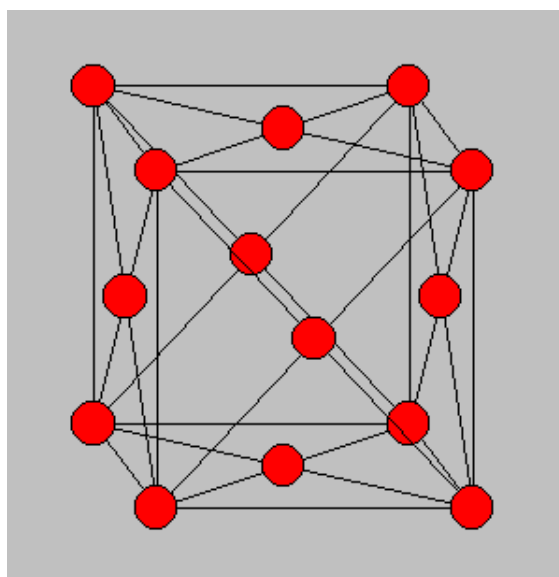


## Bandstrukturmethoden Demo 2: Nickel-Kristall (fcc)

Bandstrukturrechnung mittels FP-LAPW WIEN2k-02

- Kristallstruktur von metall. Ni → fcc
- Input für das WIEN-Programm
- Vorbereitung der Bandstrukturrechnung  
(mit weiteren wichtigen Parametern).
- Die selbstkonsistente Bandstrukturrechnung
- Ergebnisse
- Variation von RKMAX = Größe der LAPW-Basis



**WIEN2k: An Augmented Plane Wave Plus Orbitals Program for Calculating Crystal Properties**

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J. Luitz,  
Inst. of Physical and Theoretical Chemistry,  
Vienna University of Technology

