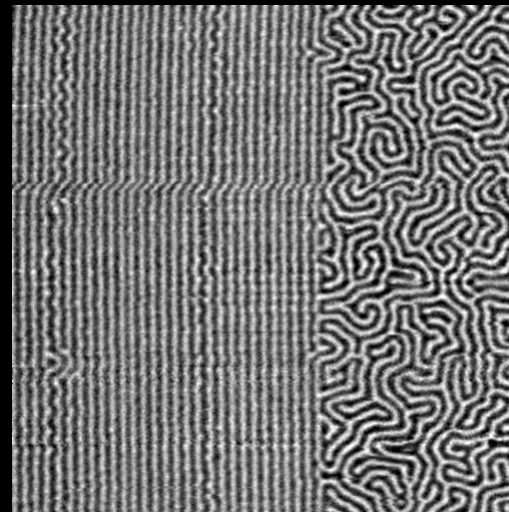


# MAGNETISM AS SEEN WITH X-RAYS



ELKE ARENHOLZ  
LAWRENCE BERKELEY NATIONAL LABORATORY  
AND  
DEPARTMENT OF MATERIAL SCIENCE AND ENGINEERING, UC BERKELEY

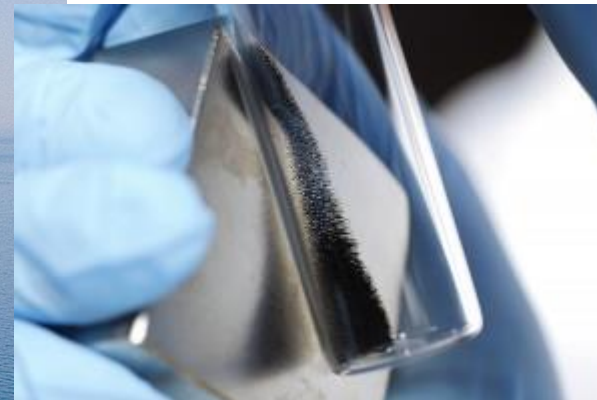
# MAGNETIC MATERIALS TODAY



**Magnetic thin films  
for  
information storage  
and processing**



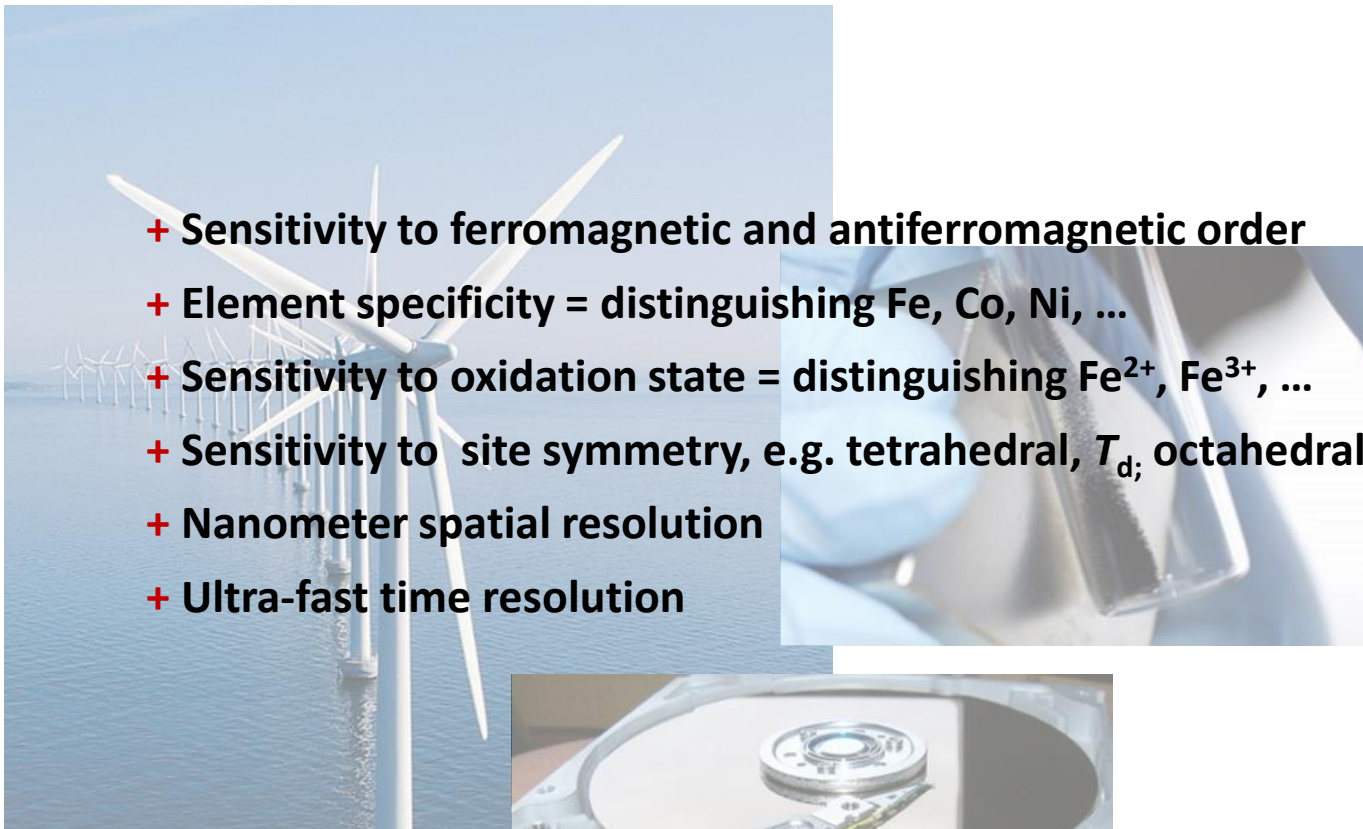
**Magnetic materials  
for  
energy applications**



**Magnetic  
nanoparticles  
for  
biomedical and  
environmental  
applications**

# MAGNETIC MATERIALS CHARACTERIZATION WISH LIST

- + Sensitivity to ferromagnetic and antiferromagnetic order
- + Element specificity = distinguishing Fe, Co, Ni, ...
- + Sensitivity to oxidation state = distinguishing  $\text{Fe}^{2+}$ ,  $\text{Fe}^{3+}$ , ...
- + Sensitivity to site symmetry, e.g. tetrahedral,  $T_d$ ; octahedral,  $O_h$
- + Nanometer spatial resolution
- + Ultra-fast time resolution



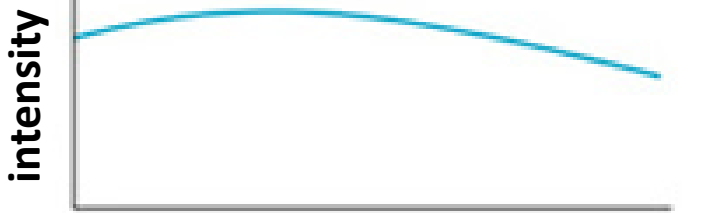
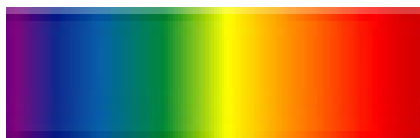
## SOFT X-RAY SPECTROSCOPY AND MICROSCOPY

# SPECTROSCOPY

Light/Photon Source

Monochromator

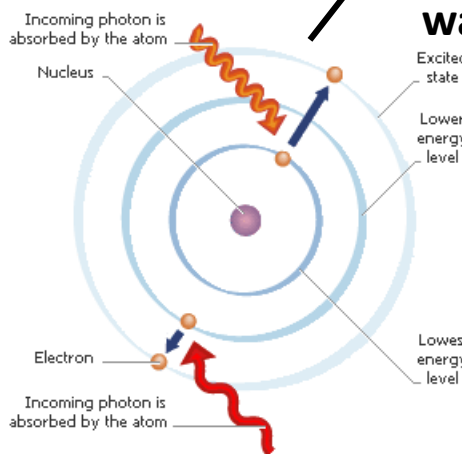
Sample



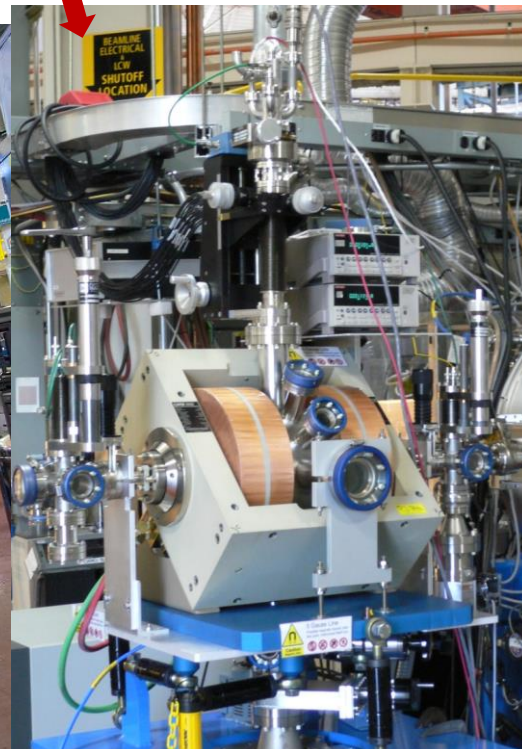
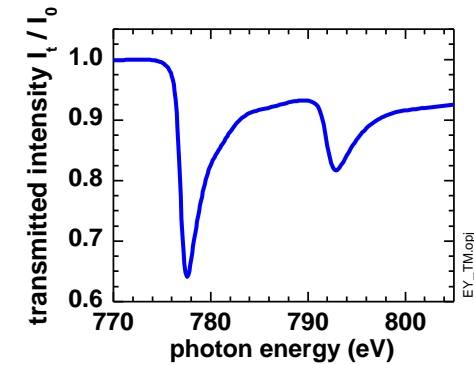
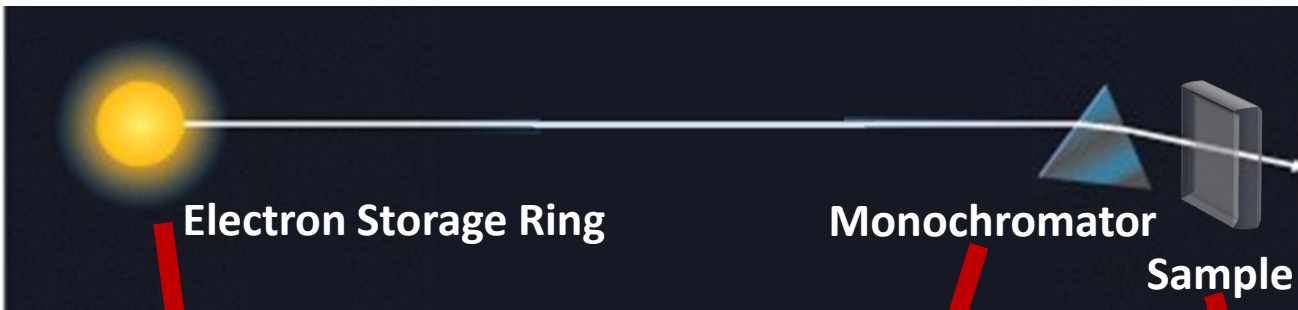
wavelength, photon energy

intensity

wavelength, photon energy

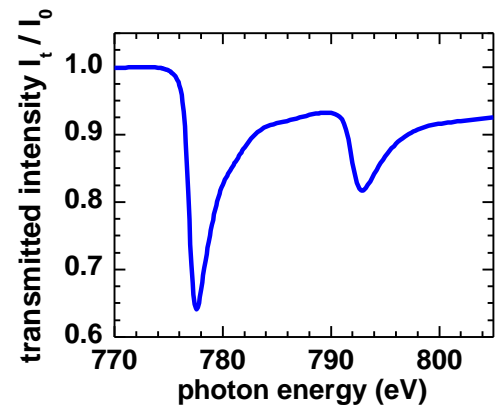
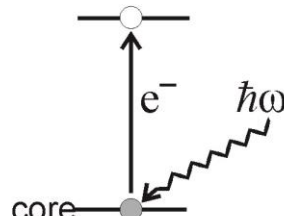
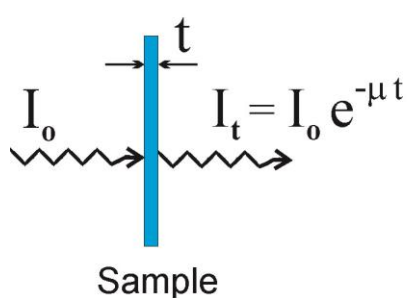


# SOFT X-RAY SPECTROSCOPY ( $h\nu \approx 500\text{-}1000\text{eV}$ , $\lambda \approx 1\text{-}2\text{nm}$ )



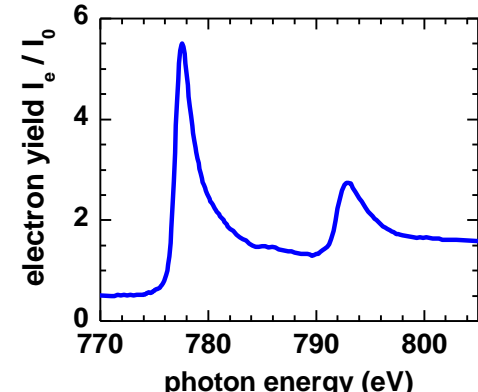
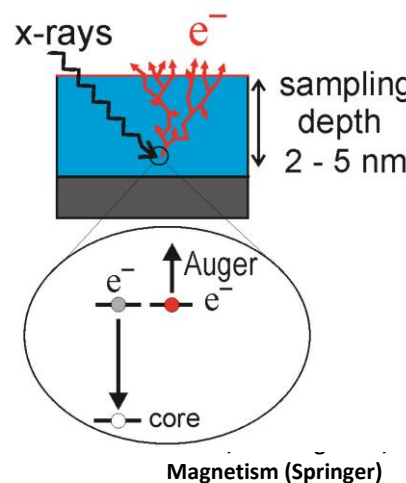
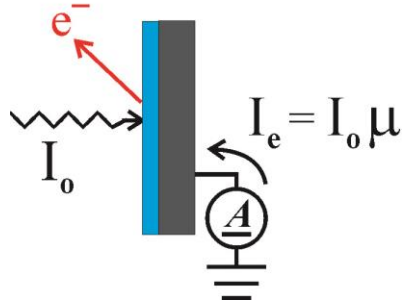
# X-RAY ABSORPTION – DETECTION MODES

## Transmission



photons absorbed

## Electron Yield

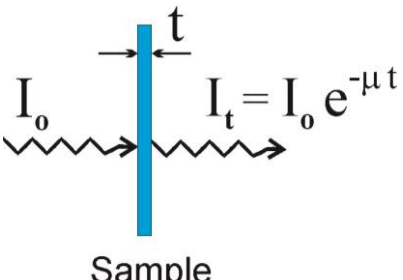


electrons generated

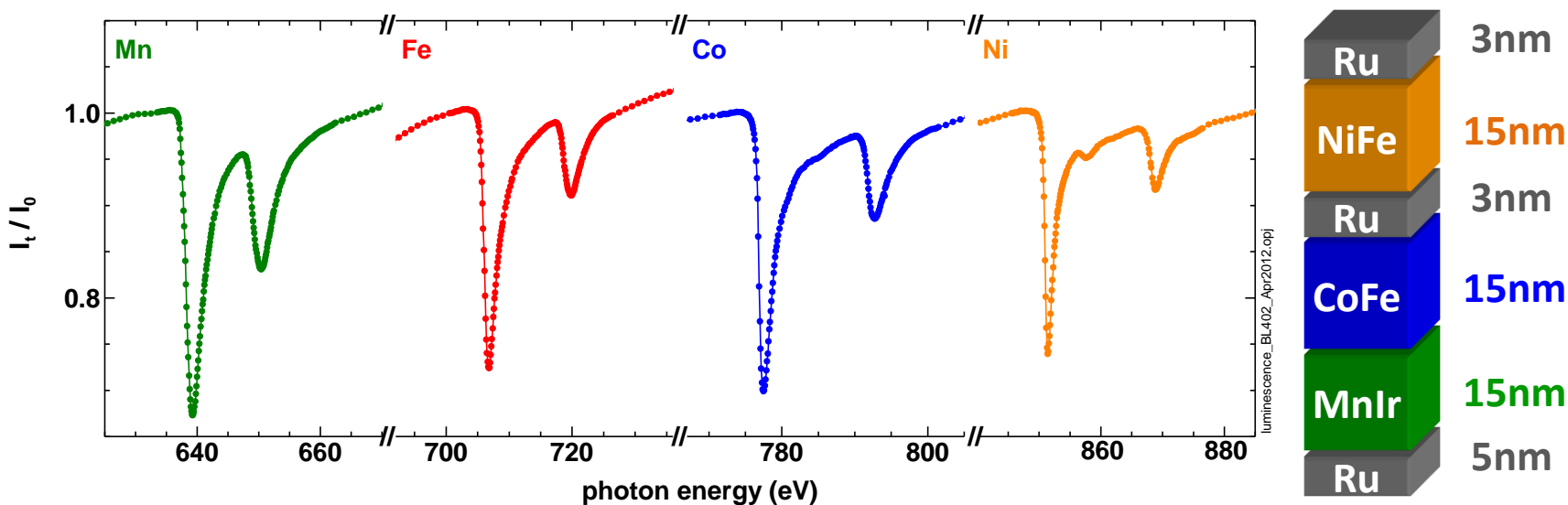
## Electron yield:

- + Absorbed photons create core holes subsequently filled by Auger electron emission
- + Auger electrons create low-energy secondary electron cascade through inelastic scattering
- + Emitted electrons  $\propto$  probability of Auger electron creation  $\propto$  absorption probability

# SOFT X-RAY ABSORPTION – PROBING DEPTH

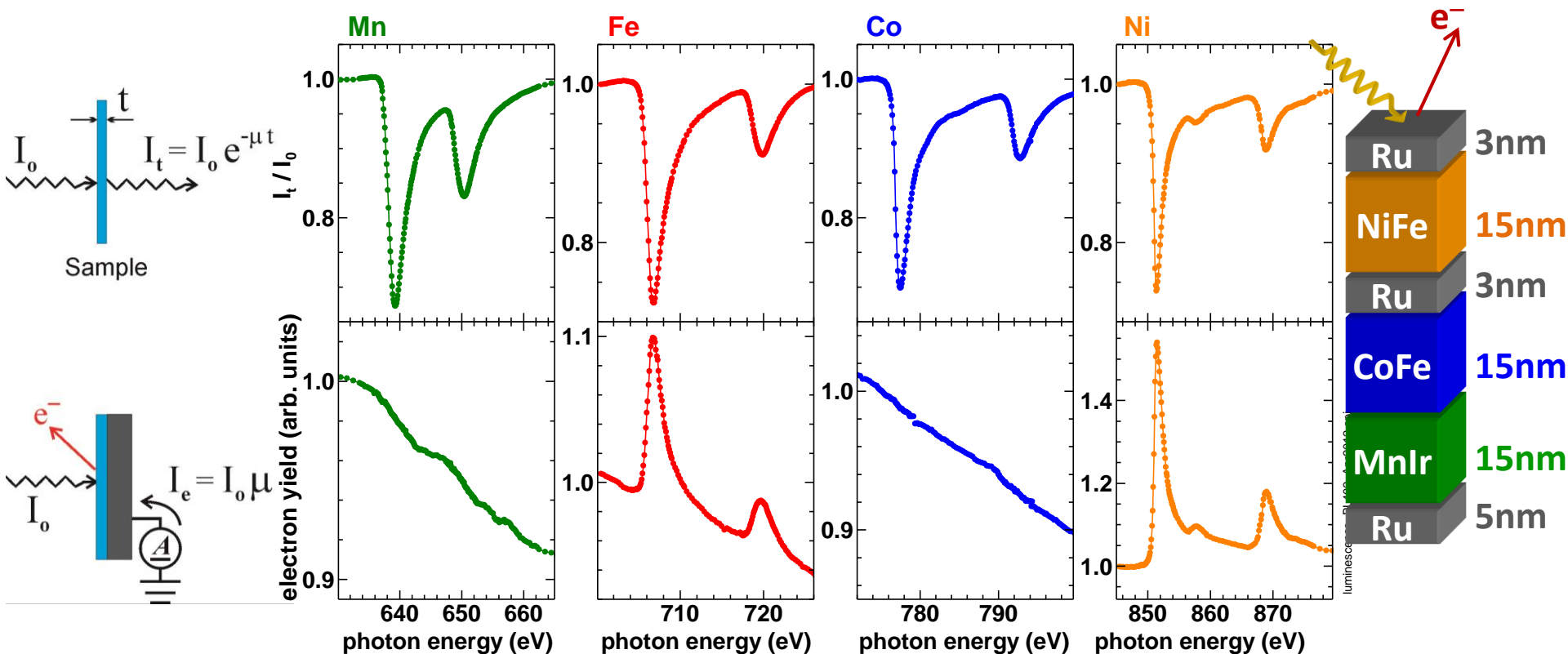


Element	10eV below $L_3$ 1/ $\mu$ [nm]	at $L_3$ 1/ $\mu$ [nm]	40 eV above $L_3$ 1/ $\mu$ [nm]
Fe	550	17	85
Co	550	17	85
Ni	625	24	85



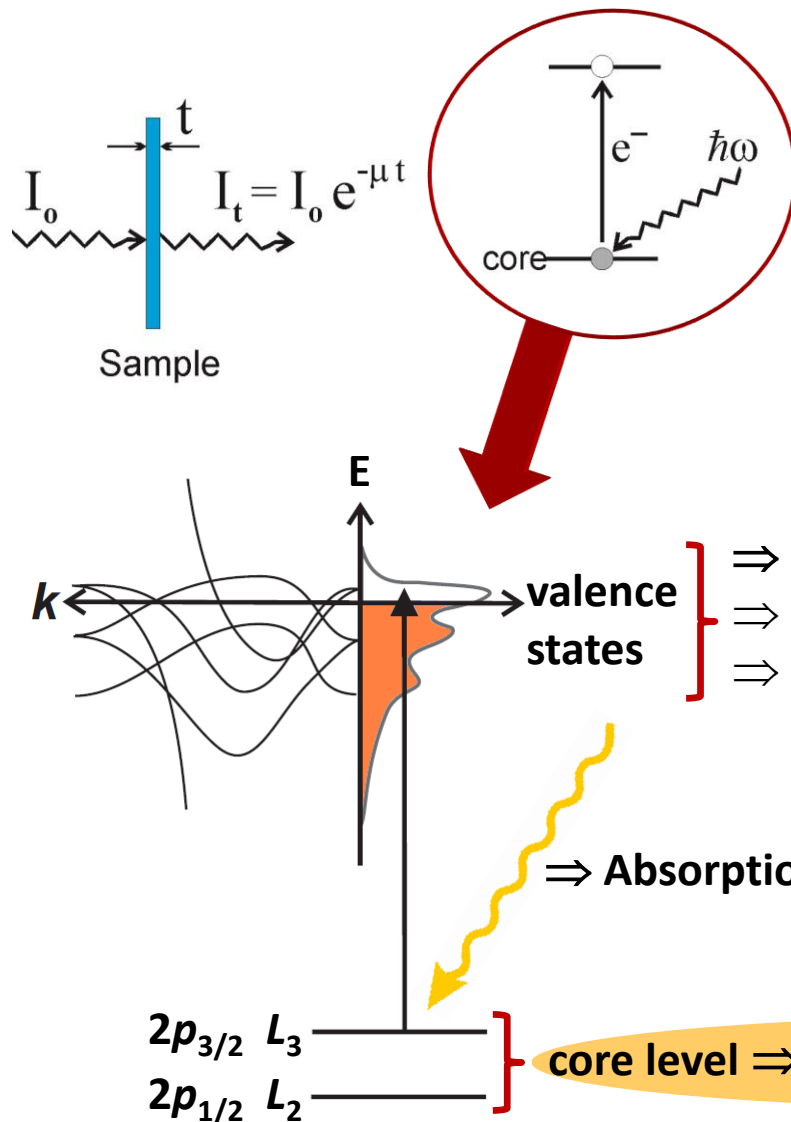
~10-20 nm layer thick films supported by substrates transparent to soft x-rays

# X-RAY ABSORPTION – DETECTION MODES AND PROBING DEPTH



- + Electron sample depth: 2-5 nm in Fe, Co, Ni  
→ 60% of the electron yield originates from the topmost 2-5 nm

