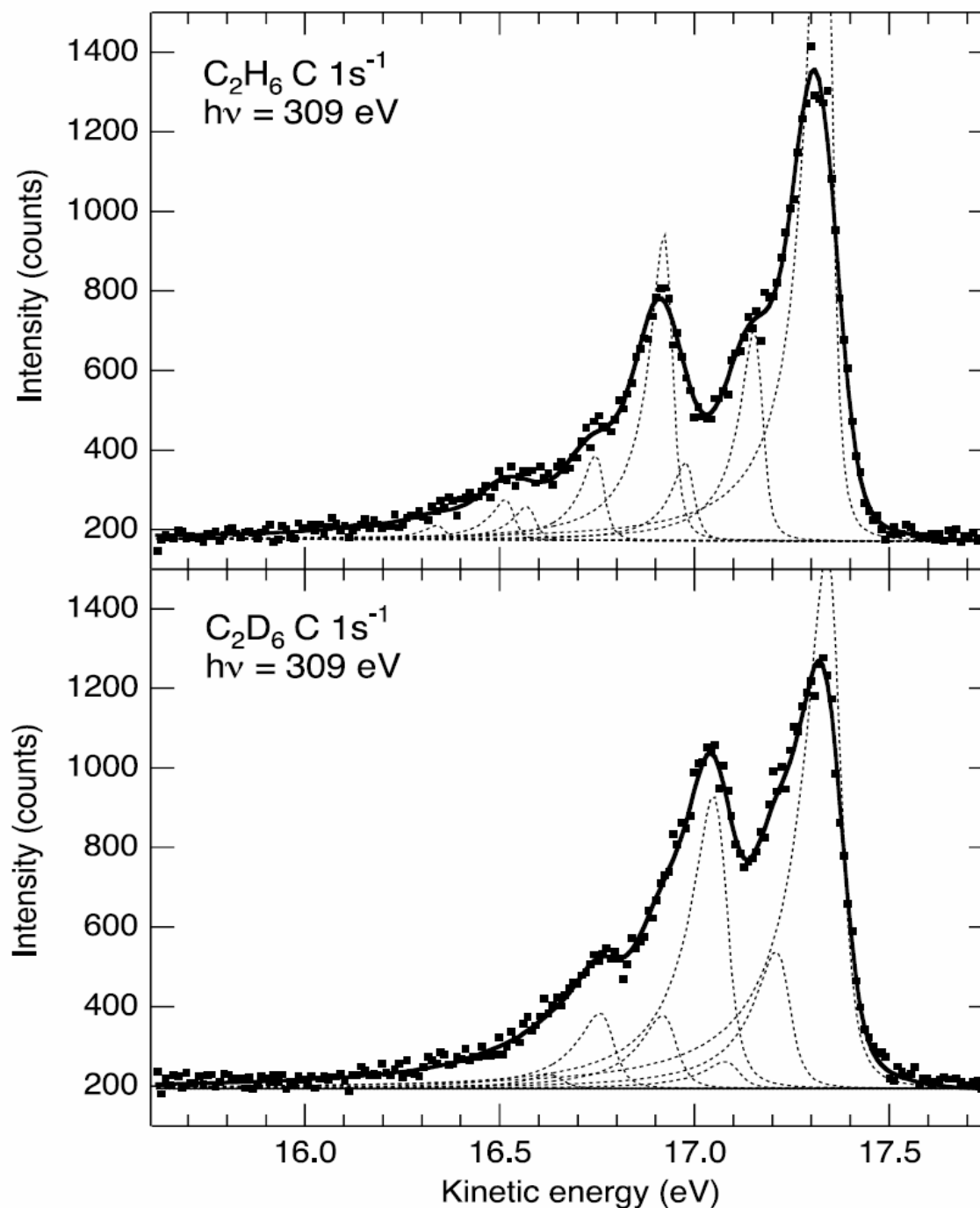
NH distance = 1.00 Å

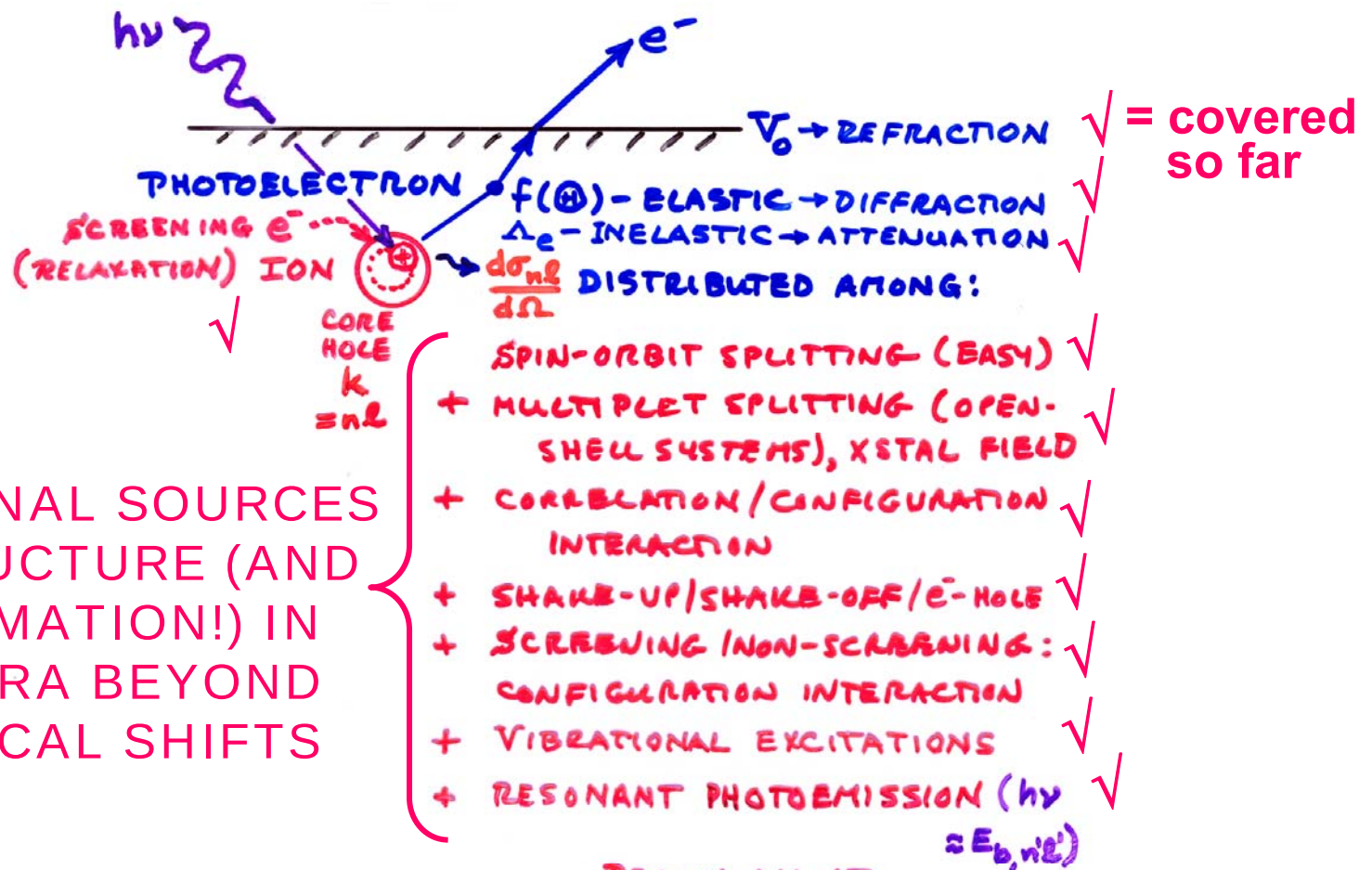
Equivalent-core



Vibrational fine structure in C 1s photoemission from ethane:
two progressions ν_a at 0.407 eV and ν_b at 0.176 eV and various excitations (ν_a, ν_b)

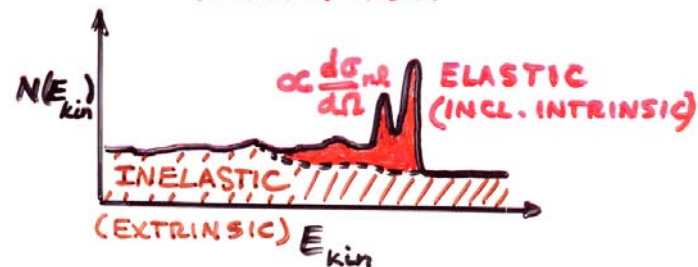
Rennie et al.,
J. Phys. At. Mol. Opt.
Phys. 32, 2691 (1999)





ADDITIONAL SOURCES OF STRUCTURE (AND INFORMATION!) IN SPECTRA BEYOND CHEMICAL SHIFTS

REALLY ALL AT ONCE, BUT SUM RULES + THEORY HELP



Outline

Surface, interface, and nanoscience—short introduction

Some surface concepts and techniques→photoemission

Synchrotron radiation: experimental aspects

Electronic structure—a brief review

**The basic synchrotron radiation techniques:
more experimental and theoretical details**

Core-level photoemission

Valence-level photoemission

 **Microscopy with photoemission:
All of the above with lateral spatial resolution
of ca. 20 nm, going down to few nm in future
(Next lecturers)**

Thank you for your attention!