## Elektronenstruktur (Z=28):

**1s<sup>2</sup> 2s<sup>2</sup> 2p<sup>6</sup> 3s<sup>2</sup> 3p<sup>6</sup> 3d<sup>8</sup> 4s<sup>2</sup>**

oder

**1s<sup>2</sup> 2s<sup>2</sup> 2p<sup>6</sup> 3s<sup>2</sup> 3p<sup>6</sup> 3d<sup>8</sup> 4s<sup>2</sup>**

- Input für das WIEN-Programm: Ni\_demo.**struct**

FCC Ni

F LATTICE,NONEQUIV.ATOMS: 1

MODE OF CALC=NREL

6.7000 6.7000 6.7000

ATOM= 1: X=0.00000000 Y=0.00000000 Z=0.00000000

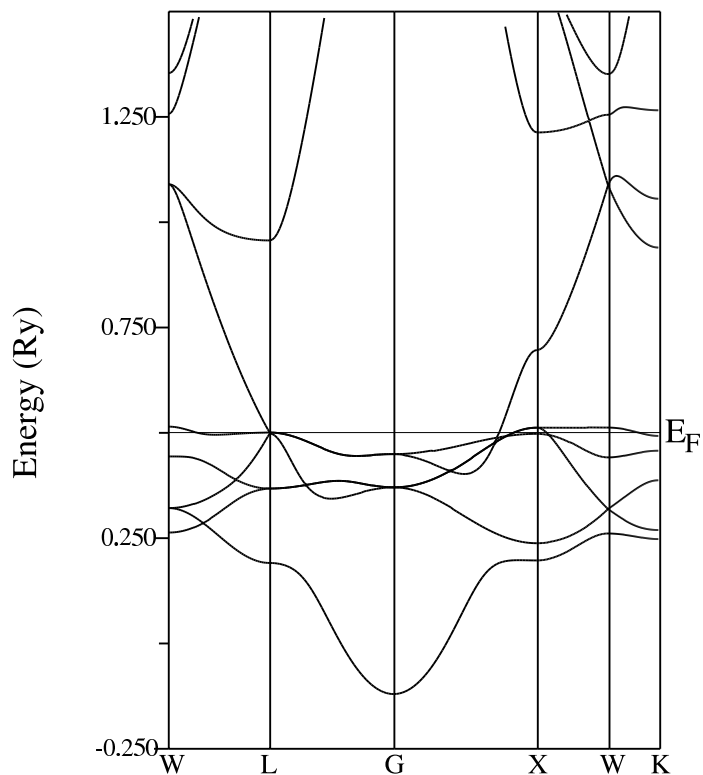
MULT= 1 ISPLIT= 2

Ni NPT= 381 R0=.000050000 RMT=2.300000000 Z:28.0

LOCAL ROT MATRIX: 1.0000000 0.0000000 0.0000000  
0.0000000 1.0000000 0.0000000  
0.0000000 0.0000000 1.0000000

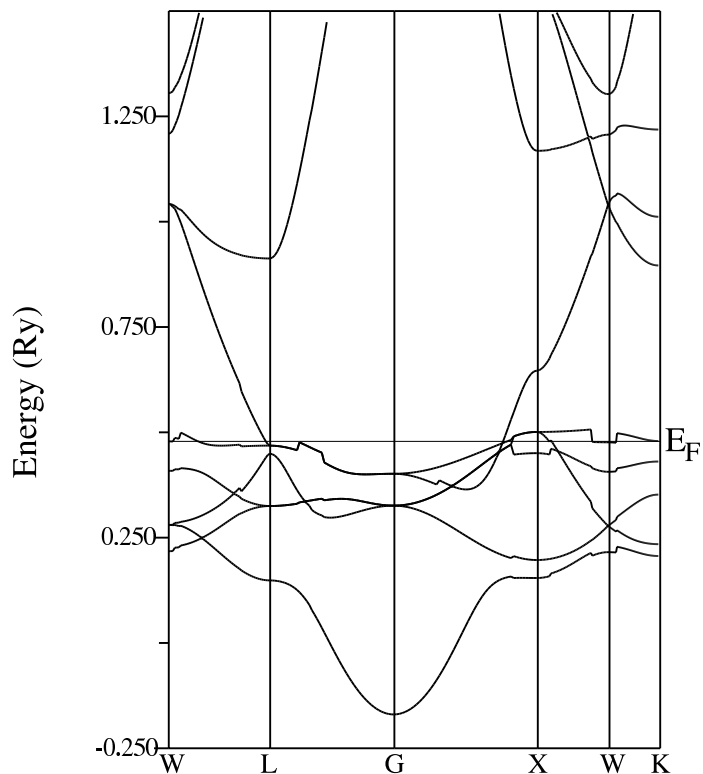
0 NUMBER OF SYMMETRY OPERATIONS

Ni\_demo atom 0 size 0.20



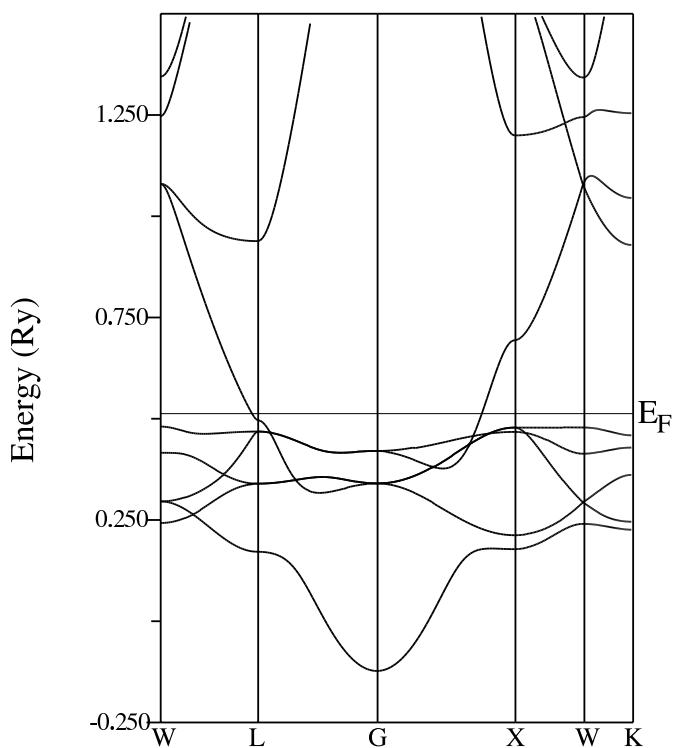
Nickel-Bandstruktur: **RKMAX = 7.0**