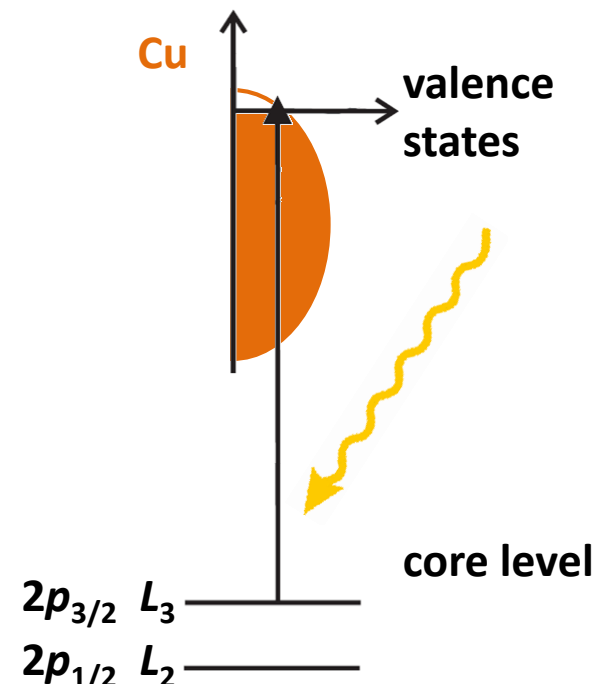
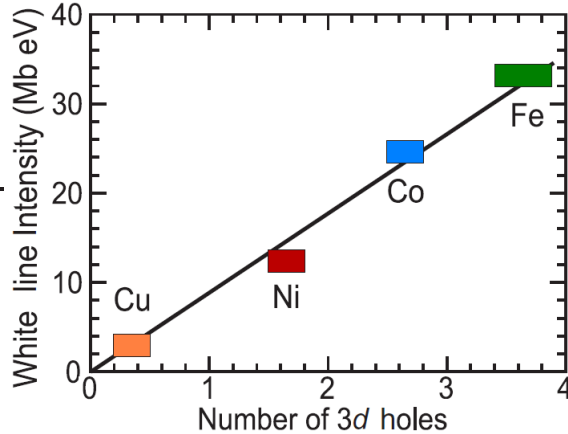
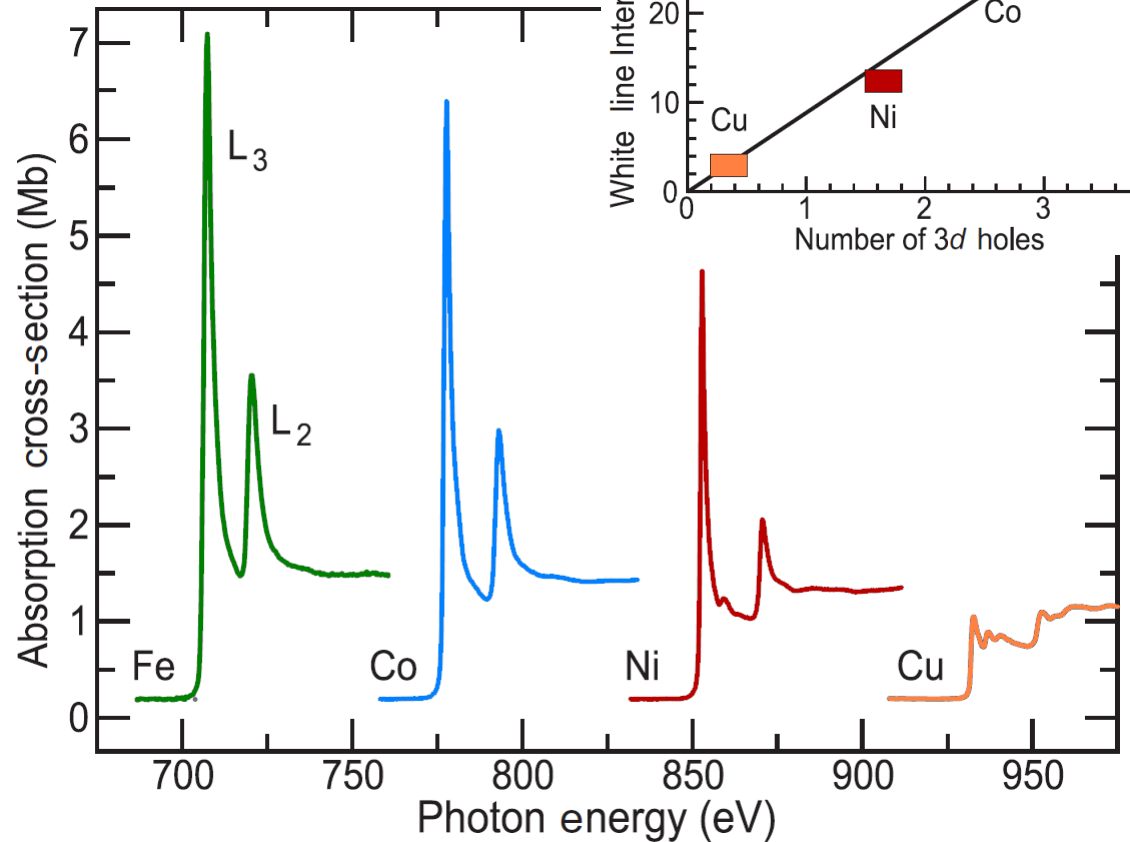
# X-RAY ABSORPTION – FUNDAMENTALS



## Experimental Concept:

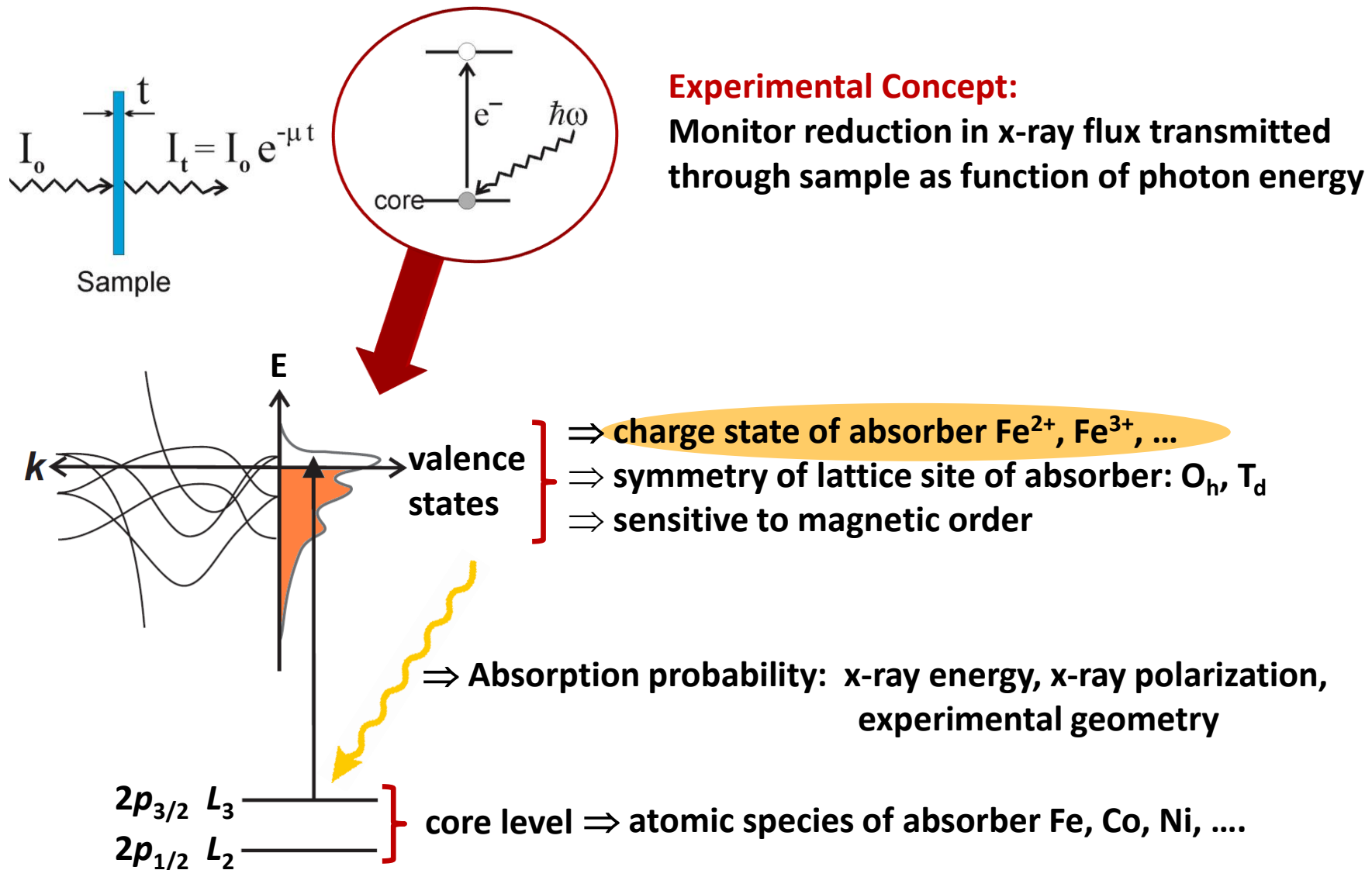
Monitor reduction in x-ray flux transmitted through sample as function of photon energy

# 'WHITE LINE' INTENSITY



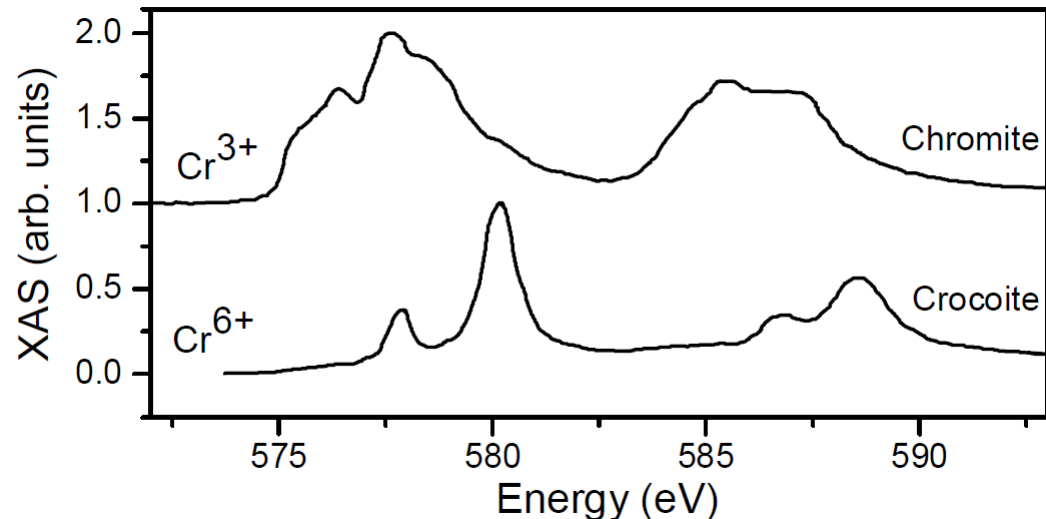
Intensity of  $L_{3,2}$  resonances is proportional to number of  $d$  states above the Fermi level, i.e. number of holes in the  $d$  band.

# X-RAY ABSORPTION – FUNDAMENTALS

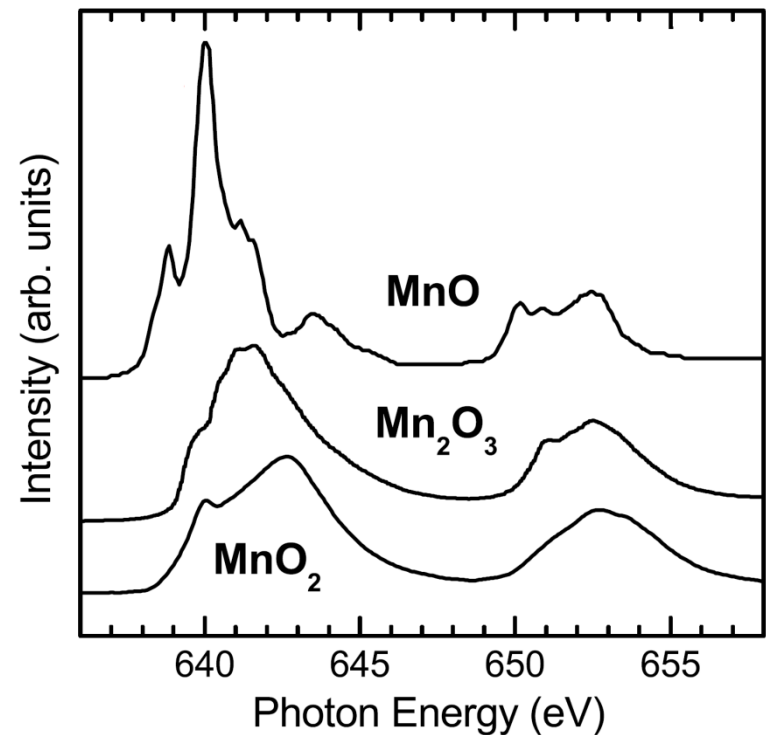


# X-RAY ABSORPTION – VALENCE STATE

Influence of the charge state of the absorber



N. Telling *et al.*,  
Appl. Phys. Lett. **95**, 163701 (2009)



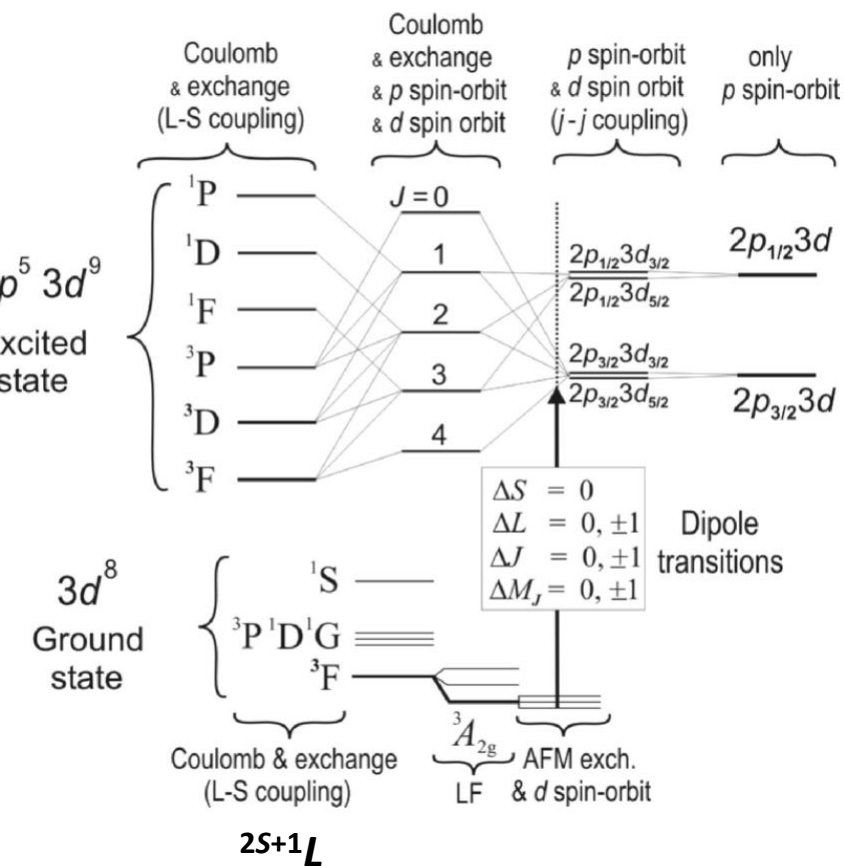
J.-S. Kang *et al.*, Phys. Rev. B **77**, 035121 (2008)

# X-RAY ABSORPTION – CONFIGURATION MODEL

$\text{Ni}^{2+}$  in  $\text{NiO}$ :  $2p^6 3d^8 \rightarrow 2p^5 3d^9$

Configuration model, e.g. *L* edge absorption :

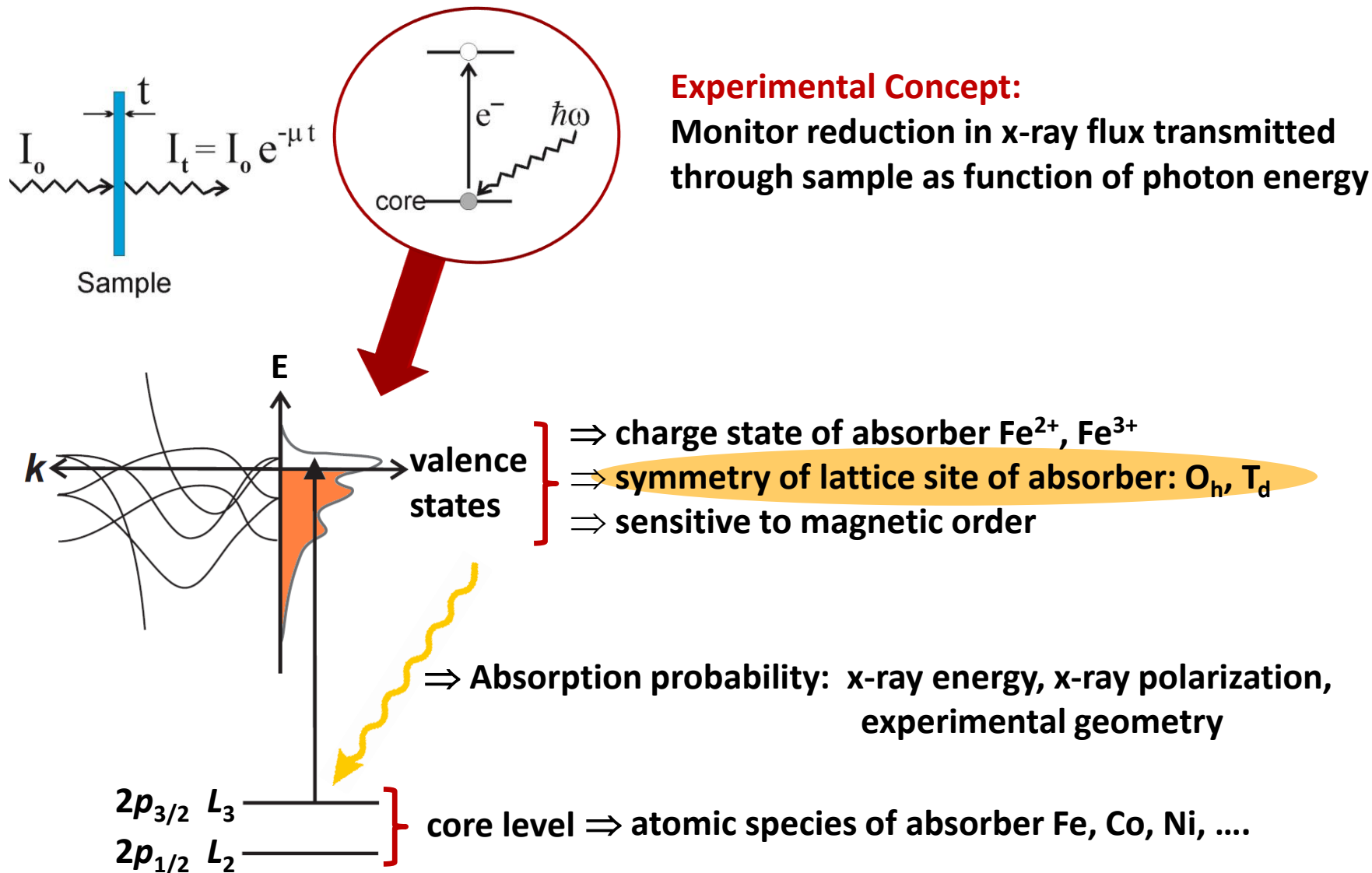
- + Excited from ground/initial state configuration,  $2p^6 3d^8$  to excited/final state configuration,  $2p^5 3d^9$
- + Omission of all full subshells (spherical symmetric)
- + Takes into account correlation effects in the ground state as well as in the excited state
- + Leads to multiplet effects/structure



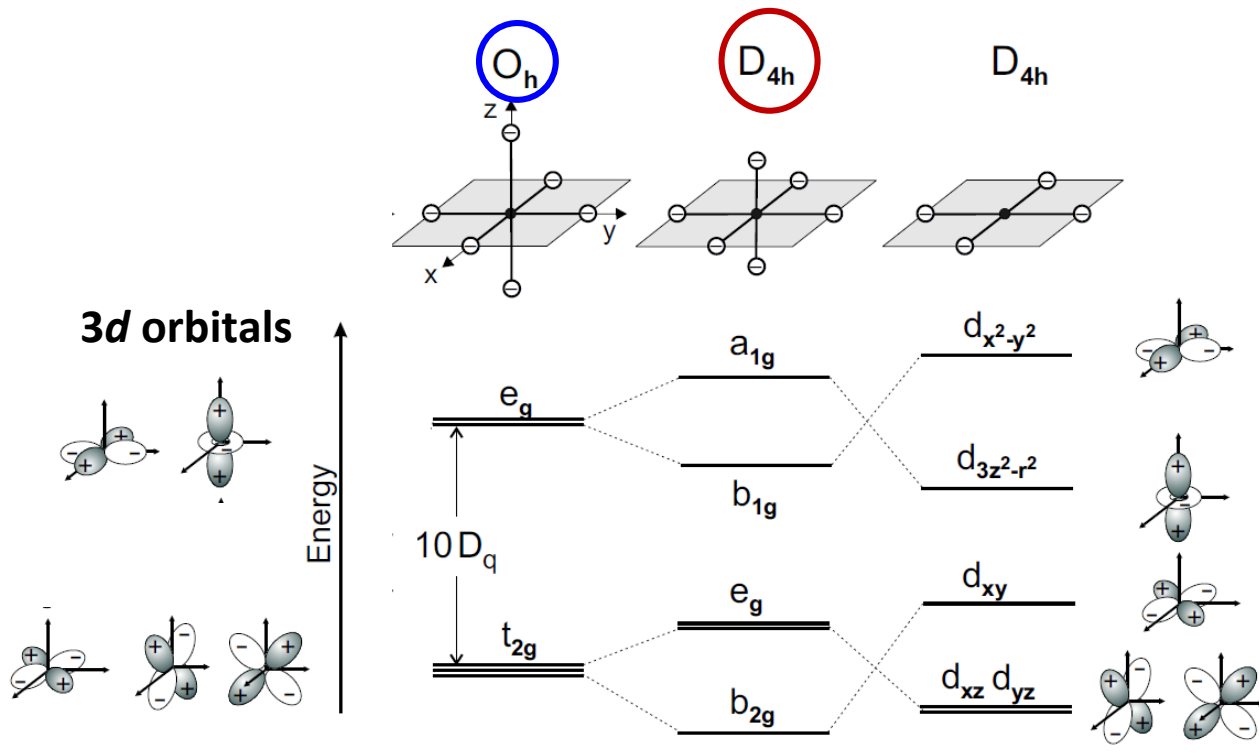
<http://www.anorg.chem.uu.nl/CTM4XAS/>

J. Stöhr, H.C. Siegmann,  
Magnetism (Springer)

# X-RAY ABSORPTION – FUNDAMENTALS



# SENSITIVITY TO SITE SYMMETRY: $\text{Ti}^{4+} L_{3,2}$



+ Electric dipole transitions:  $d^0 \rightarrow 2p^5 3d^1$

+ Crystal field splitting  $10Dq$  acting on 3d orbitals:

**Octahedral symmetry:**

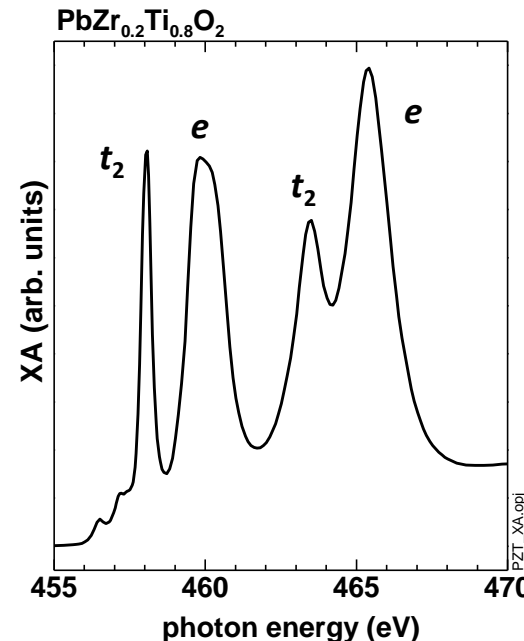
**$e$  orbitals towards ligands**  $\rightarrow$  higher energy

**$t_2$  orbitals between ligands**  $\rightarrow$  lower energy

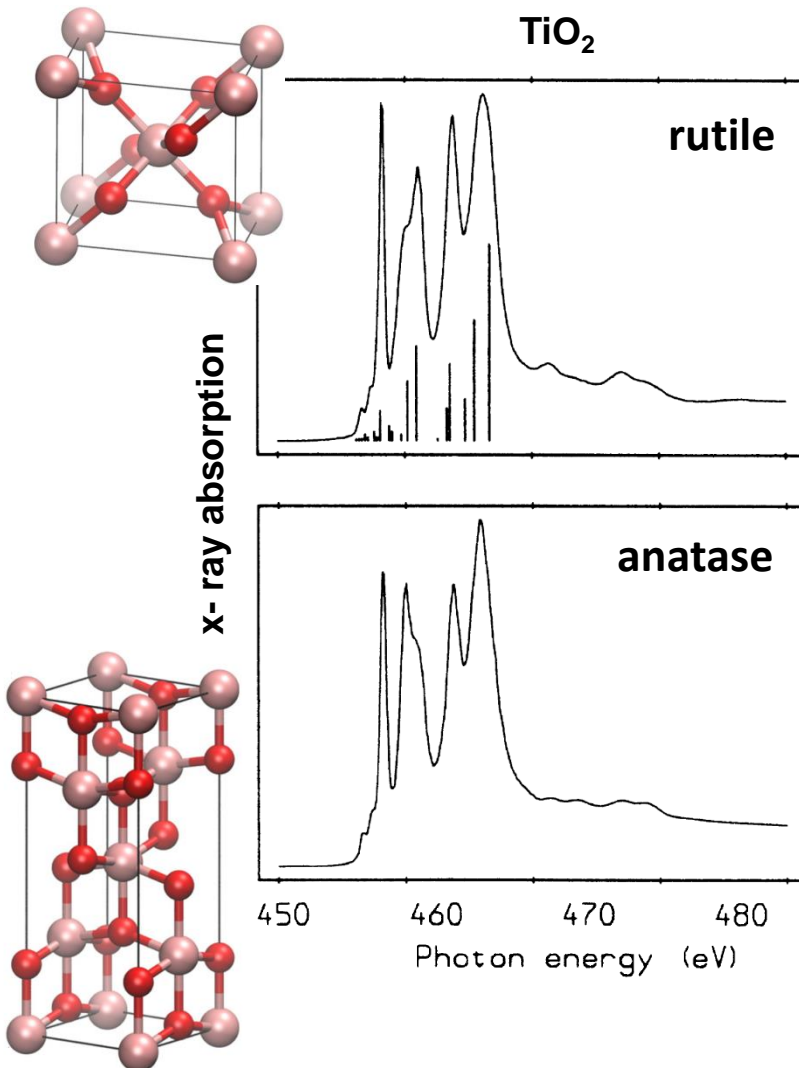
**Tetragonal symmetry:**

**$e$  orbitals**  $\rightarrow b_2 = d_{xy}, e = d_{yz}, d_{yz}$

**$t_2$  orbitals**  $\rightarrow b_1 = d_{x^2-y^2}, a_1 = d_{3z^2-r^2}$



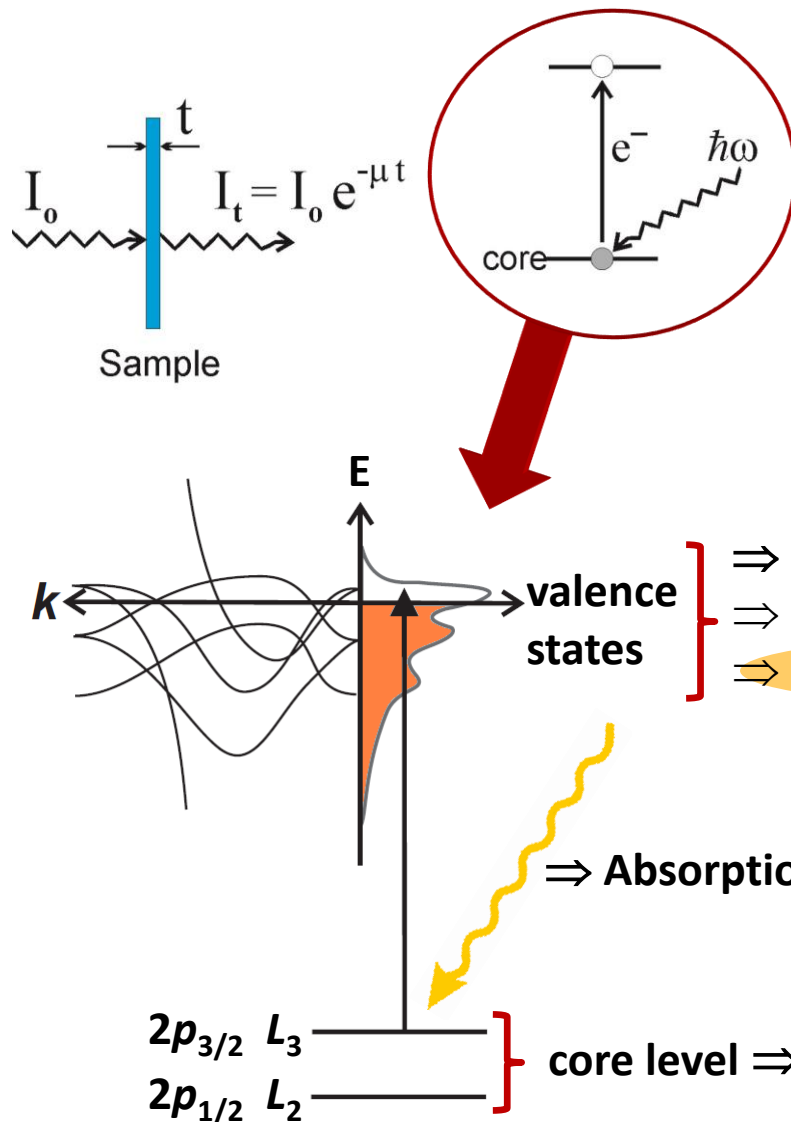
# X-RAY ABSORPTION – LATTICE SYMMETRY



Influence of lattice site symmetry at the absorber

G. Van der Laan  
Phys. Rev. B 41, 12366 (1990)

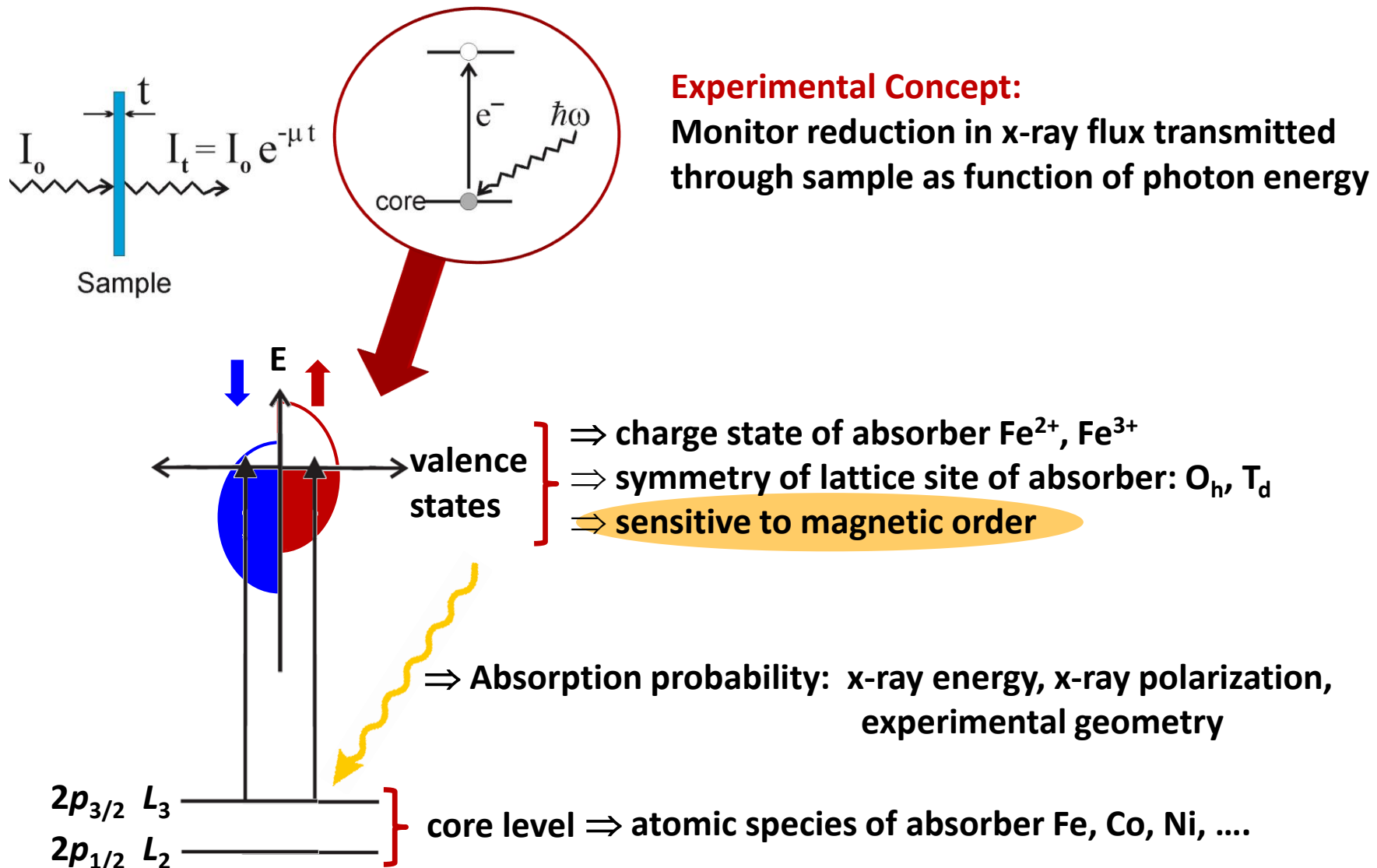
# X-RAY ABSORPTION – FUNDAMENTALS



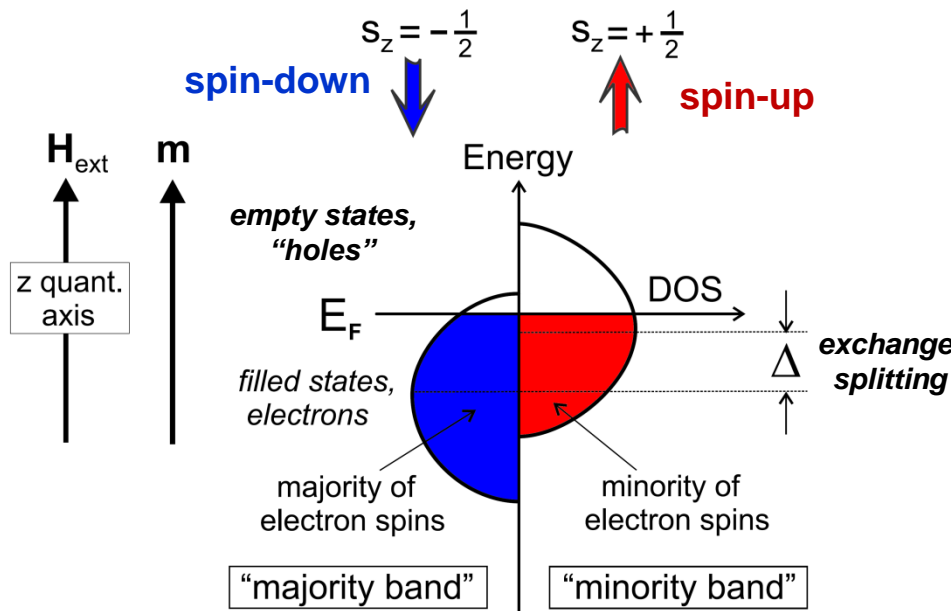
## Experimental Concept:

Monitor reduction in x-ray flux transmitted through sample as function of photon energy

# X-RAY ABSORPTION – FUNDAMENTALS



# STONER MODEL FOR FERROMAGNETIC METALS



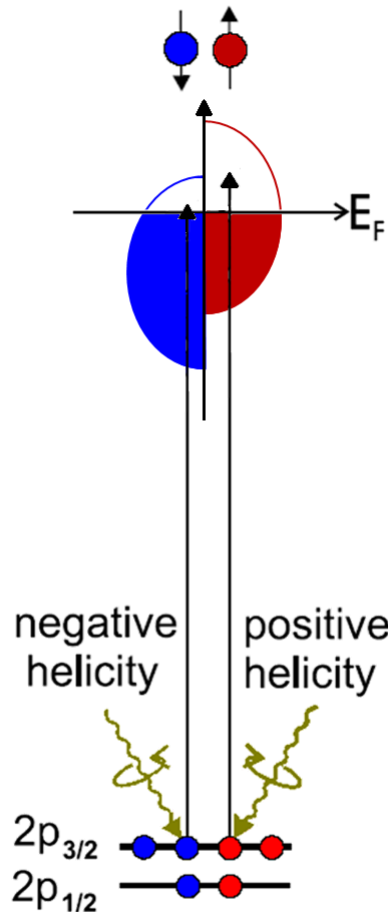
J. Stöhr, H.C. Siegmann,  
Magnetism (Springer)

**3d shell**

- + Magnetic moments in Fe, Co, Ni well described by Stoner model:  $d$ -bands containing up and down spins shifted relative to each other by exchange splitting
- + Spin-up and spin-down bands filled according to Fermi statistics
- + Magnetic moment  $|m|$  determined by difference in number of electrons in majority and minority bands

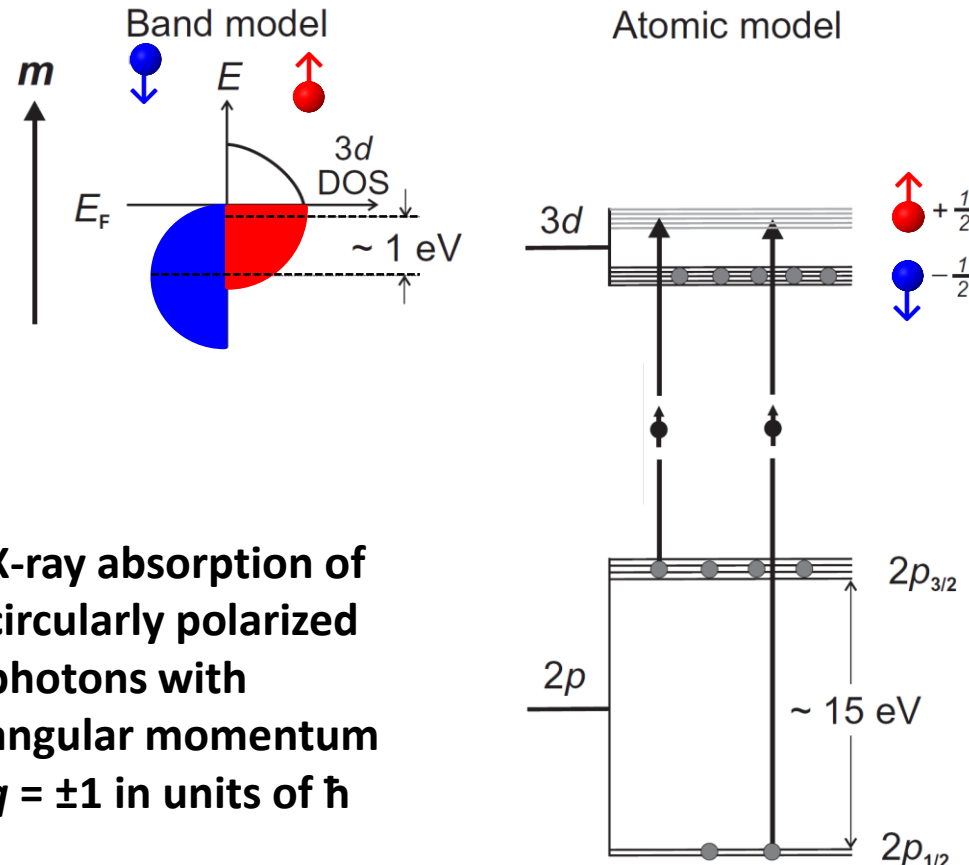
$$|m| \propto \mu_B (n_e^{\text{maj}} - n_e^{\text{min}})$$

# STONER MODEL FOR FERROMAGNETIC METALS



- + Calculate transition probabilities from filled  $2p_{3/2}$  and  $2p_{1/2}$  states to empty states in  $d$ -band for circularly polarized x rays using **Fermi's golden rule**
- + Right/left circularly polarized photons with angular momentum  $q = +/-1$  in units of  $\hbar$
- + Important: Spin of the excited electron remains the same

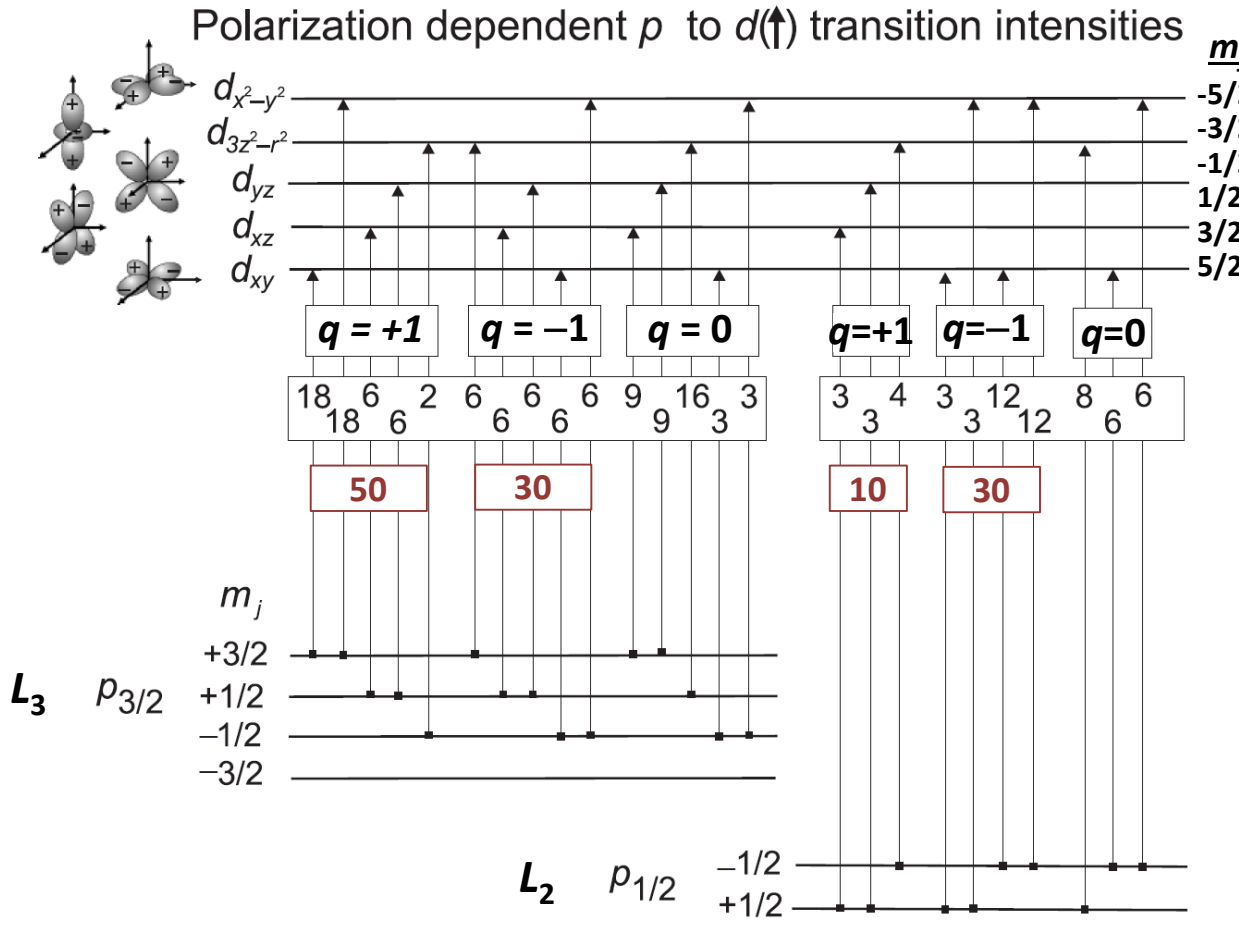
# ORIGIN OF X-RAY MAGNETIC CIRCULAR DICHROISM



- + Consider strong ferromagnet with one filled spin band:
  - All spin down  $d$  states filled
  - Spin up  $d$  states partially filled
- + This specific case:  
Only spin up electron excited

J. Stöhr, H.C. Siegmann,  
Magnetism (Springer)

# ORIGIN OF X-RAY MAGNETIC CIRCULAR DICHROISM

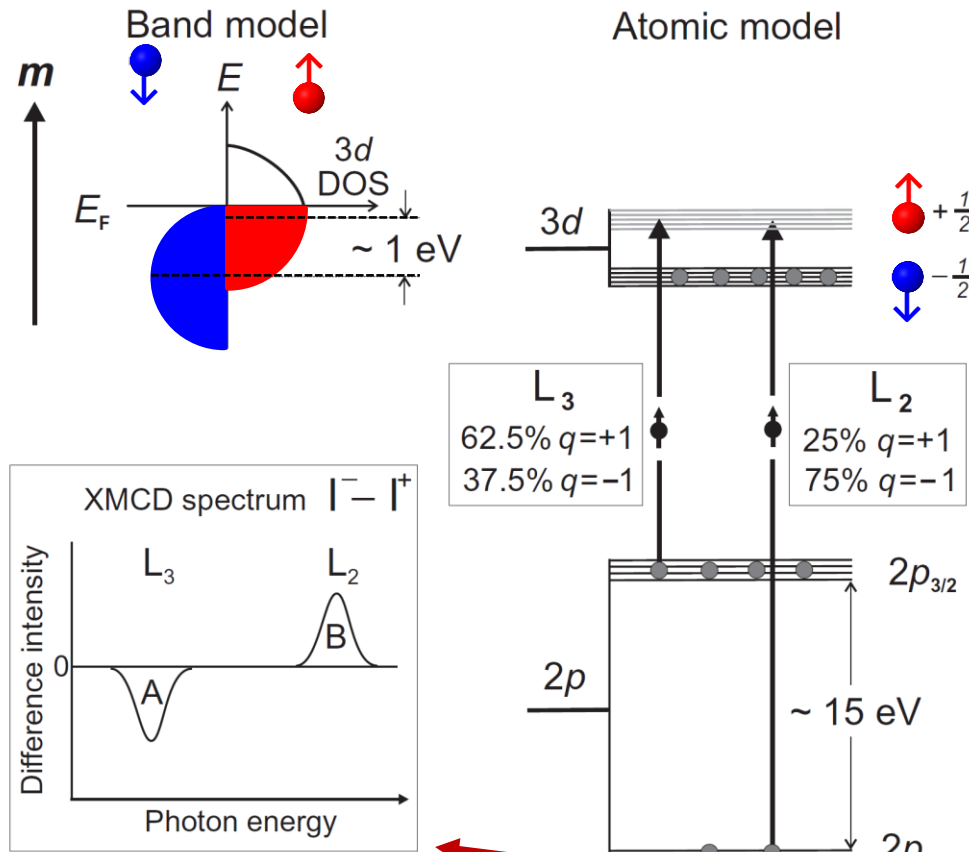


J. Stöhr, H.C. Siegmann,  
Magnetism (Springer)

$L_3$ : X rays with  $q = +/−1$  excite 62.5%/37.5% of the spin up electrons

$L_2$ : X rays with  $q = +/−1$  excite 25%/75% of the spin up electrons

# ORIGIN OF X-RAY MAGNETIC CIRCULAR DICHROISM



$L_3$ : X rays with  $q = +1$  excite 62.5% spin up electrons

X rays with  $q = -1$  excite 37.5% spin up electrons

$$\Delta = 25\%$$

$L_2$ : X rays with  $q = +1$  excite 25% spin up electrons

X rays with  $q = -1$  excite 75% spin up electrons

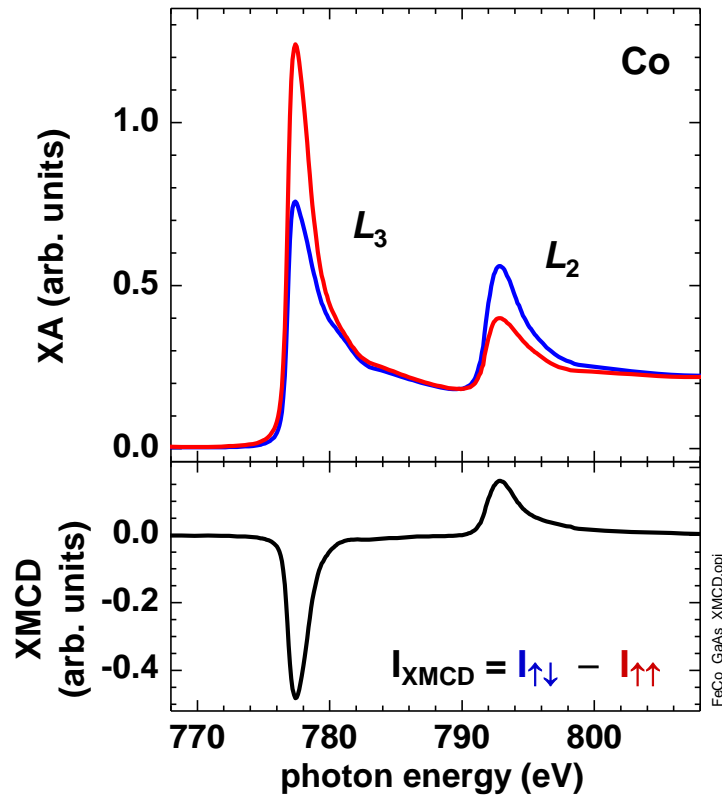
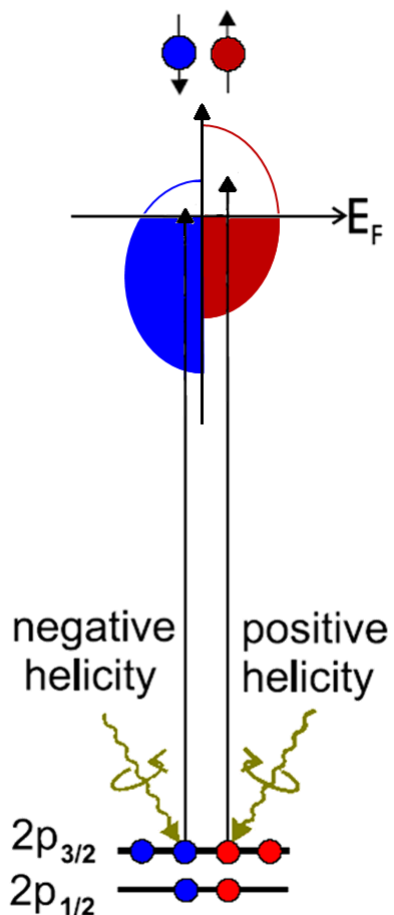
$$\Delta = 50\%$$

Taking into account 2x higher population of  $2p_{3/2}$  state as compared to  $2p_{1/2}$  state:

⇒ Identical magnitude XMCD at  $L_3$  and  $L_2$  with opposite sign

Contributions of  $p \rightarrow d$  transitions dominate  $p \rightarrow s$  transitions by factor >20.