Ni\_demo atom 0 size 0.20



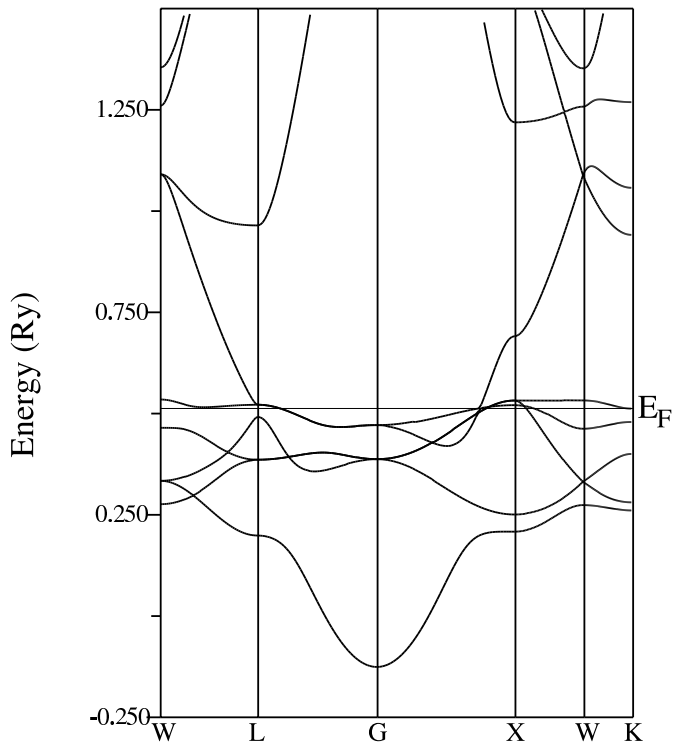
Nickel-Bandstruktur: **RKMAX = 5.0**

Ni\_demo atom 0 size 0.20



Nickel-Bandstruktur: spin-polarisiert UP

Ni\_demo atom 0 size 0.20



Nickel-Bandstruktur: spin-polarisiert DOWN

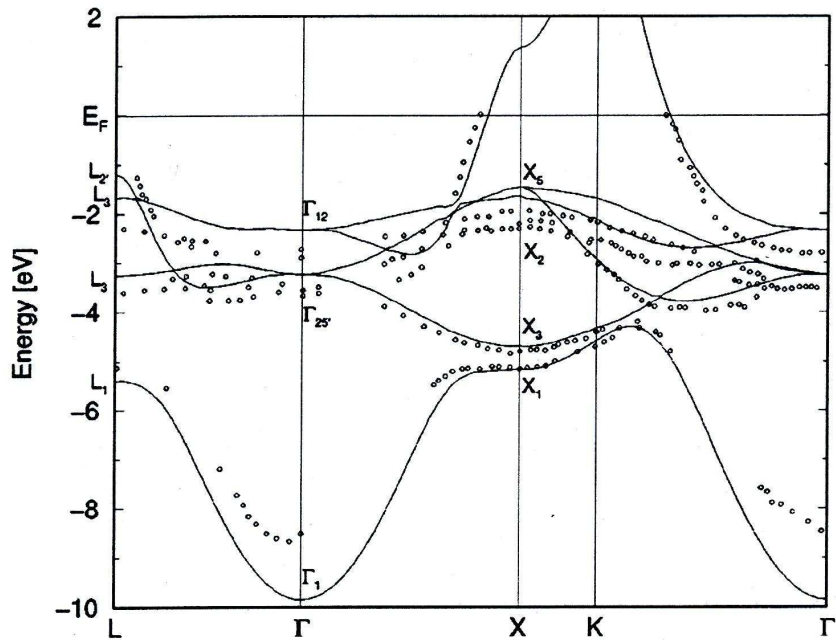


Figure 1: Bulk copper DFT-LDA band structure (solid line), compared with photoemission data (points).

Theory: Marini et al., Phys. Rev. B **64**, 195125 (2001),  
experiment: Courths and Hüfner, Phys. Rep. **112**, 53 (1984).

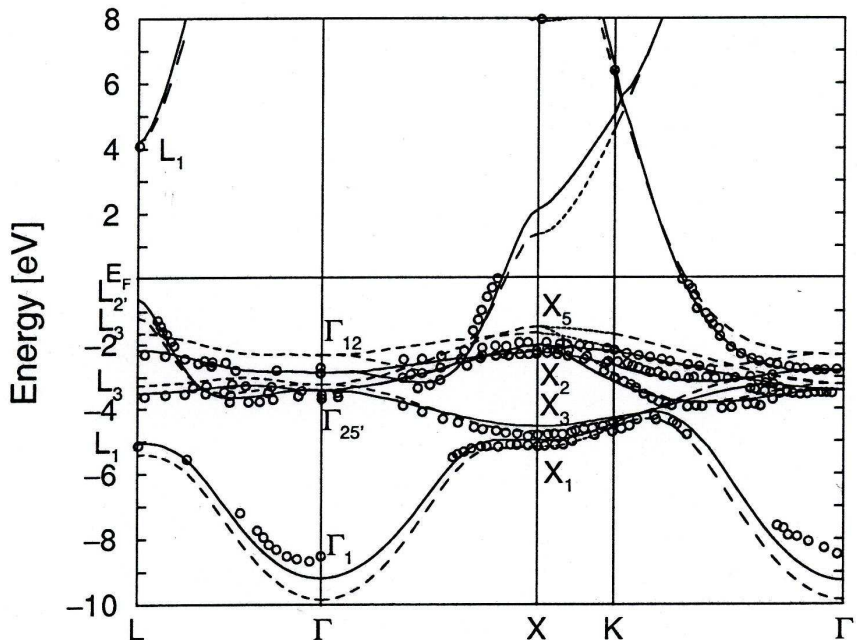


Figure 2: Solid line: GW results for the bulk copper band structure, compared with the DFT-LDA results (dashed line), and with experimental photoemission data.

Theory: Marini et al., Phys. Rev. Lett. **88**, 016403 (2002),  
experiment: Courths and Hüfner, Phys. Rep. **112**, 53 (1984).