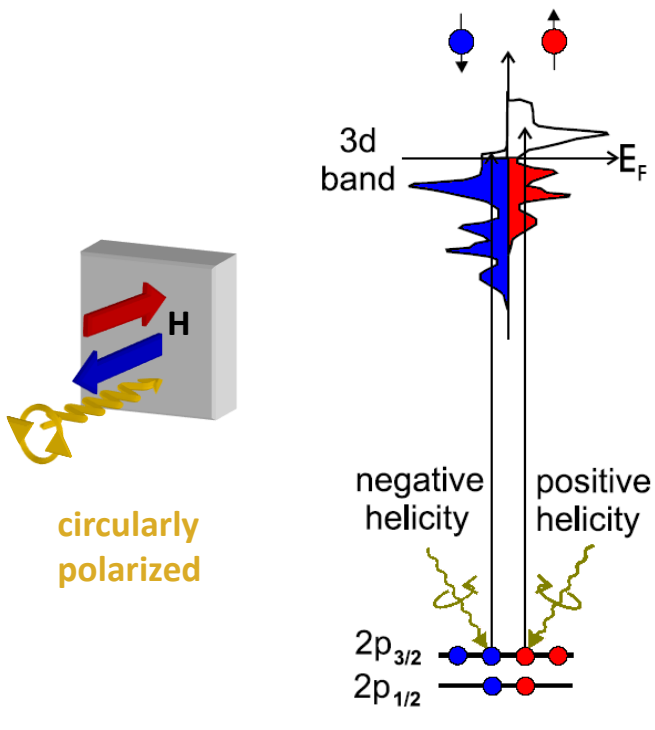
# X-RAY MAGNETIC CIRCULAR DICHROISM (XMCD)



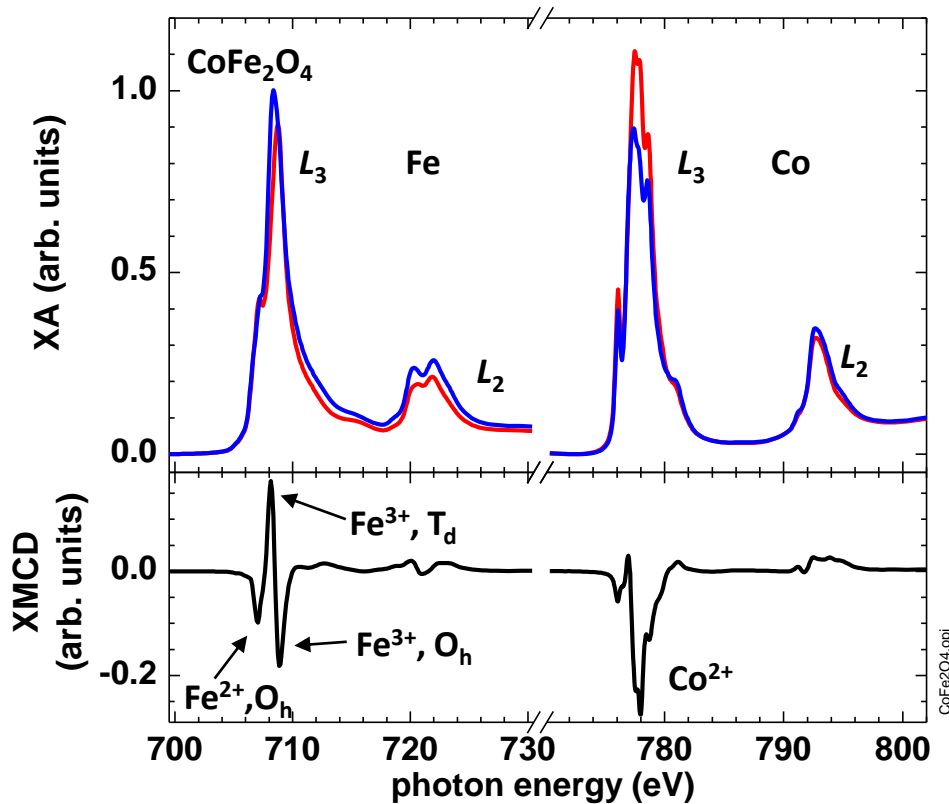
Magnitude of XMCD depends on

- + expectation value of 3d magnetic moment
- + degree of circular photon polarization,  $P_{\text{circ}}$
- + geometry

# X-RAY MAGNETIC CIRCULAR DICHROISM (XMCD)



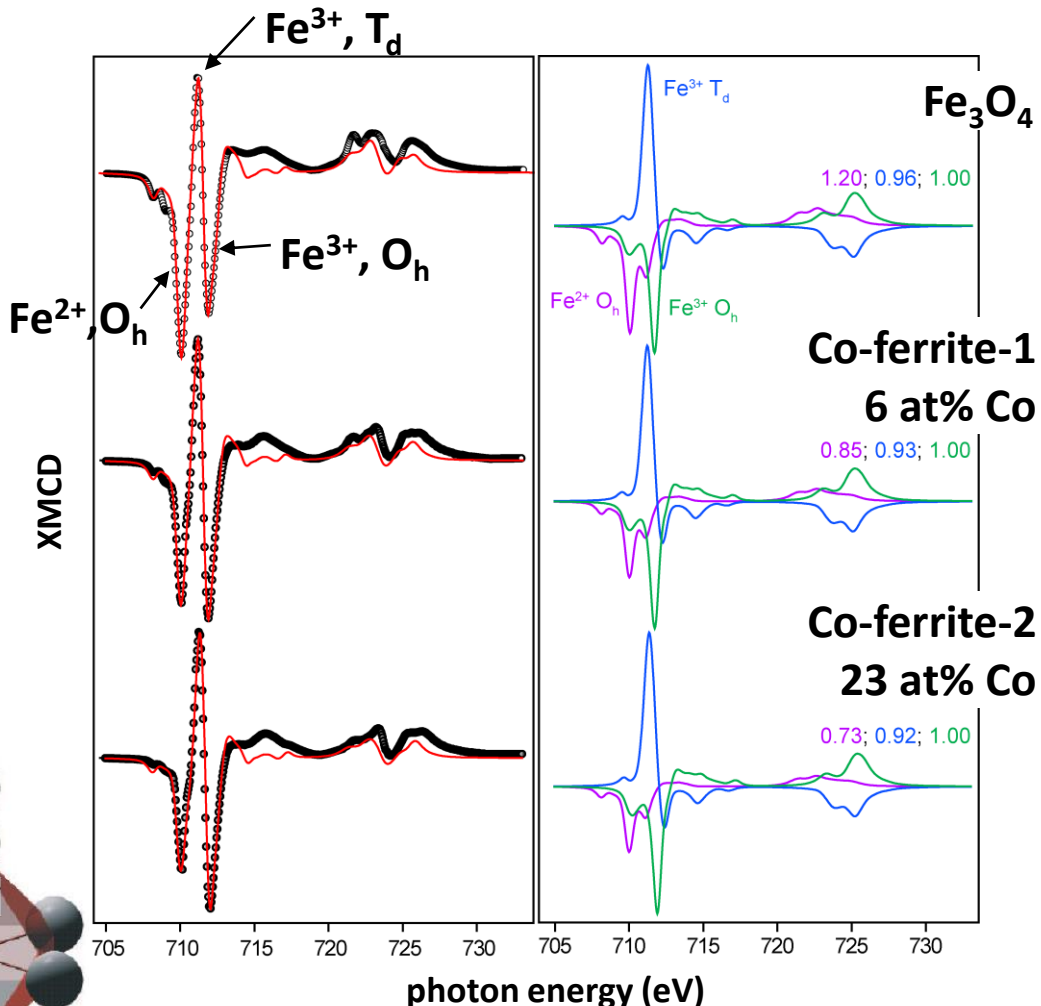
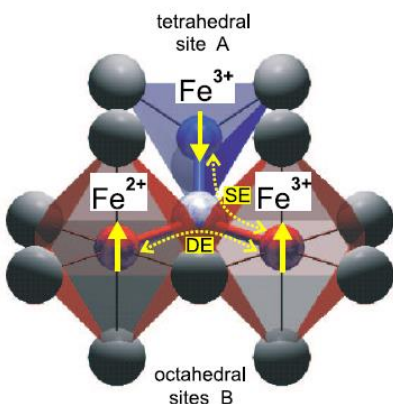
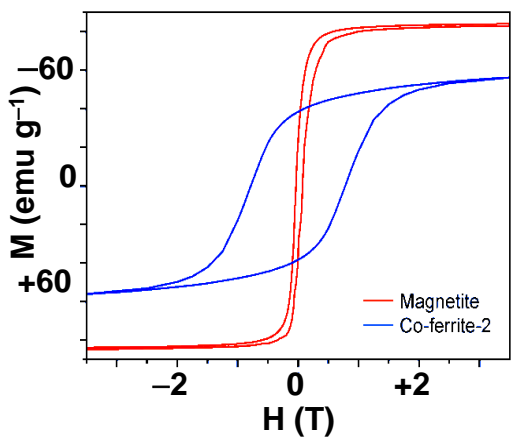
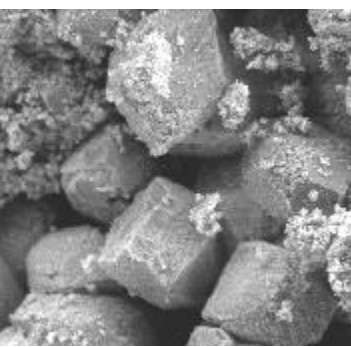
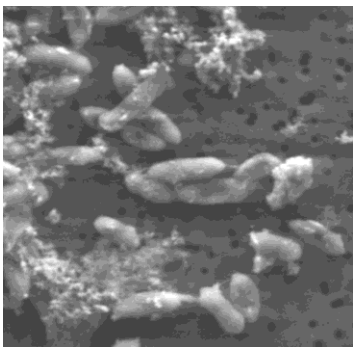
J. Stöhr, H.C. Siegmann,  
Magnetism (Springer)



- + XMCD provides magnetic information resolving
  - elements Fe, Co, ...
  - valence states:  $\text{Fe}^{2+}$ ,  $\text{Fe}^{3+}$ , ...
  - lattice sites: octahedral,  $\text{O}_h$ , tetrahedral,  $\text{T}_d$ ,

# MAGNETIC BIONANOSPINELS

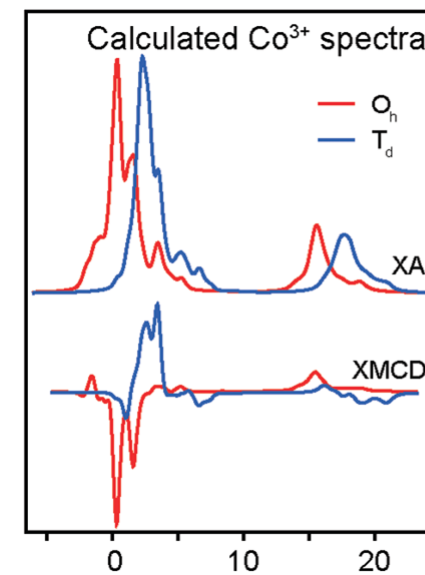
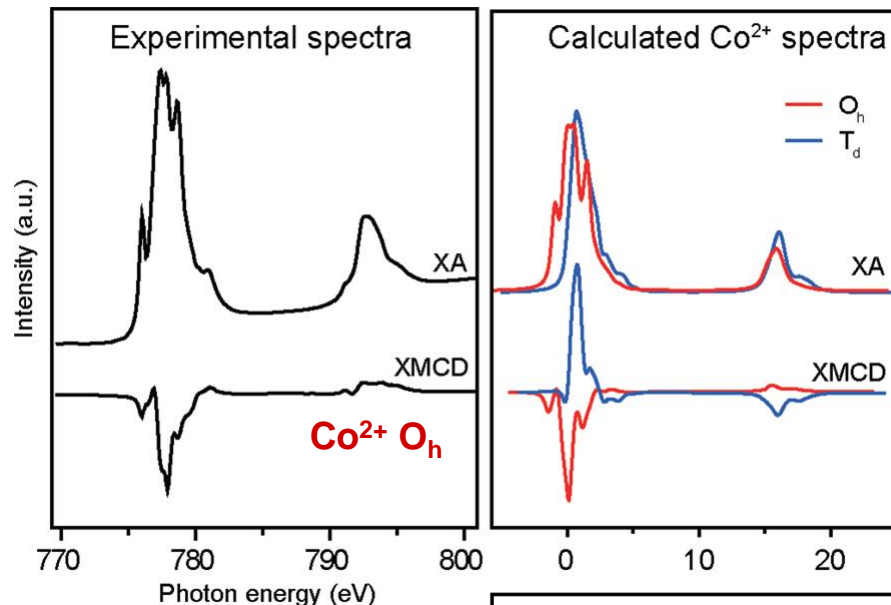
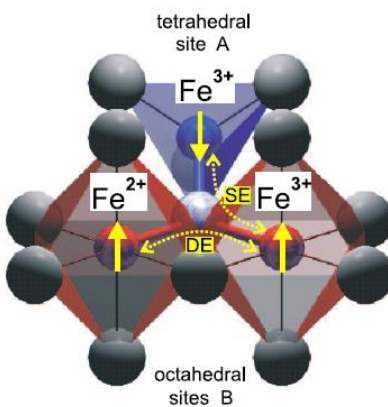
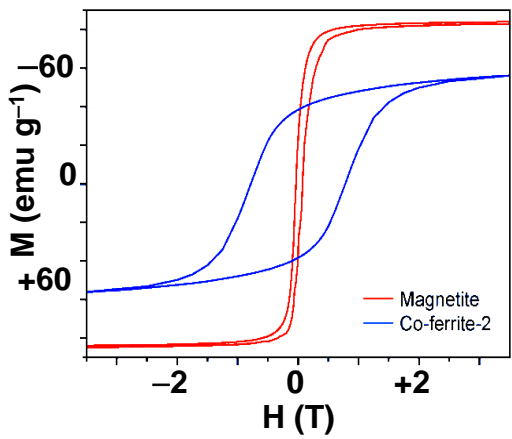
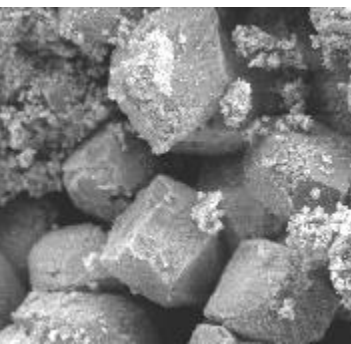
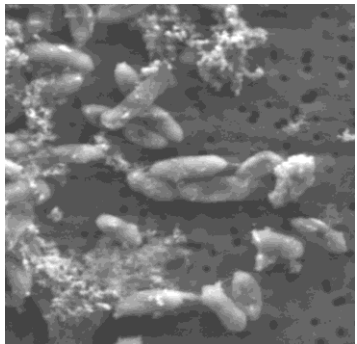
+ *Geobacter sulfurreducens* bacteria form magnetite nanocrystals (15nm) via extracellular reduction of amorphous Fe(III)-bearing minerals



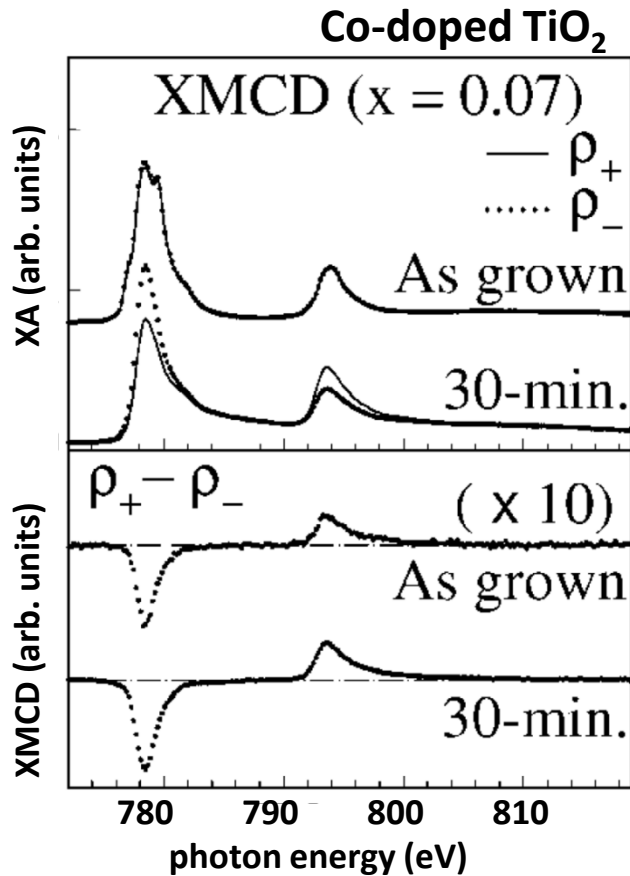
V. Cocker et al.,  
Eur. J. Mineral. 19, 707–716 (2007)

# MAGNETIC BIONANOSPINELS

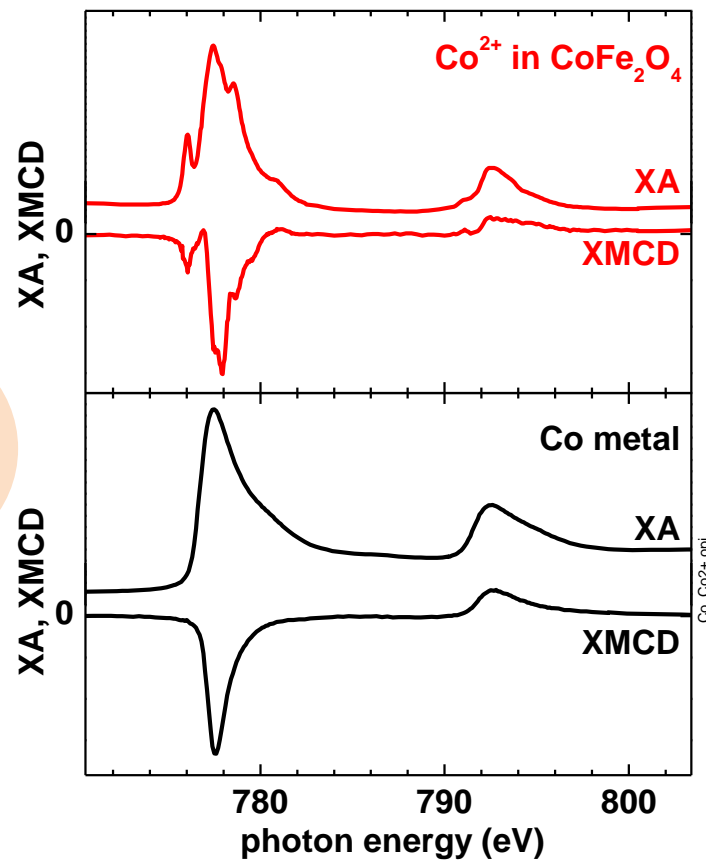
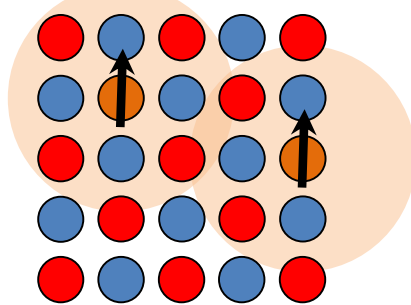
+ **Geobacter sulfurreducens bacteria form magnetite nanocrystals (15nm) via extracellular reduction of amorphous Fe(III)-bearing minerals**



# Co-DOPED TiO<sub>2</sub>



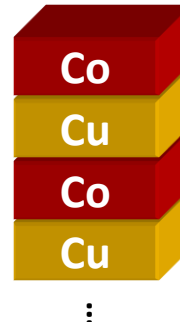
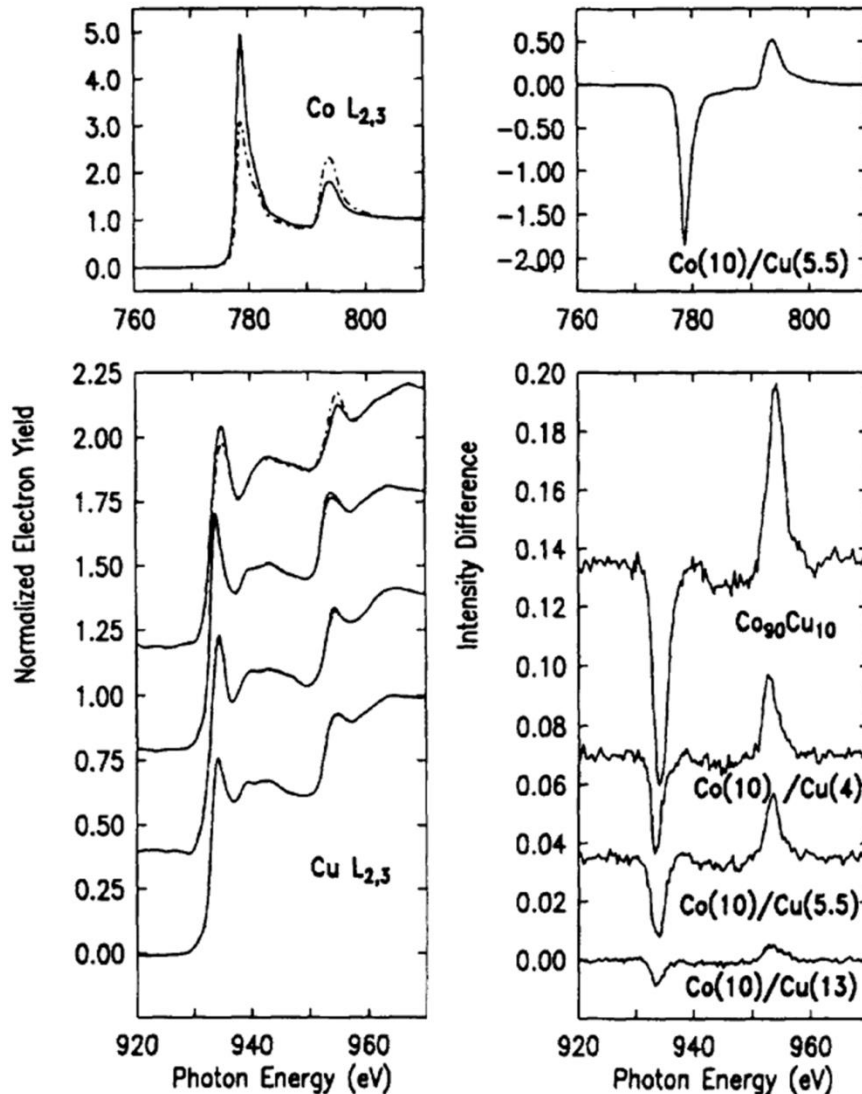
dilute magnetic semiconductors



J.-Y. Kim *et al.*,  
 Phys. Rev. Lett. **90**, 017401 (2003)

- + Comparing XMCD spectra with model compounds and/or calculations
- ⇒ Identifying magnetic phases

# INDUCED MOMENTS AT Co/Cu INTERFACES

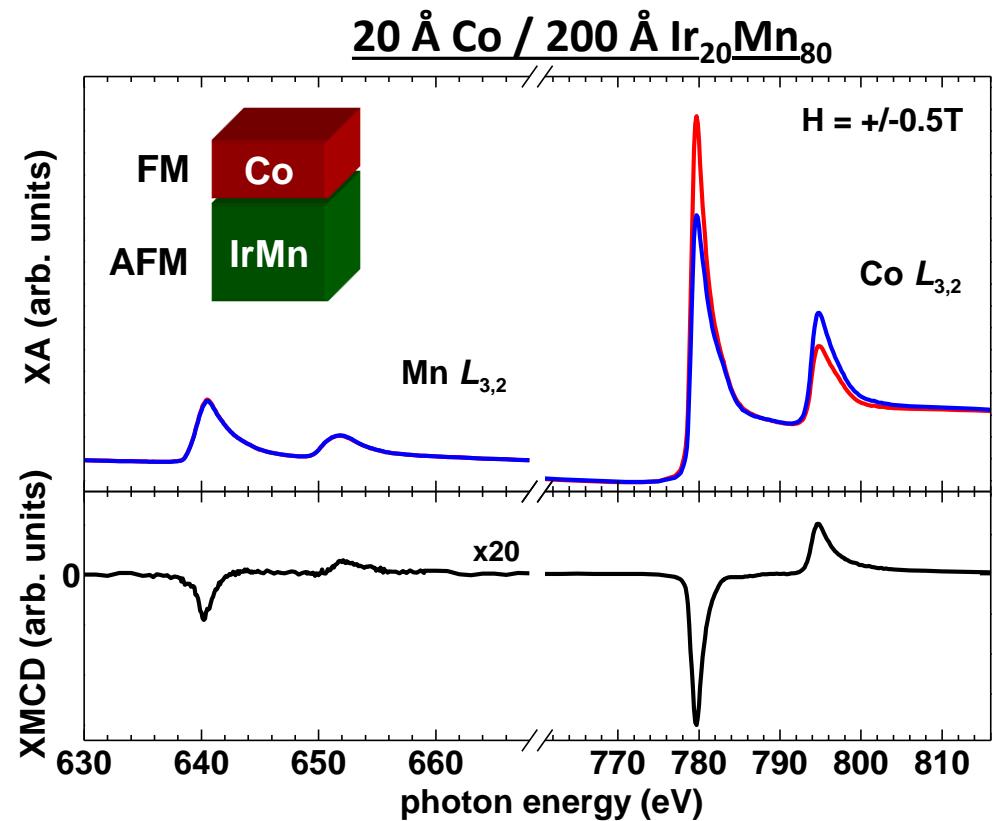


+ The element-specificity makes XMCD measurements an ideal tool to determine induced moments at interfaces between magnetic and non-magnetic elements.

M. G. Samant *et al.*,  
Phys. Rev. Lett. 72, 1112 (1994)

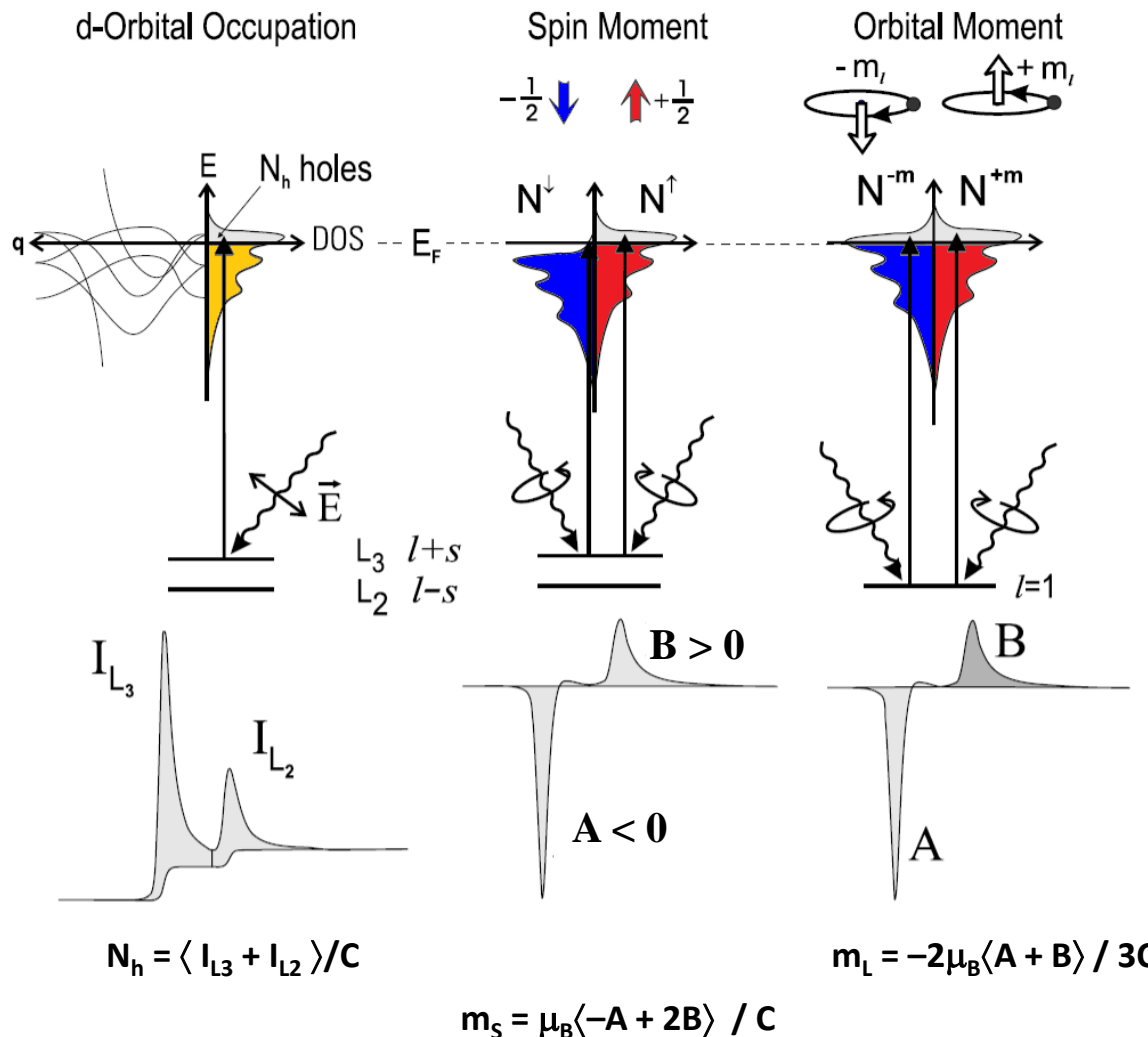
# MAGNETIC INTERFACES

- + Weak Mn XMCD signal  
⇒ Uncompensated Mn at Co/IrMn interface
- + Same sign of XMCD signal for Co and Mn  
⇒ Parallel coupling of Co and Mn moments
- + Nominal thickness of uncompensated interface moments:  $(0.5 \pm 0.1) \text{ML}$



H. Ohldag *et al.*,  
Phys. Rev. Lett. **91**, 017203 (2003)

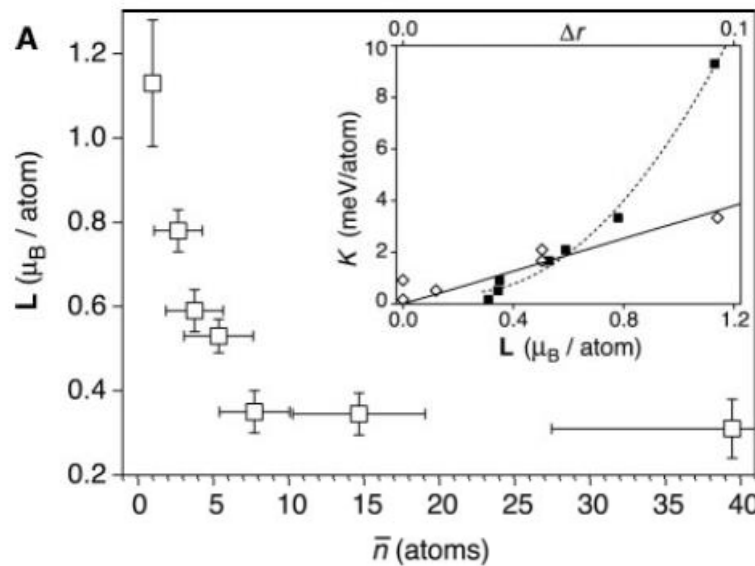
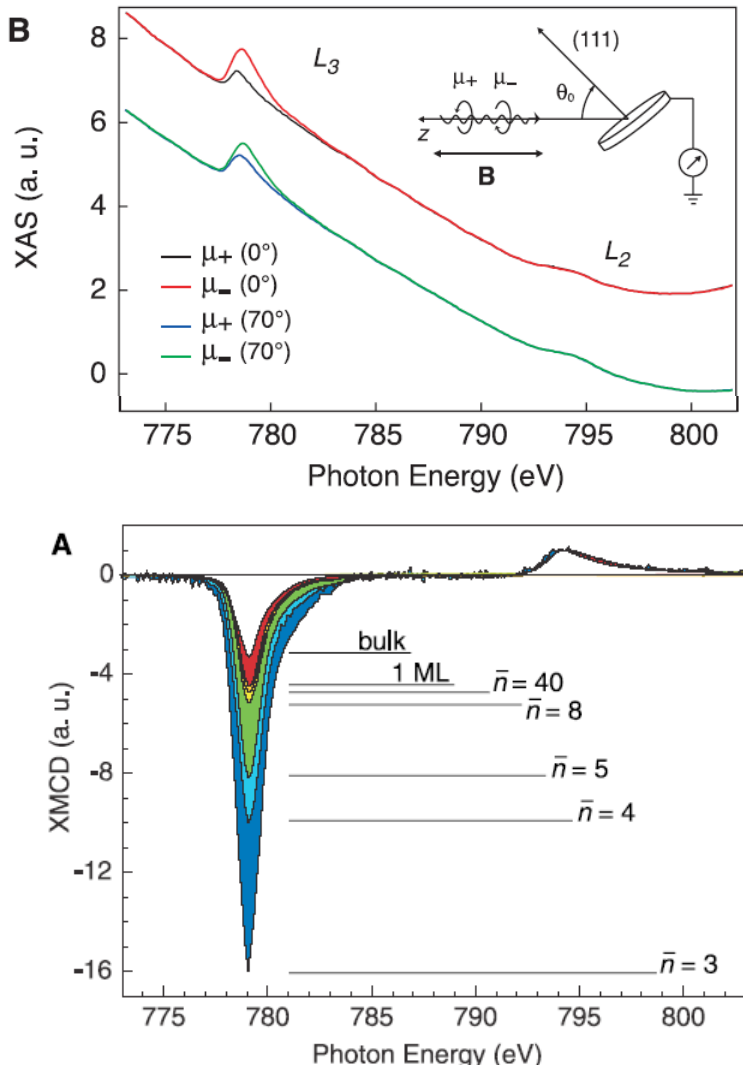
# SUM RULES



• Theoretically derived sum rules correlate XMCD spectra with spin and orbital moment providing unique tool for studying magnetic materials.

J. Stöhr, H.C. Siegmann,  
Magnetism (Springer)

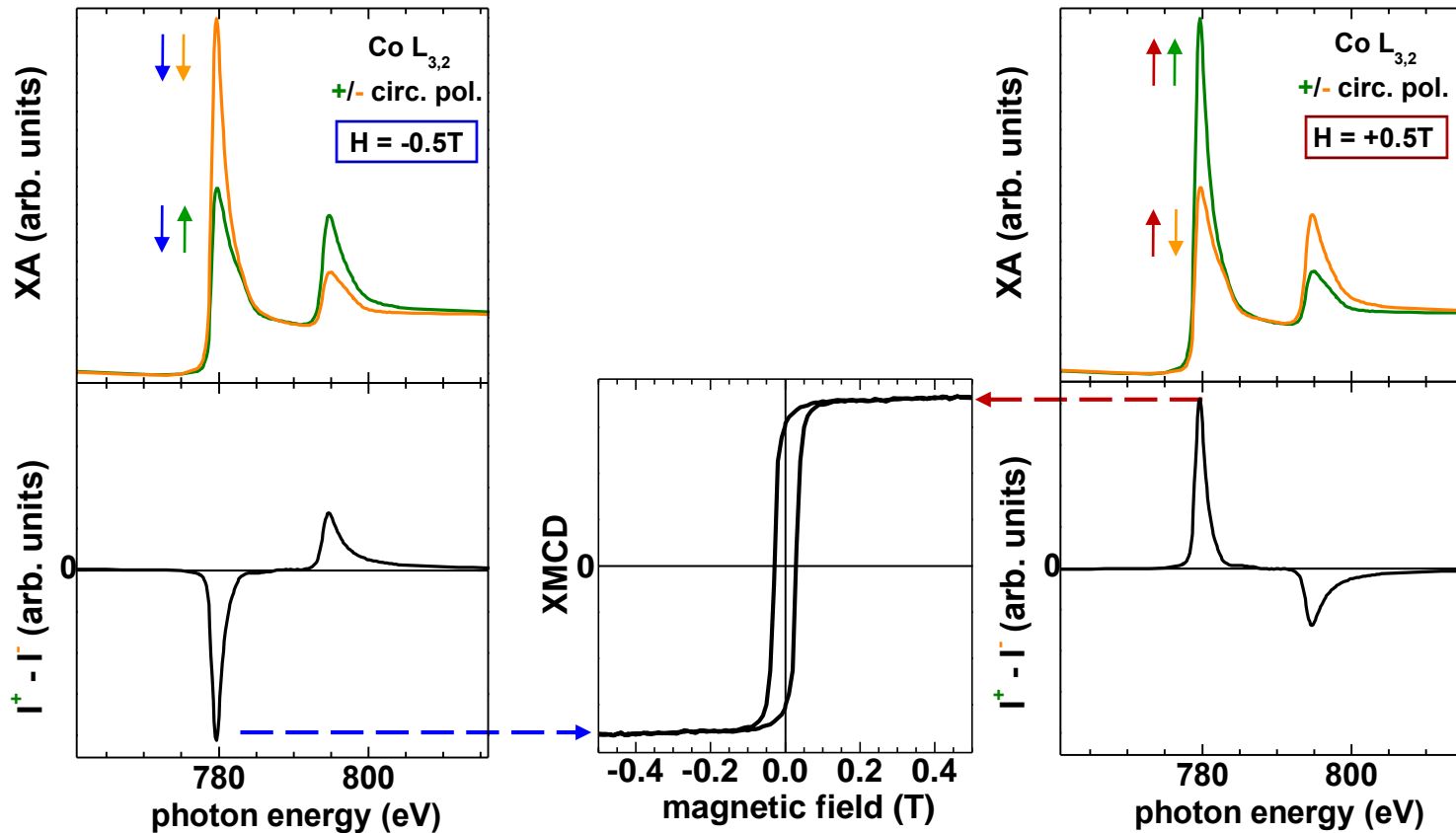
# ORBITAL MOMENT OF CO NANOPARTICLES



- + Strong variation of orbital and spin magnetic moment observable as change in relative  $L_3$  and  $L_2$  intensity in XMCD spectrum.
- + Co atoms and nanoparticles on Pt have enhanced orbital moments up to  $1.1 \mu_B$

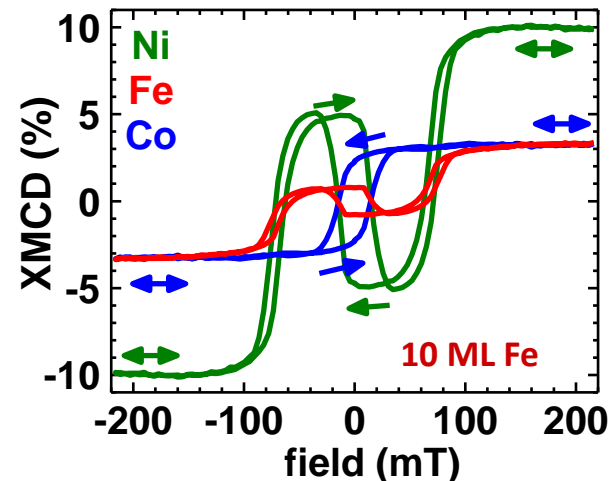
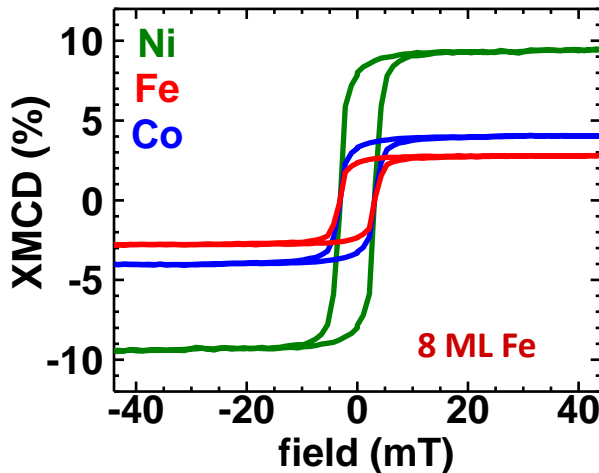
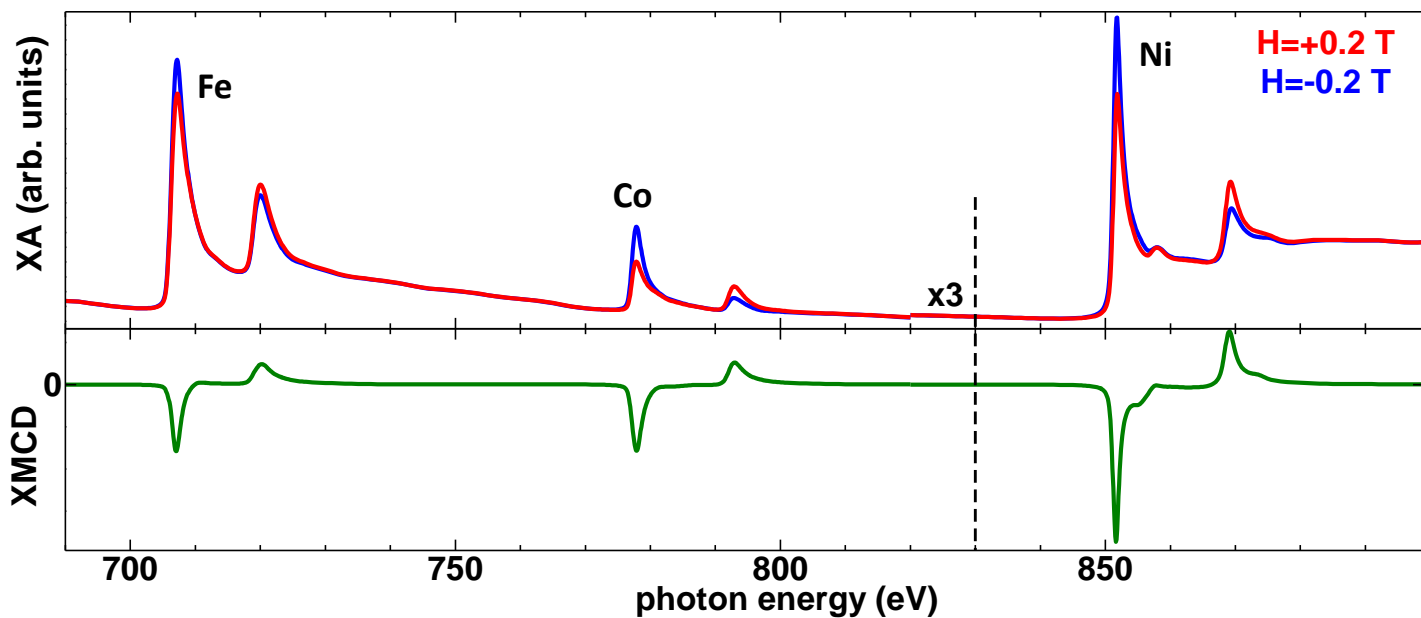
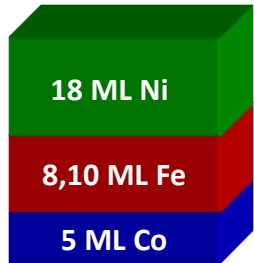
P. Gambardella *et al.*,  
Science 300, 1130 (2003)

# ELEMENT-SPECIFIC MAGNETIZATION REVERSAL

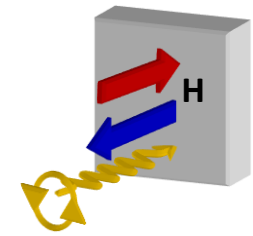


- + Monitoring field dependence of XMCD
- Element-specific information on magnetization reversal in complex magnetic nanostructures.

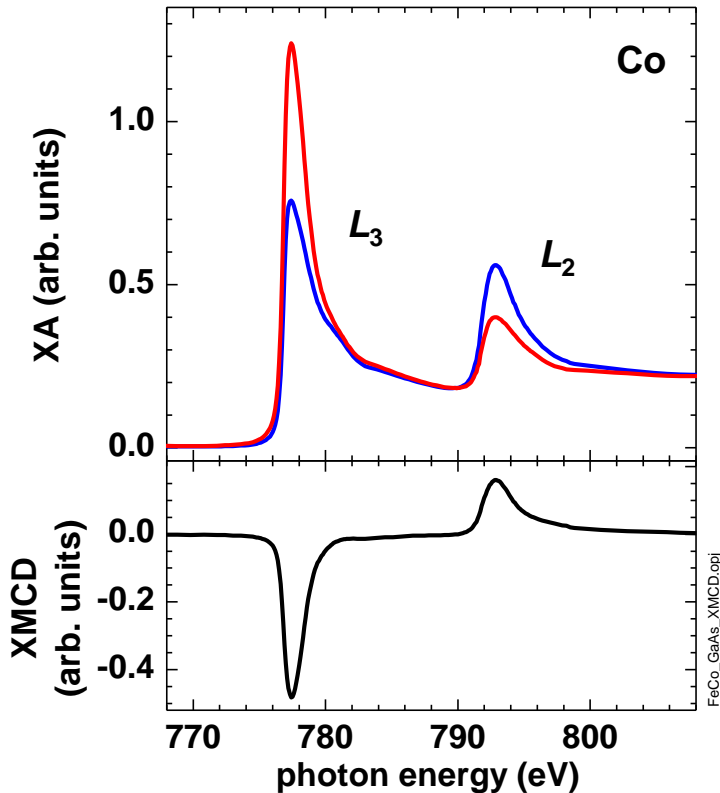
# ELEMENT-SPECIFIC MAGNETIZATION REVERSAL



# X-RAY FERROMAGNETIC RESONANCE

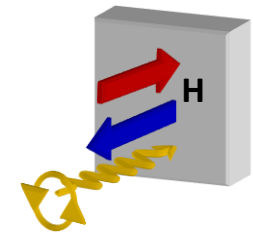


circularly  
polarized

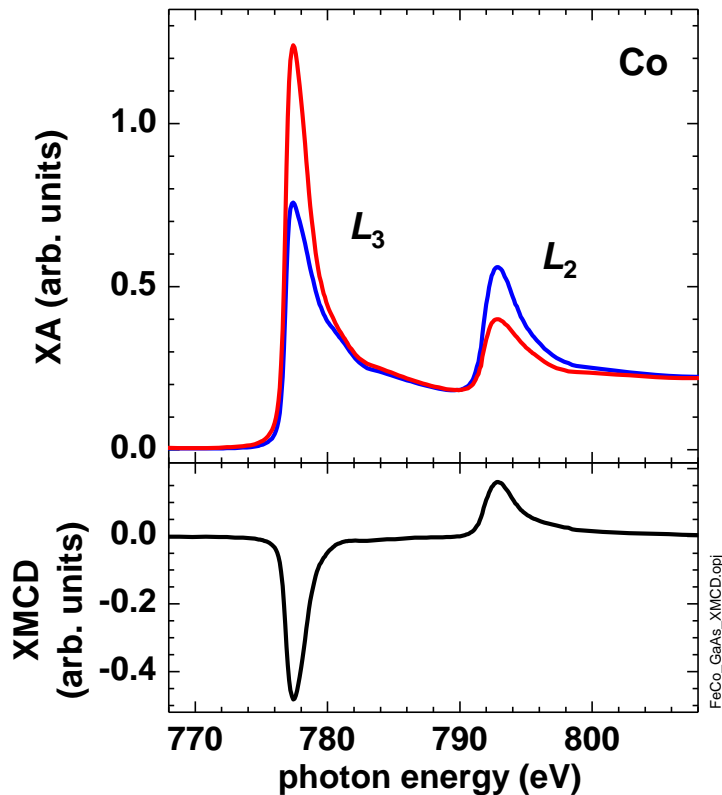


- + XMCD is the difference in x-ray absorption between antiparallel and parallel orientation of magnetic moment and photon spin.
- + The XMCD magnitude reflects the magnetic moment aligned parallel to the x ray beam.

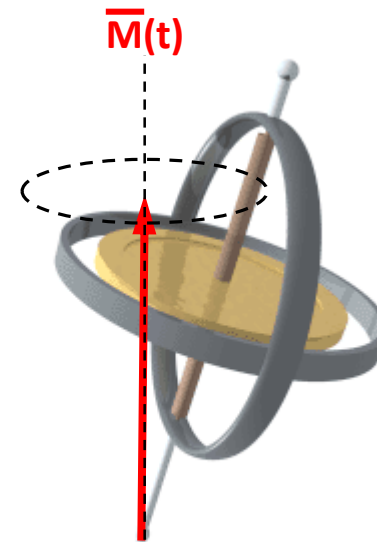
# X-RAY FERROMAGNETIC RESONANCE



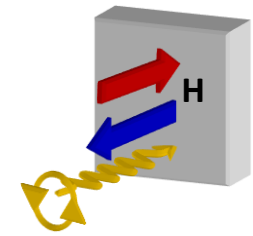
circularly  
polarized



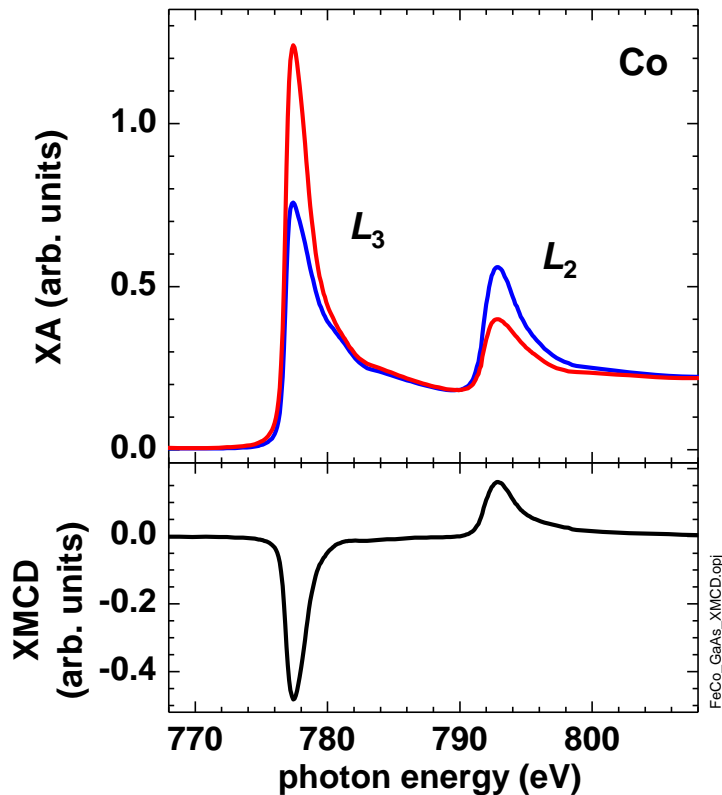
+ In fact:  
Magnetic moments are not  
fully aligned with applied fields  
but precess around them.



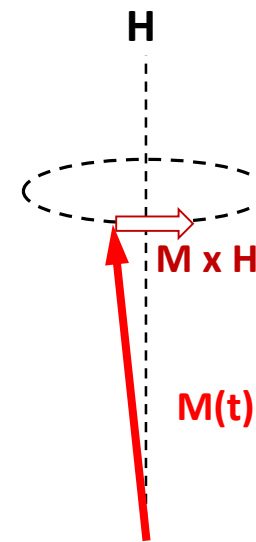
# X-RAY FERROMAGNETIC RESONANCE



circularly  
polarized

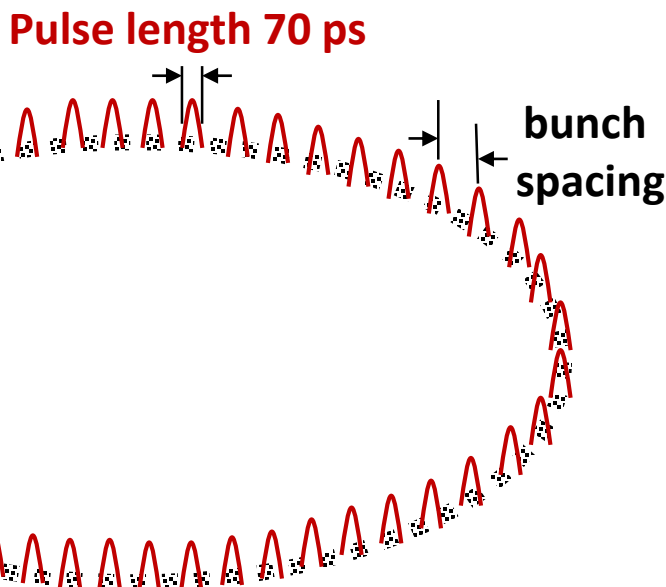


- + In fact:  
Magnetic moments are not fully aligned with applied fields but precess around them.



- + Is it possible to measure the precession of magnetic moments making use of the pulsed nature of synchrotron radiation and XMCD?

# X-RAY FERROMAGNETIC RESONANCE

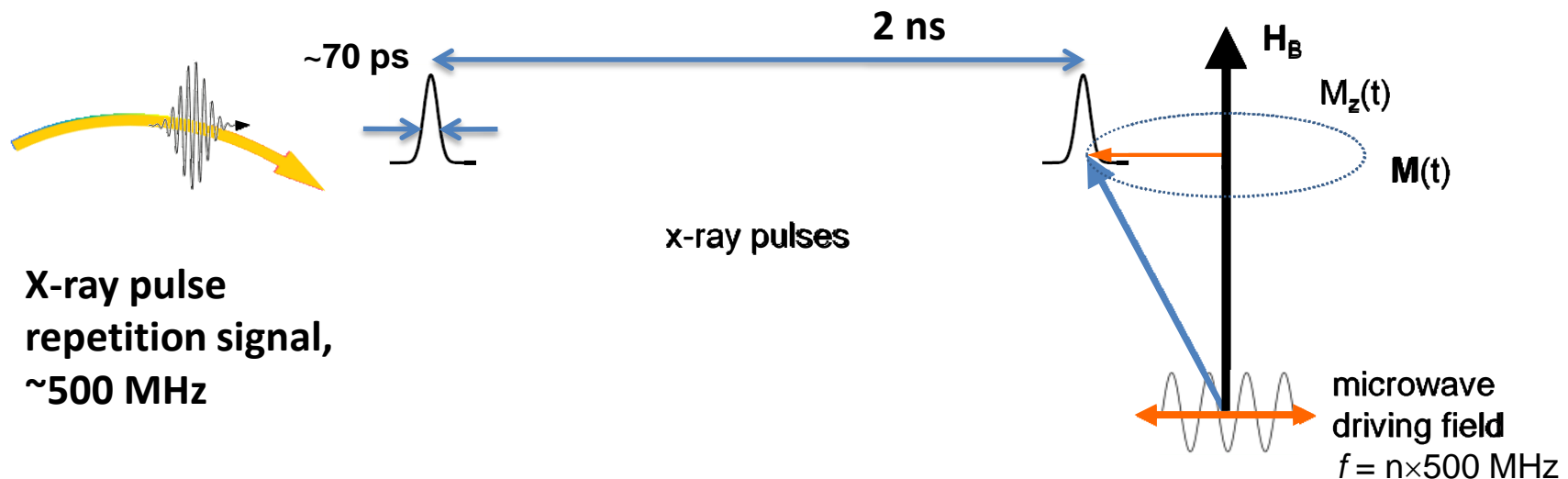


Pulsed nature of synchrotron radiation  
Example: Advanced Light Source

- + 256-320 bunches for 500mA beam current
- + Bunch spacing: 2 ns
- + Pulse length 70ps

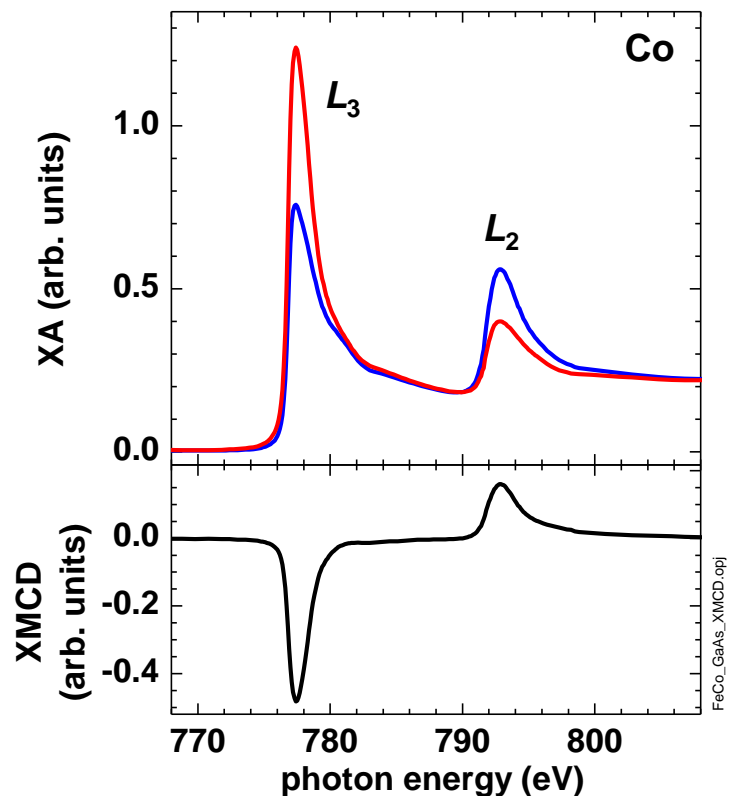
# X-RAY FERROMAGNETIC RESONANCE

Dynamic XMCD measurement, i.e. synchronize x-ray pulses with FMR precession

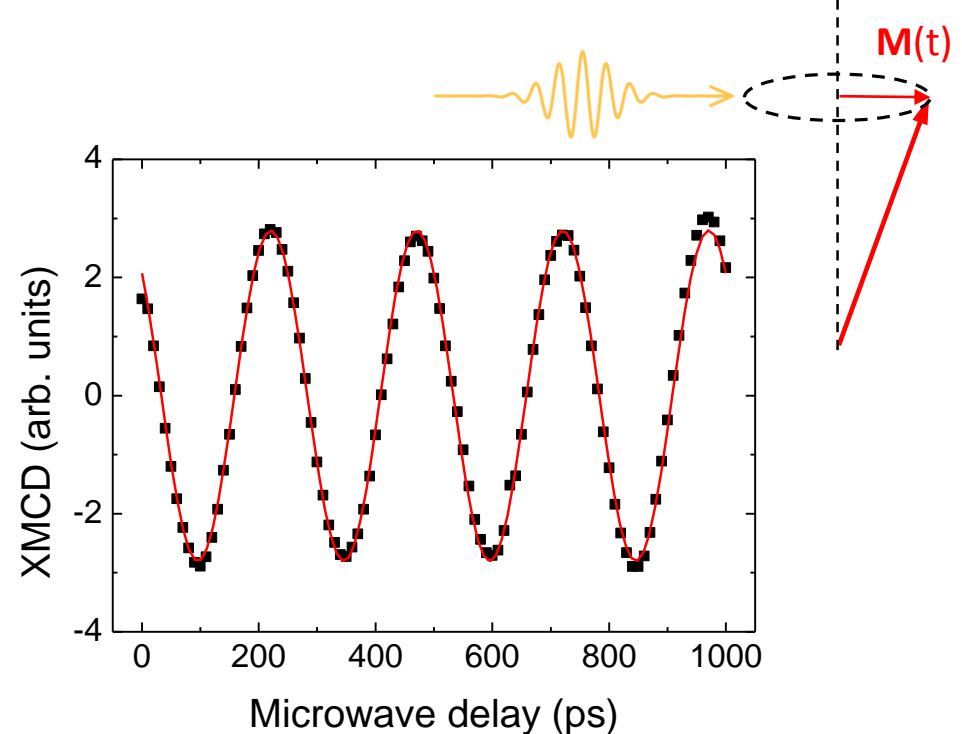


# X-RAY FERROMAGNETIC RESONANCE

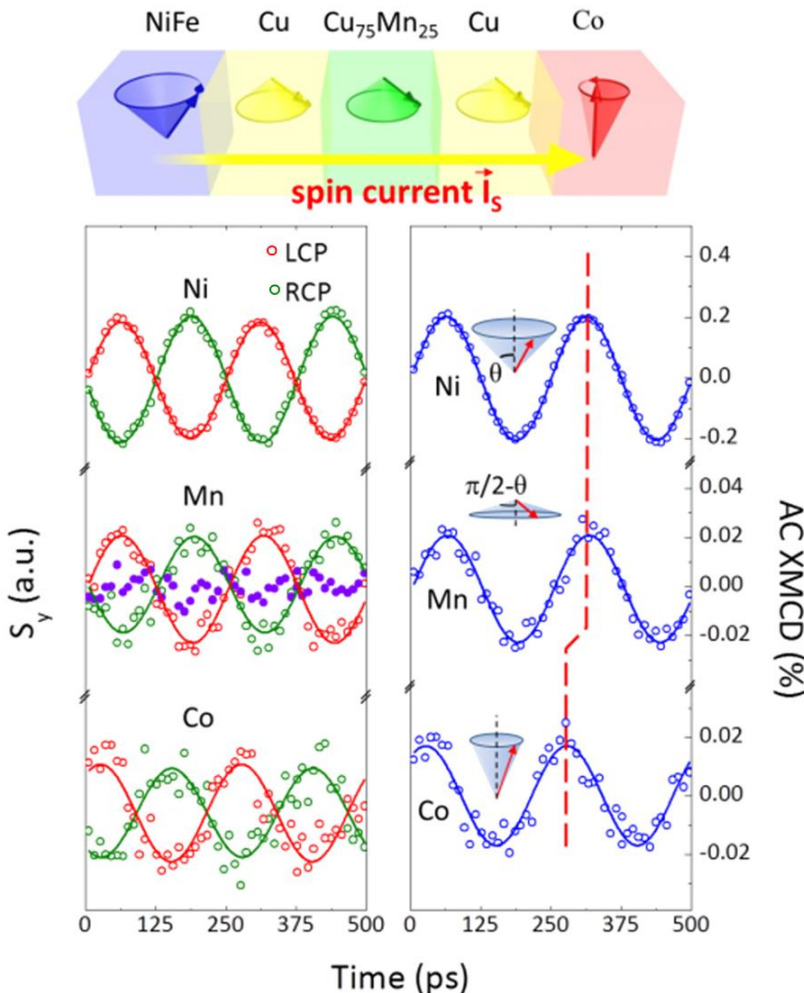
Static XMCD



Dynamic XMCD

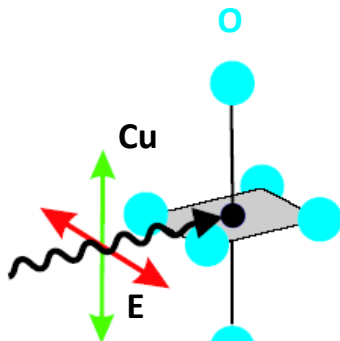


# X-RAY FERROMAGNETIC RESONANCE

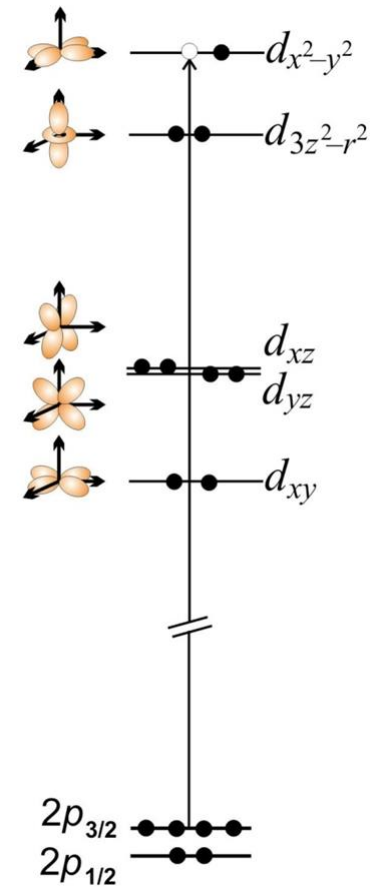
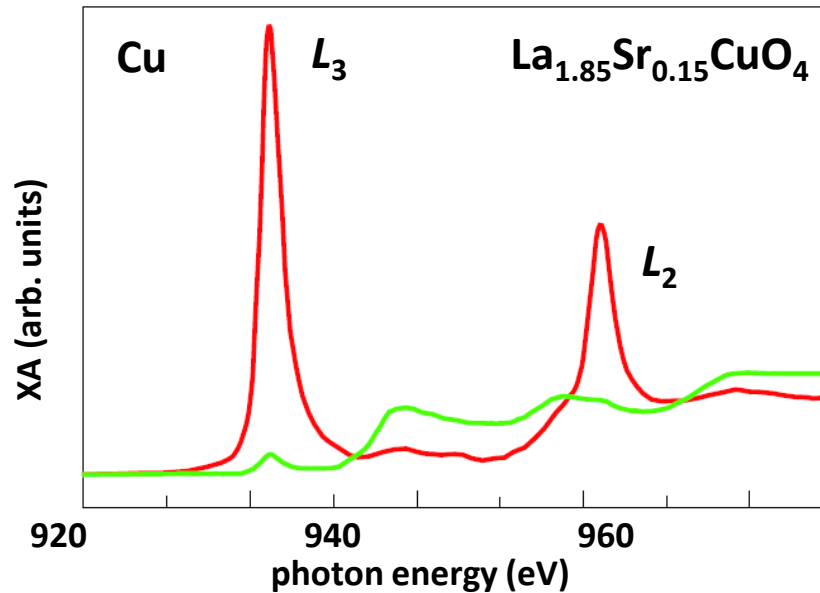


- + Precession is resonantly excited in the NiFe layer with an 4 GHz RF field.
- + The resonance field of the CO layer is higher, i.e. no precession is excited in the Co layer.
- + Precession in Py, Cu<sub>75</sub>Mn<sub>25</sub>, and Co layers are probed by XMCD using left- and right-circularly polarized x-rays at Ni, Mn, and Co edges, respectively.
- + The Cu<sub>75</sub>Mn<sub>25</sub> spin precession is a direct indicator of the AC spin current through the structure.

# X-RAY LINEAR DICHROISM



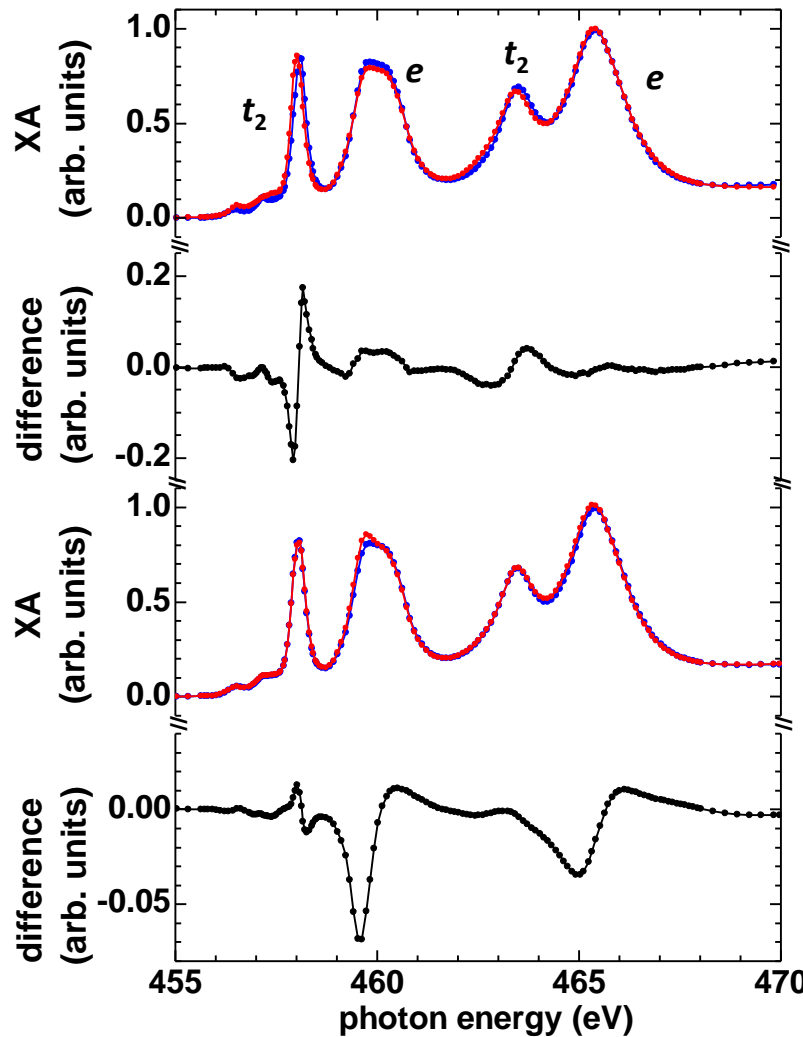
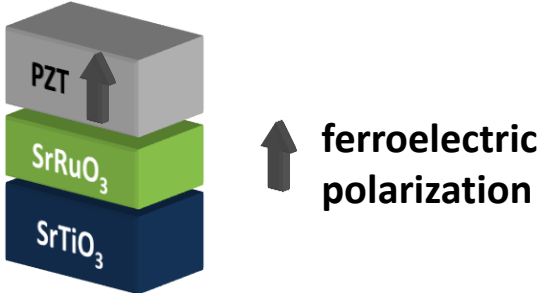
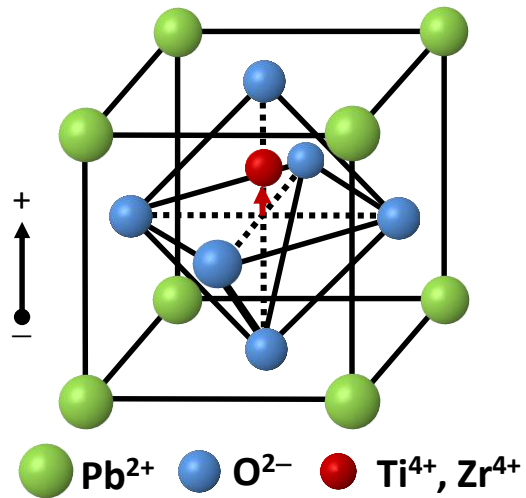
C. T. Chen et al.  
PRL 68, 2543 (1992)



## X-Ray Linear Dichroism:

- + Difference in x-ray absorption for different linear polarization direction relative to crystalline and/or spin axis.
- + Due to the anisotropic charge distribution about the absorbing atom caused by bonding and/or magnetic order.
- + “Search Light Effect”: X-ray absorption of linear polarized x rays proportional to density of empty valence states in direction of electric field vector E.

# STRUCTURAL CHANGES IN $\text{PbZr}_{0.2}\text{Ti}_{0.8}\text{O}_3$



E. Arenholz *et al.*,  
Phys. Rev. B **82**, 140103 (2010)

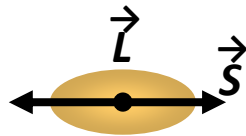
- Spontaneous electric polarization due to off-center shift of  $\text{Ti}^{4+}, \text{Zr}^{4+}$  associated with tetragonal distortion  $\Leftrightarrow$  linear dichroism

- Reversing ferroelectric polarization changes XA  $\Leftrightarrow$  Change in tetragonal distortion

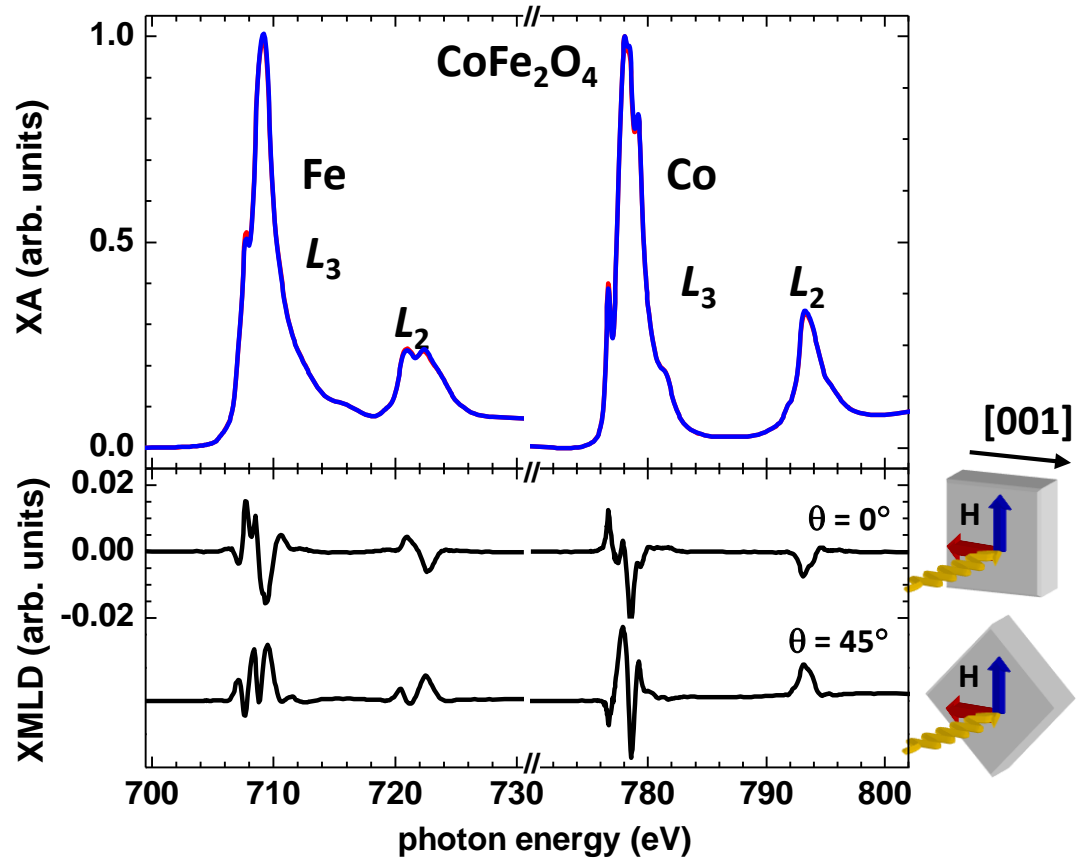
# X-RAY MAGNETIC LINEAR DICHROISM



Isotropic  $d$  electron charge density  
⇒ No polarization dependence



Magnetically aligned system  
⇒ Spin-orbit coupling distorts  
charge density  
⇒ Polarization dependence

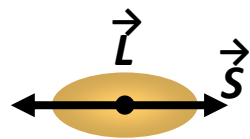


- +  $I_{\text{XMLD}} = I_{||} - I_{\perp} \propto \langle m^2 \rangle$ ,  $\langle m^2 \rangle$  = expectation value of square of atomic magnetic moment
- + XMLD allows investigating ferri- and ferromagnets as well as antiferromagnets
- + XMLD spectral shape and angular dependence are determined by magnetic order and lattice symmetry

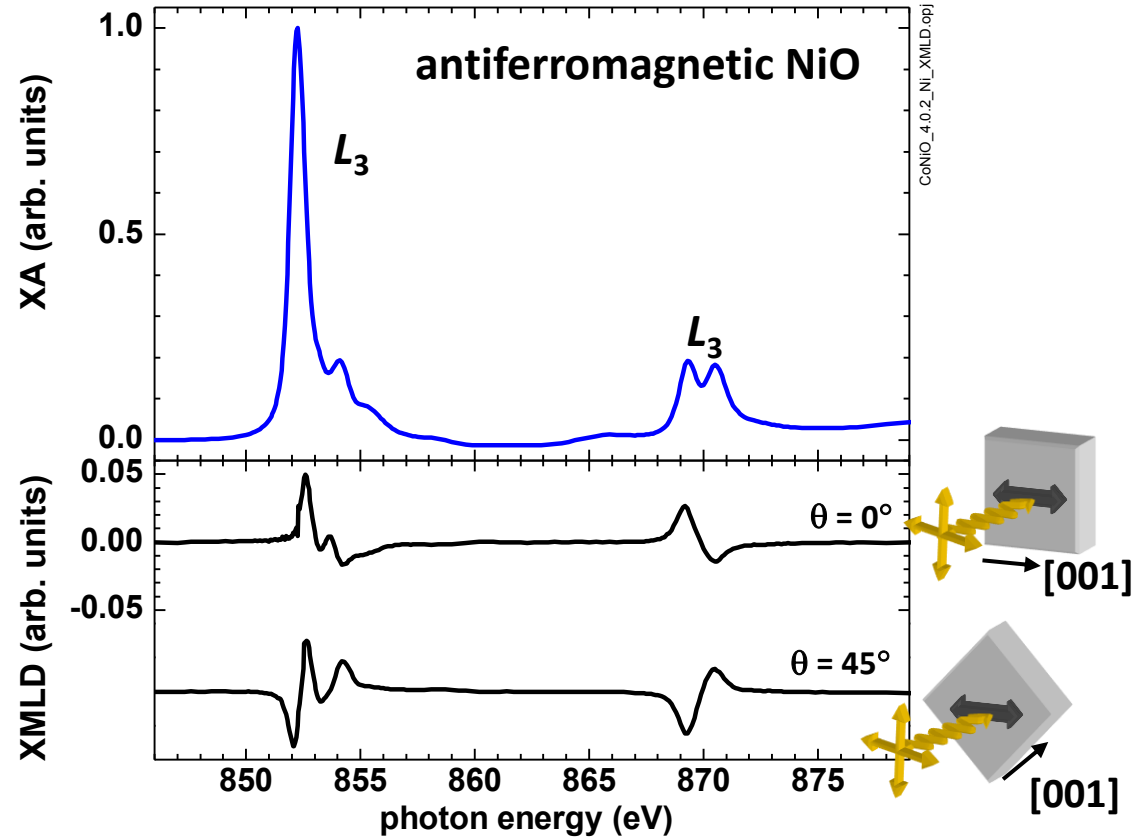
# X-RAY MAGNETIC LINEAR DICHROISM



Isotropic  $d$  electron charge density  
⇒ No polarization dependence

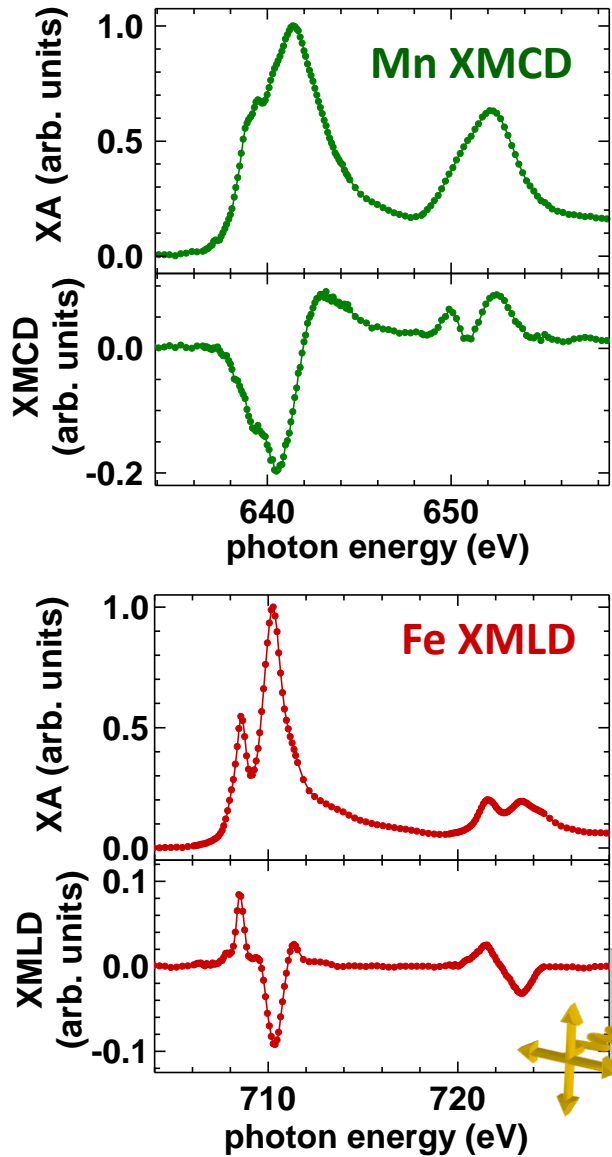


Magnetically aligned system  
⇒ Spin-orbit coupling distorts  
charge density  
⇒ Polarization dependence

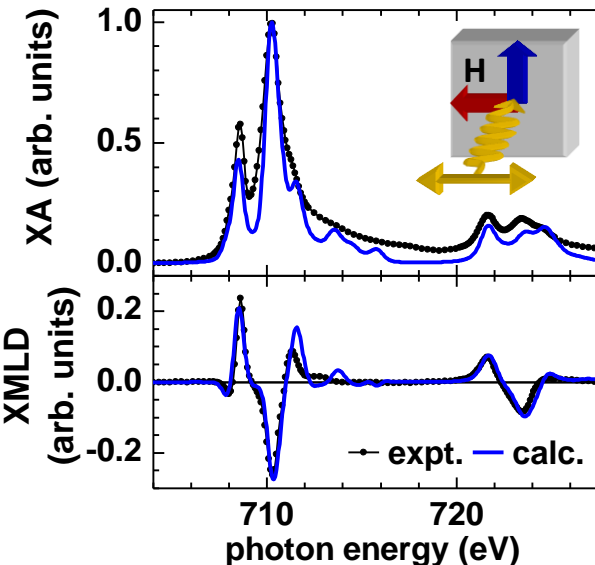


- +  $I_{\text{XMLD}} = I_{||} - I_{\perp} \propto \langle m^2 \rangle$ ,  $\langle m^2 \rangle$  = expectation value of square of atomic magnetic moment
- + XMLD allows investigating ferri- and ferromagnets as well as antiferromagnets
- + XMLD spectral shape and angular dependence are determined by magnetic order and lattice symmetry

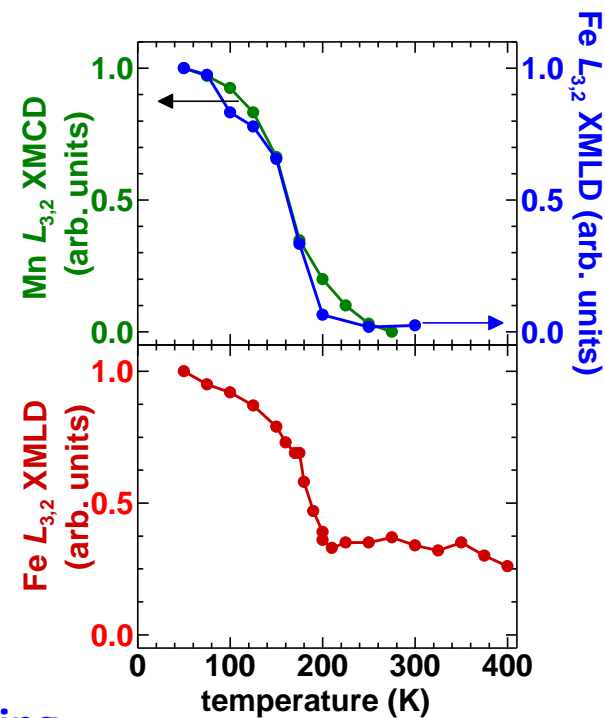
# MAGNETIC COUPLING AT INTERFACES



$\text{La}_{0.7}\text{Sr}_{0.3}\text{MnO}_3$  (LSMO)  
ferromagnet  
 $\text{La}_{0.7}\text{Sr}_{0.3}\text{FeO}_3$  (LSFO)  
antiferromagnet

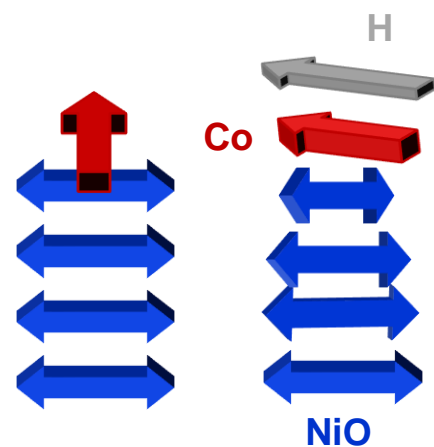
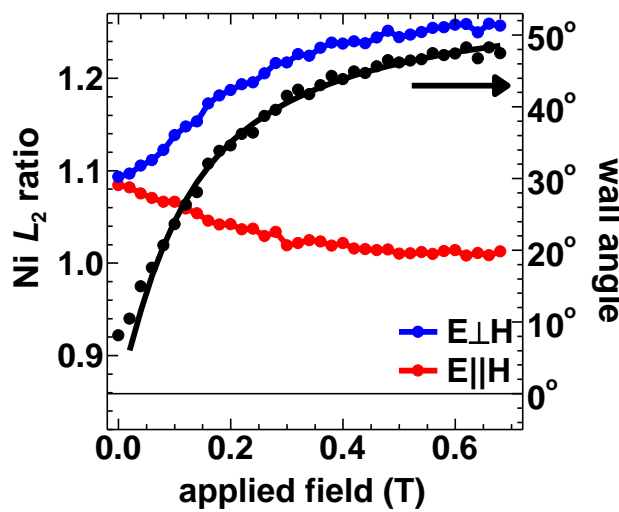
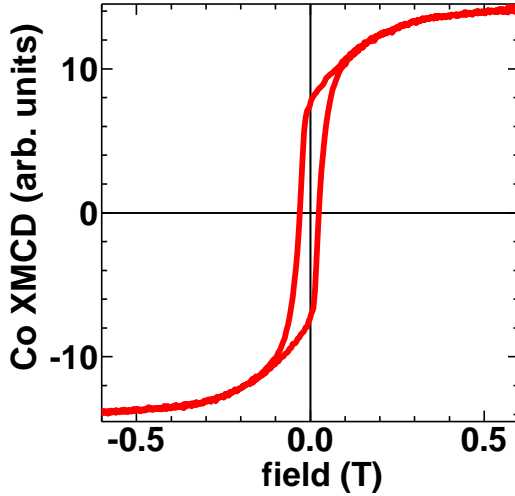
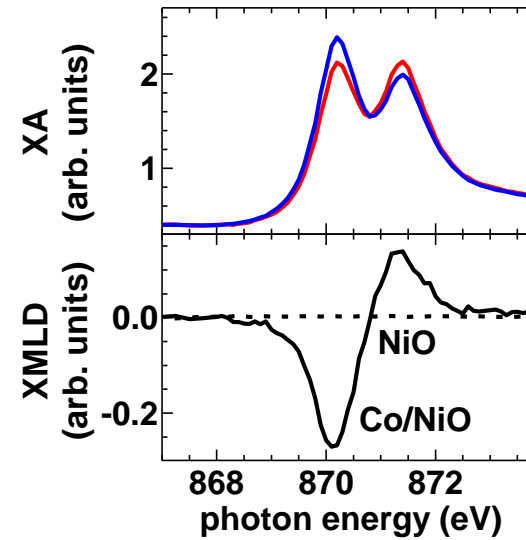
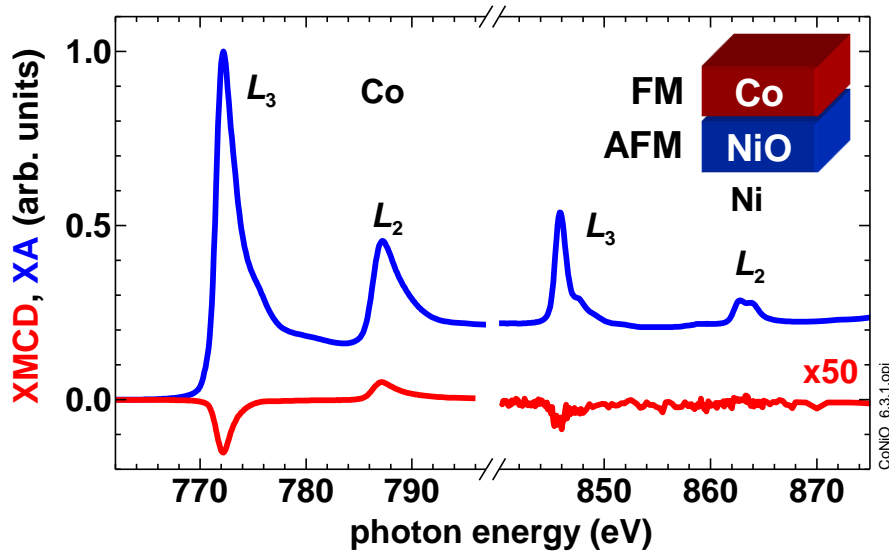
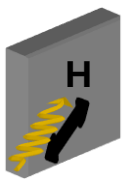


→ Perpendicular coupling  
at LSMO/LSFO interface



E. Arenholz *et al.*,  
Appl. Phys. Lett. **94**, 072503 (2009)

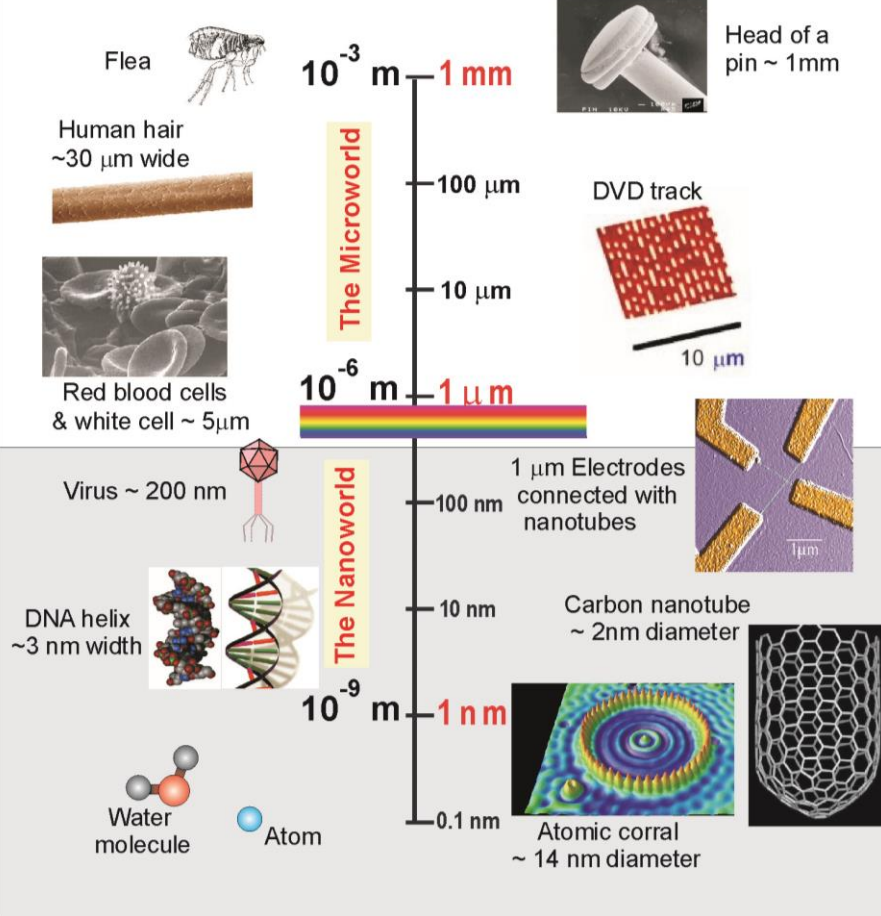
# PLANAR DOMAIN WALL



A. Scholl *et al.*,  
Phys. Rev. Lett. **92**, 247201 (2004)

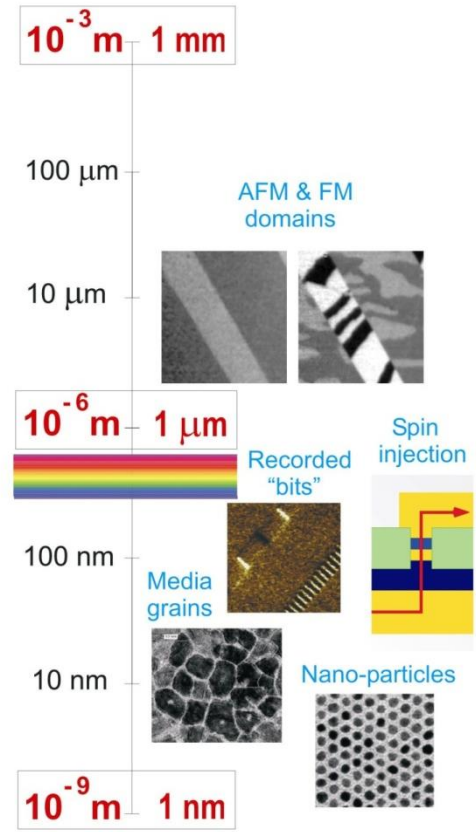
# MAGNETIC MICROSCOPY

## Nature



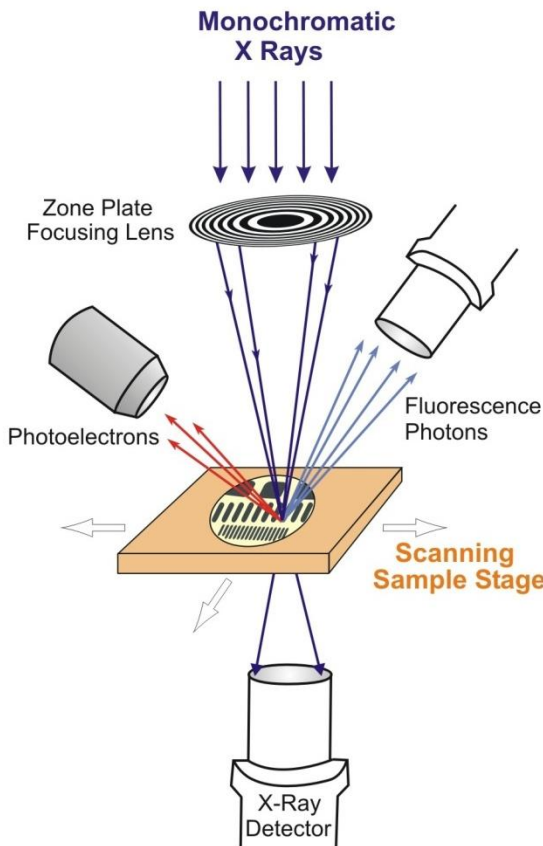
## Technology

### The Nanoworld

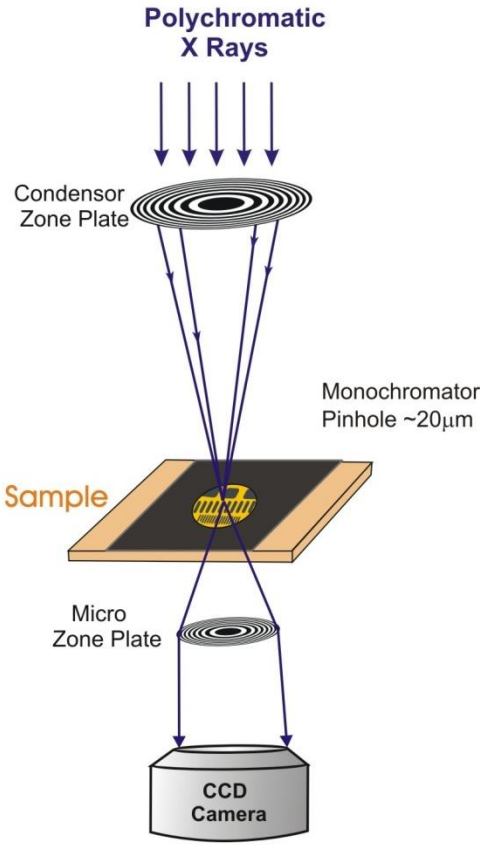


# MAGNETIC MICROSCOPY

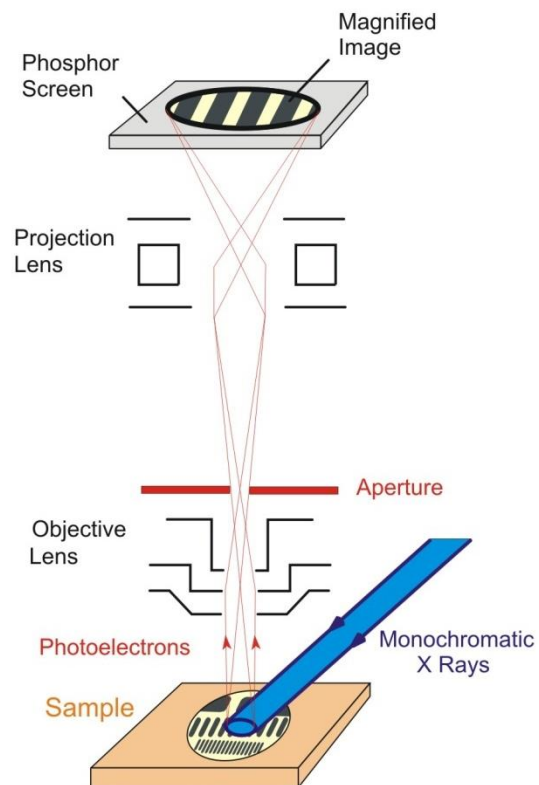
Scanning Transmission X-ray Microscopy  
STXM



Transmission X-ray Microscopy  
TXM



X-Ray Photoemission Electron Microscopy  
XPEEM

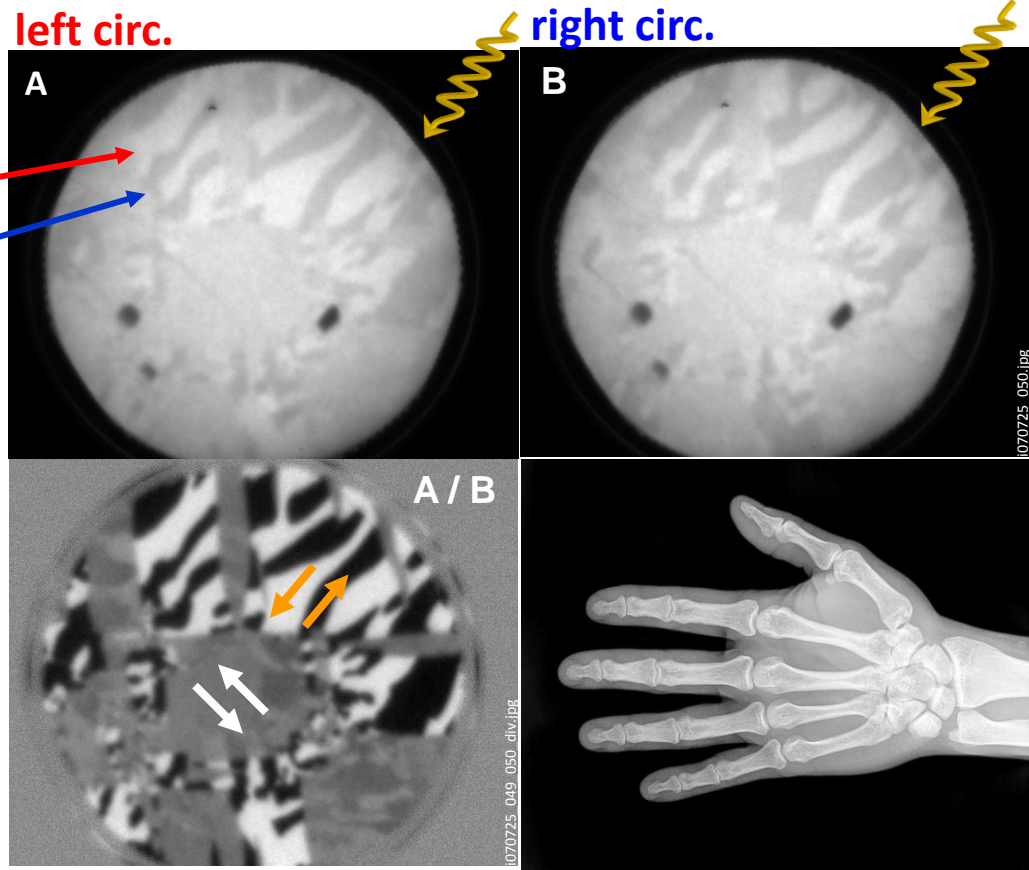
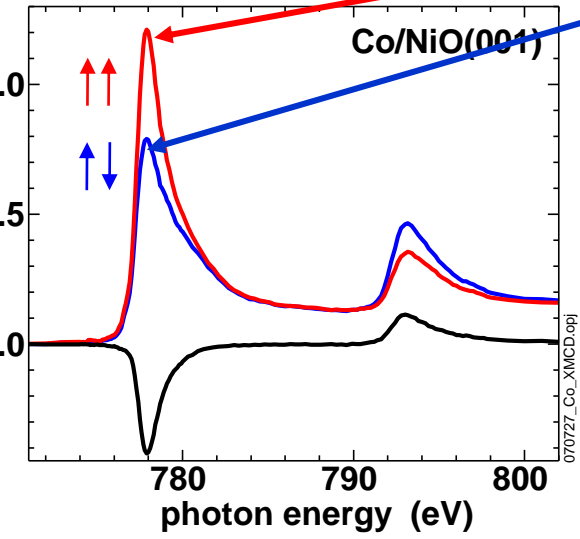


10-50 nm spatial resolution

J. Stöhr, H.C. Siegmann,  
Magnetism (Springer)

# IMAGING MAGNETIC DOMAINS USING X-RAYS

XMC<sub>D</sub>, XA (arb. units)

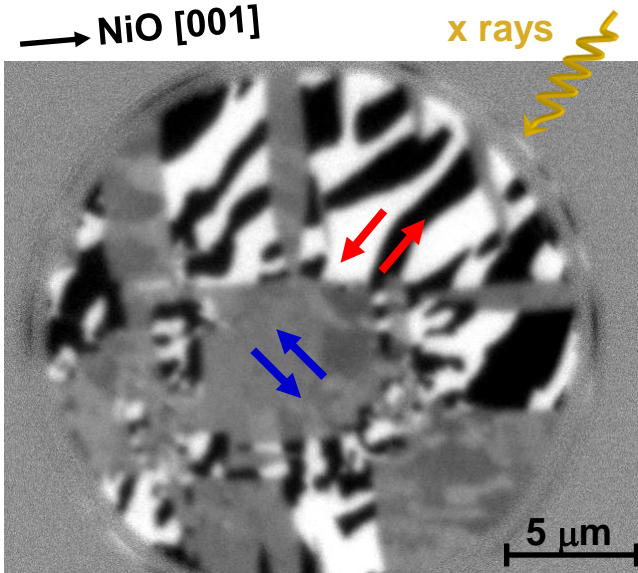


E. Arenholz *et al.*,  
Appl. Phys. Lett. **93**, 162506 (2008)

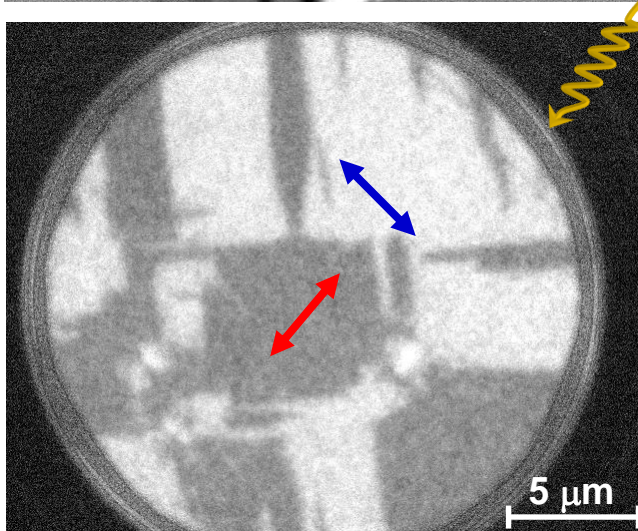
- + Images taken with left and right circularly polarized x-rays at photon energies with XMCD, i.e. Co  $L_3$  edge, provide magnetic contrast and domain images.

# MAGNETIC COUPLING AT Co/NiO INTERFACE

Co XMCD



Ni XMLD



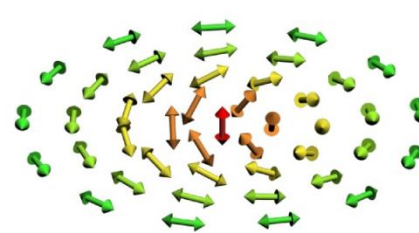
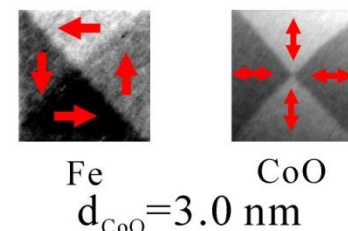
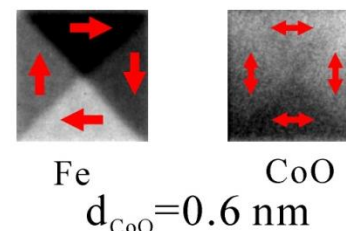
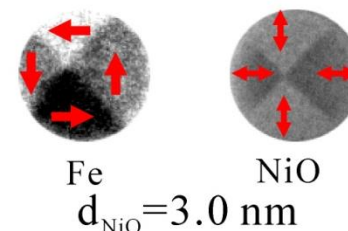
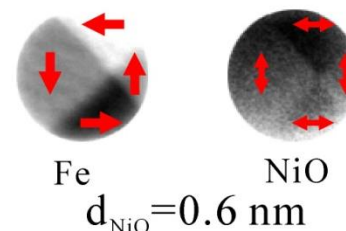
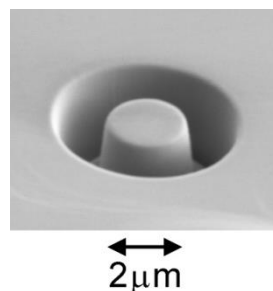
probing in-plane

- + Taking into account the geometry dependence of the Ni XMLD signal  
⇒ Perpendicular coupling of Co and NiO moments at the interface.

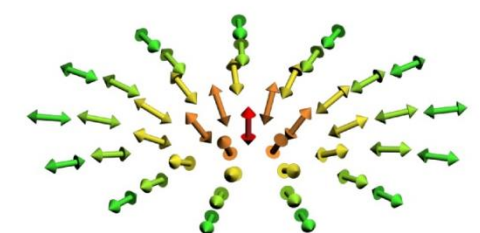
E. Arenholz *et al.*,  
Appl. Phys. Lett. **93**, 162506 (2008)

# MAGNETIC VORTICES

- + First direct observation of vortex state in antiferromagnetic CoO and NiO disks in Fe/CoO and Fe/NiO bilayers using XMCD and XMLD.
- + Two types of AFM vortices:
  - conventional curling vortex as in ferromagnets
  - divergent vortex, forbidden in ferromagnets
  - thickness dependence of magnetic interface coupling



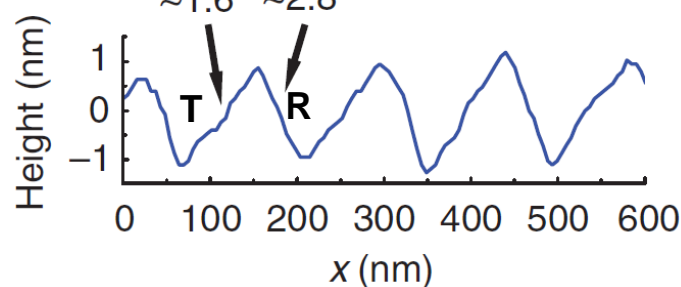
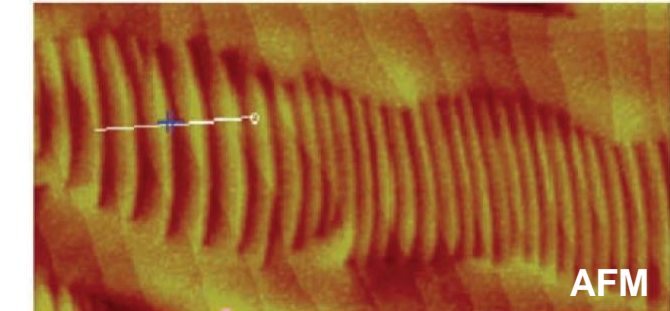
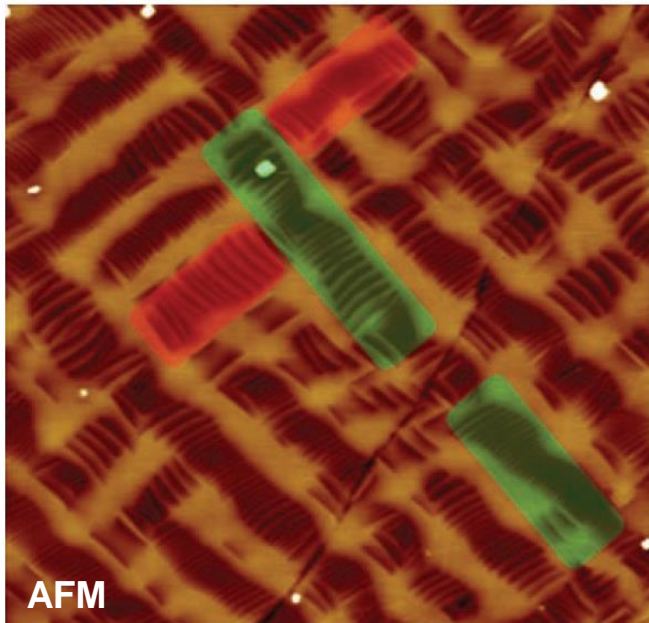
conventional curling vortex



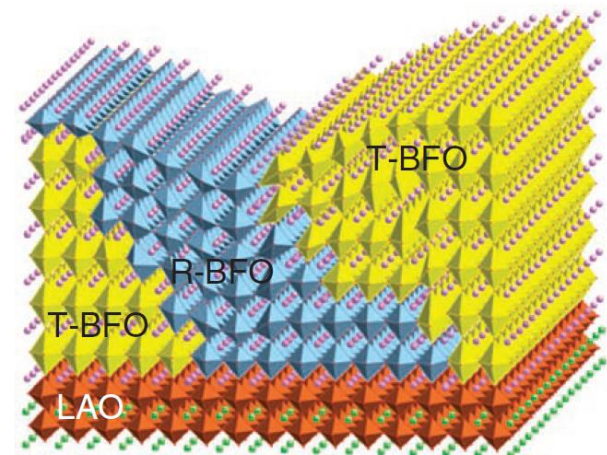
divergent vortex

J. Wu et al.,  
Nature Phys. 7, 303 (2011)

# NANOSCALE MAGNETIC PHASES

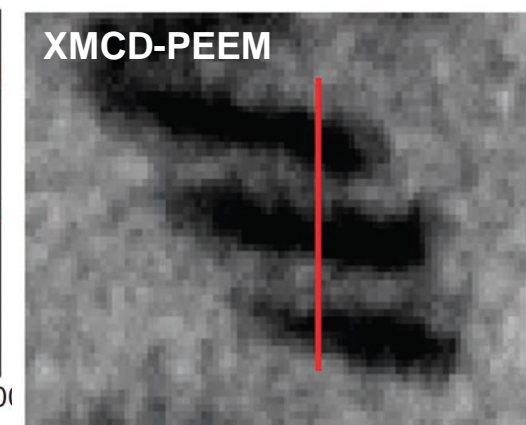
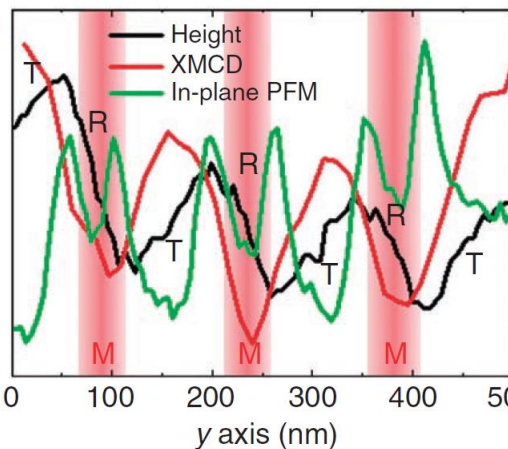
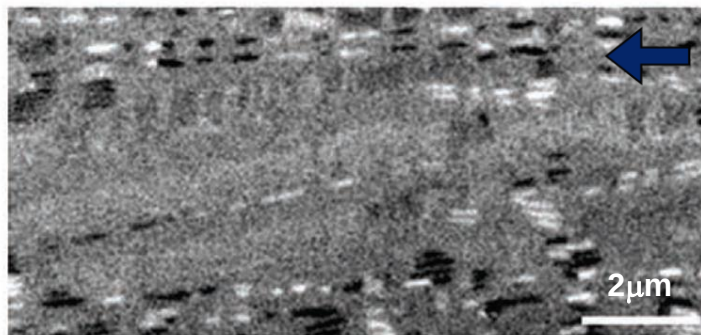
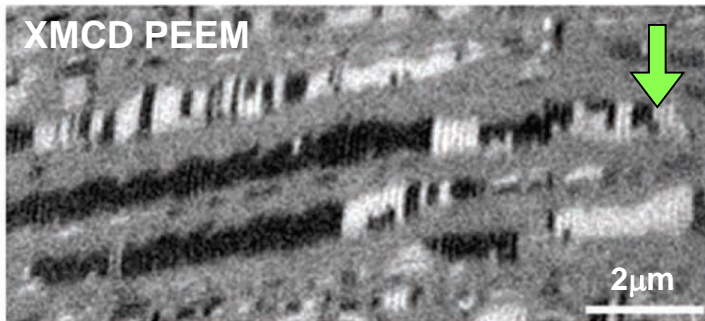
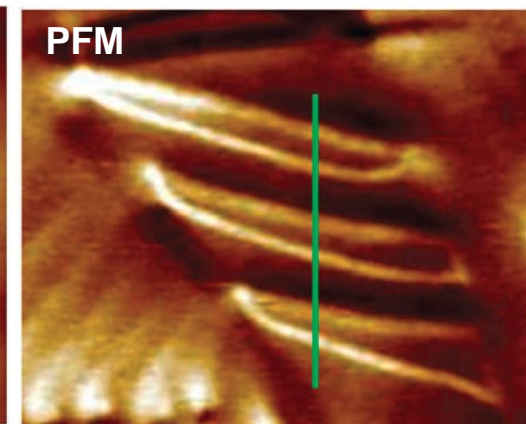
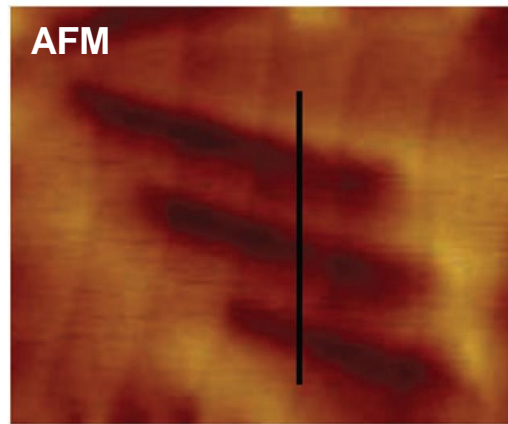
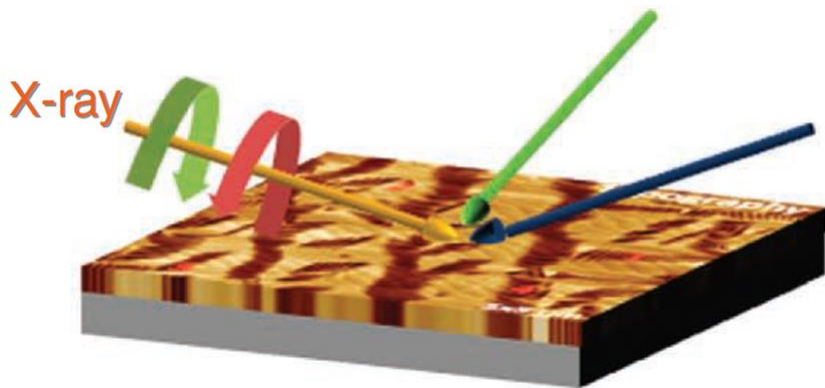


- +  $\text{BiFeO}_3$  – multiferroic = ferroelectric + antiferromagnetic
- + Compressive strain on rhombohedral phase (R-phase) induced by substrate
- ⇒ tetragonal-like phase (T-phase)
- + Partial relaxation of epitaxial strain
- ⇒ Formation of a nanoscale mixture of T- and R-phases



Q. He et al., Nature Comm. 2, 225 (2011)

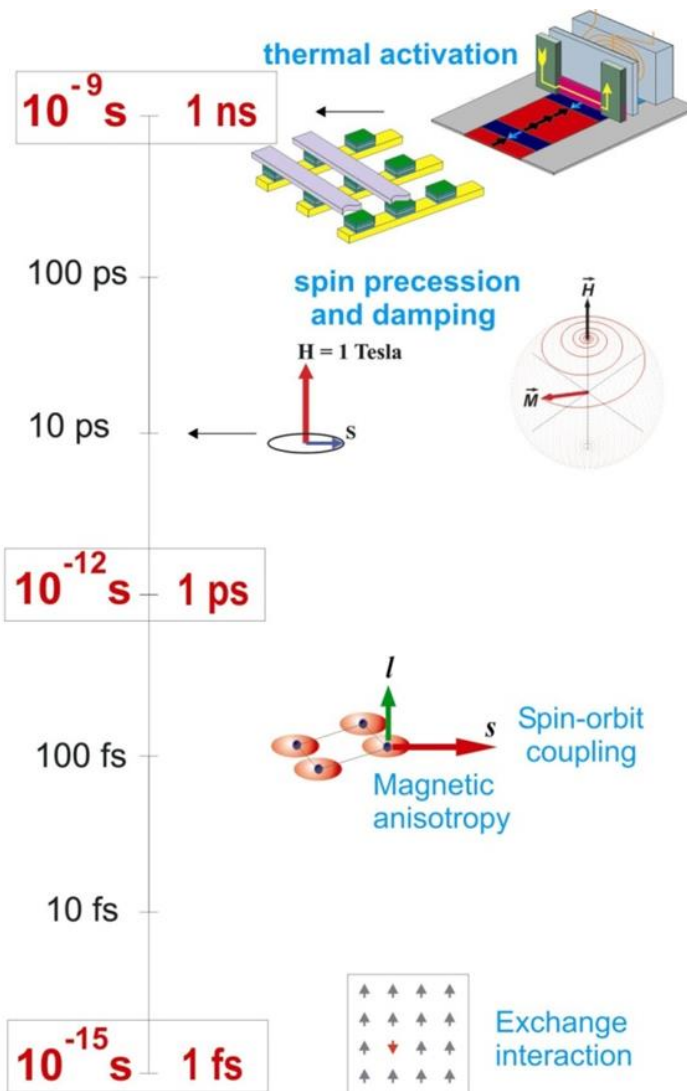
# NANOSCALE MAGNETIC PHASES



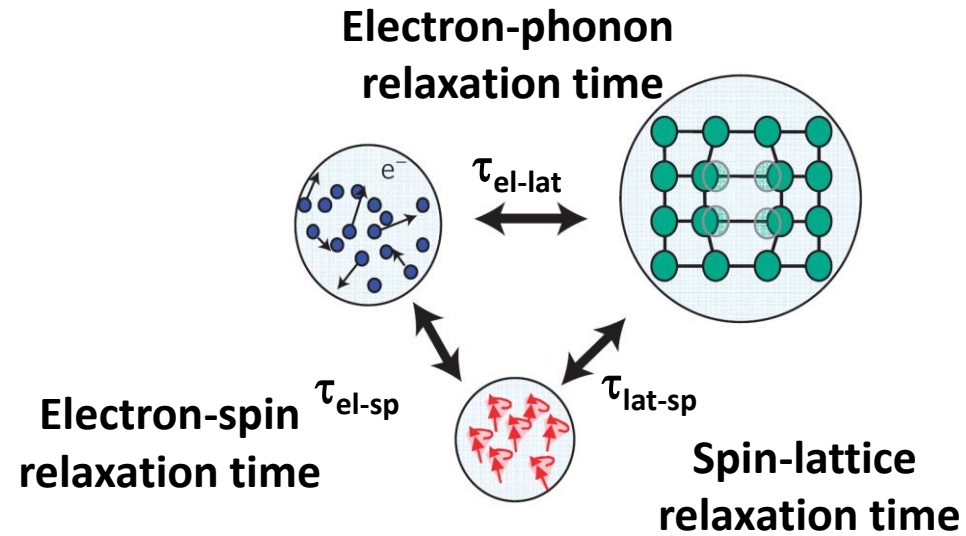
• Highly distorted R-phase is the source of enhanced magnetic moment in the XMCD image.

Q. He et al.,  
Nature Comm. 2, 225 (2011)

# ULTRAFAST MAGNETISM

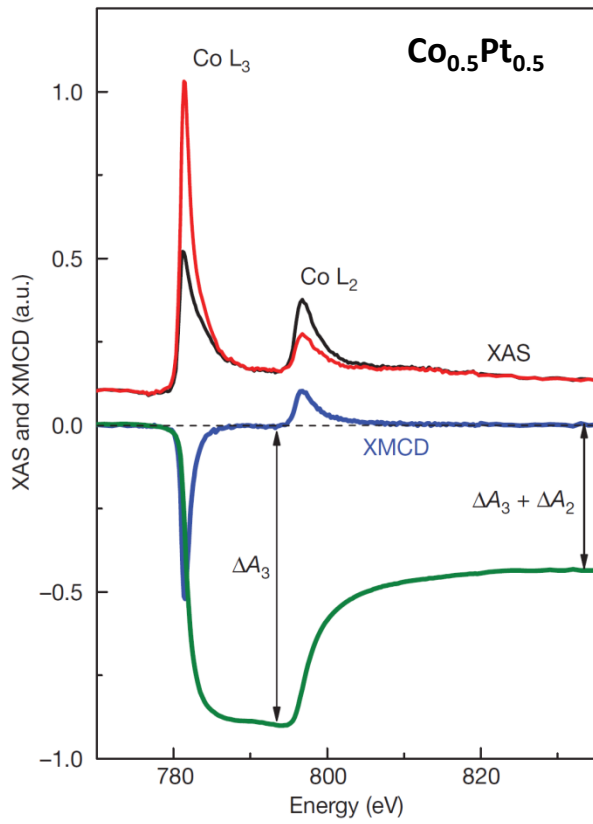


- + Energy reservoirs in a ferromagnetic metal
- + Deposition of energy in one reservoir
- Non-equilibrium distribution and subsequent relaxation through energy and angular momentum exchange



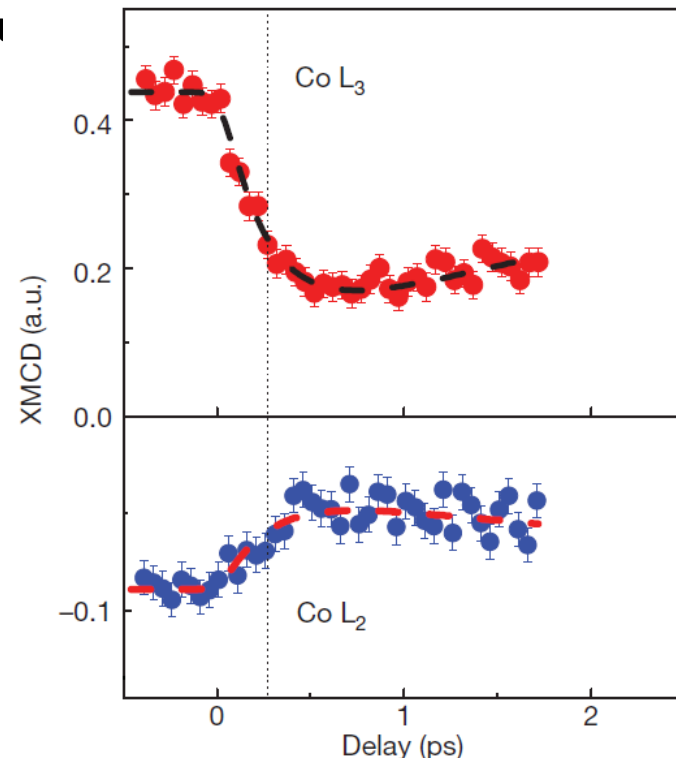
J. Stöhr, H.C. Siegmann,  
Magnetism (Springer)

# ULTRAFAST DYNAMICS OF SPIN AND ORBITAL MOMENTS

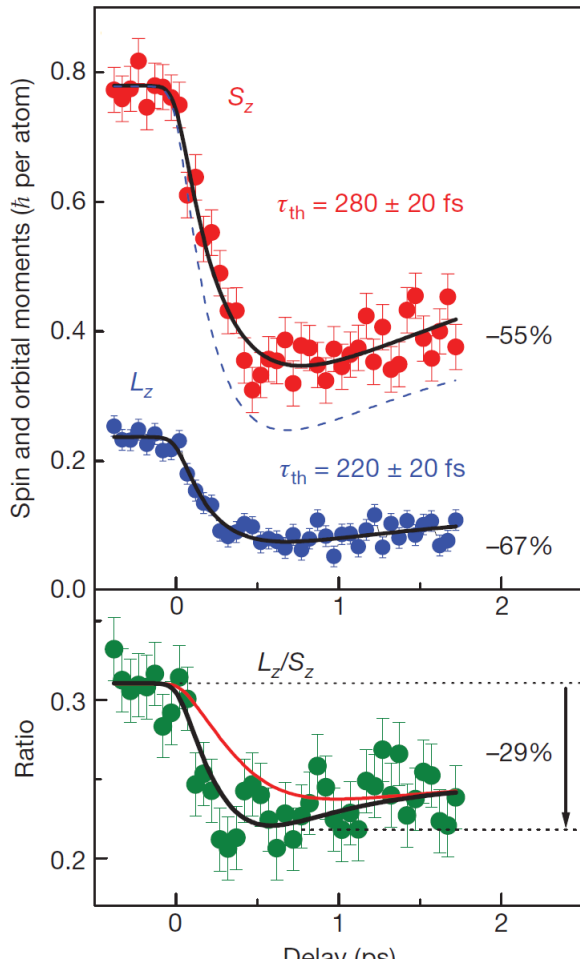


C. Boeglin, et al.,  
Nature **465**, 458 (2010)

- + Orbital ( $L$ ) and spin ( $S$ ) magnetic moments can change with total angular momentum is conserved.
- + Efficient transfer between  $L$  and  $S$  through spin–orbit interaction in solids
- + Transfer between  $L$  and  $S$  occurs on fs timescales.
  
- +  $\text{Co}_{0.5}\text{Pt}_{0.5}$  with perpendicular magnetic anisotropy
- + 60 fs optical laser pulses change magnetization
- + Dynamics probed with XMCD using 120fs x-ray pulses
  
- + Linear relation connects  $\text{Co } L_3$  and  $L_2$  XMCD with  $L_z$  and  $S_z$  using sum rules

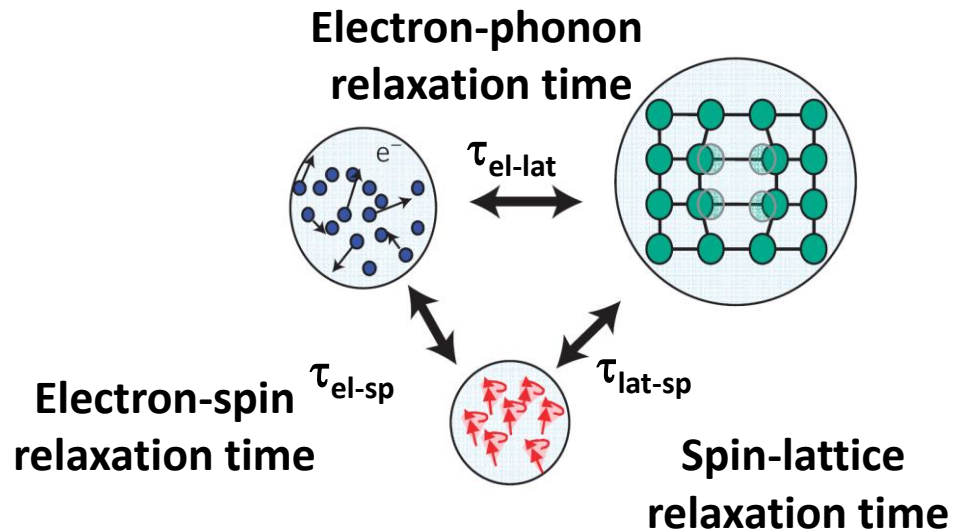


# ULTRAFAST DYNAMICS OF SPIN AND ORBITAL MOMENTS

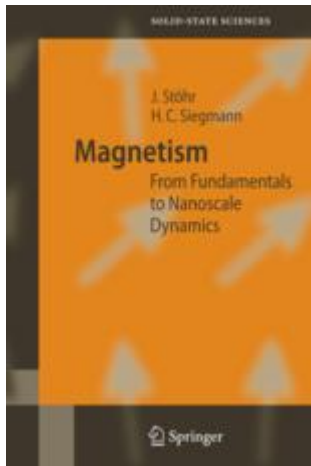


C. Boeglin, et al.,  
Nature **465**, 458 (2010)

- + Thermalization: Faster decrease of orbital moment
- + Theory: Orbital magnetic moment strongly correlated with magnetocrystalline anisotropy
- + Reduction in orbital moment  
 $\Leftrightarrow$  Reduction in magnetocrystalline anisotropy
- + Typically observed at elevated temperatures in static measurements as well



# REFERENCES AND FURTHER READING



**J. Stöhr, H.C. Siegmann**  
**Magnetism— From Fundamentals to Nanoscale Dynamics**  
**Springer**