

Modern Methods in Heterogeneous Catalysis Research: Theory and Experiment



Low Energy Electron Diffraction - LEED

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Literature:

G. Ertl, J. Küppers, Low Energy Electrons and Surface Chemistry, VCH, Weinheim (1985).

M. <u>Henzler</u>, W. <u>Göpel</u>, Oberflächenphysik des Festkörpers, Teubner, Stuttgart (1991).

M.A. Van Hove, W.H. Weinberg, C.-M. Chan, Low-Energy Electron Diffraction, Experiment, Theory and Surface Structure Determination, Springer Series in Surface Sciences G. Ertl, R. Gomer eds., Springer, Berlin (1986).

M. Horn-von Hoegen, Zeitschrift für Kristallographie 214 (1999) 1-75.

1. Introduction, General Surface science, UHV, p~10⁻¹⁰ mbar

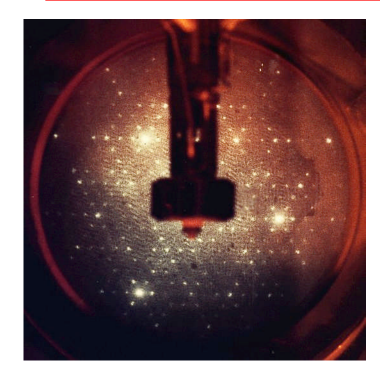
De Broglie wavelength: $\lambda = h/(mv)$

For electrons: $\lambda = \sqrt{150 / E_0}$ E_0 in eV, λ in Å.

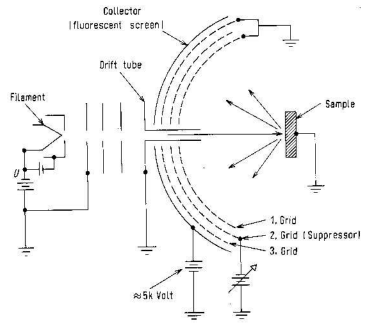
For 100 eV-electrons: $\lambda(100) = 1.22 \text{ Å}$ (low energy)

corresponds to atomic dimensions,

similar to XRD



Si(111)-(7x7)

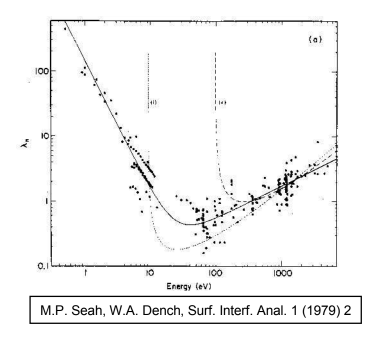


LEED display system

Ertl/Küppers fig. 9.7, p. 210

LEED is surface sensitive

Low energy electrons interact strongly with matter: electron mean free path λ_e is small. Only e^- scattered from near surface can leave the surface, surface sensitive



The observation of a LEED pattern does not guarantee that the whole surface is ordered!

Coherence of e^- -beam limited by ΔE and beam divergence. Coherence length = diameter of coherently scattering area.

The coherence length of a standard LEED optics is only 10 – 20 nm!

1st approximation: Scattering from 2-D lattice.

Analogy to optical grating.

Constructive interference: Enhancement of intensity only in certain directions:

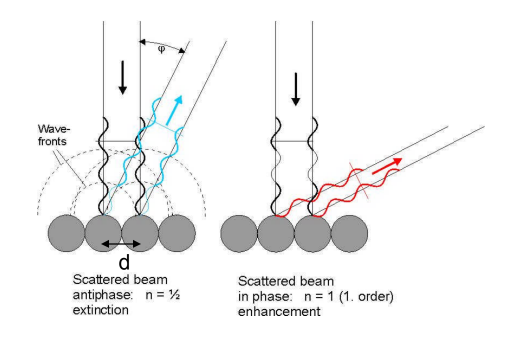
$$n \lambda = d \sin \varphi$$

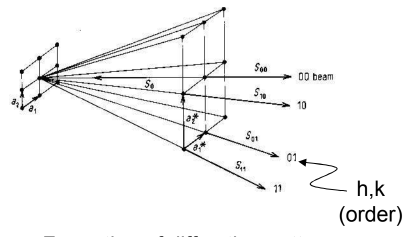
For 2D arrangement (plane lattice): scattering conditions have to be fulfilled in both directions

Note:

If the lattice constant(s) a_1 (a_2) <u>in</u>crease, the scattering angle for the beam h (k) <u>de</u>creases.

This is the reason for the reciprocity of the real and the s.c. reciprocal lattice.





Formation of diffraction pattern

Ertl/Küppers fig. 9.12, p. 217

Useful: Introduction of reciprocal lattice

Real lattice vectors $\mathbf{a_1}$, $\mathbf{a_2}$ Reciprocal lattice vectors $\mathbf{a_1}^*$, $\mathbf{a_2}^*$

Definitions: a_1^* perpendicular to a_2^* perpendicular to a_1^*

 $a_1^* = 1/(a_1 \sin \gamma)$ $a_2^* = 1/(a_2 \sin \gamma)$ γ angle between a_1 and a_2

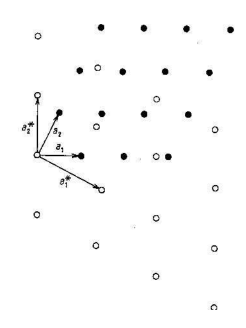
Constructive interference for:

$$\mathbf{a_1} (\mathbf{s} - \mathbf{s_0}) = \mathbf{h} \lambda$$

 $\mathbf{a_2} (\mathbf{s} - \mathbf{s_0}) = \mathbf{k} \lambda$

(Laue conditions for 2 dimensions)

Real 2D system: 3rd Laue condition always fulfilled.



Example

Ertl/Küppers fig. 9.11, p 216

It follows for the direction of beams:

$$1/\lambda (\mathbf{s} - \mathbf{s_0}) = 1/\lambda \Delta \mathbf{s} = \mathbf{h} \ \mathbf{a_1}^* + \mathbf{k} \ \mathbf{a_2}^* = \mathbf{g}$$

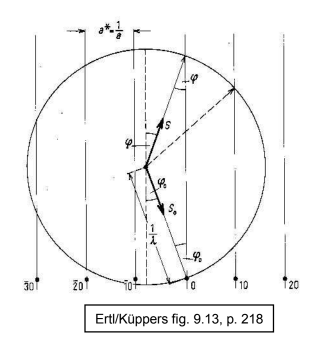
 $\mathbf{g} = \text{reciprocal lattice vector}$

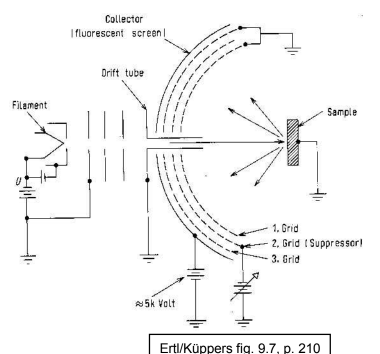
Ewald sphere construction

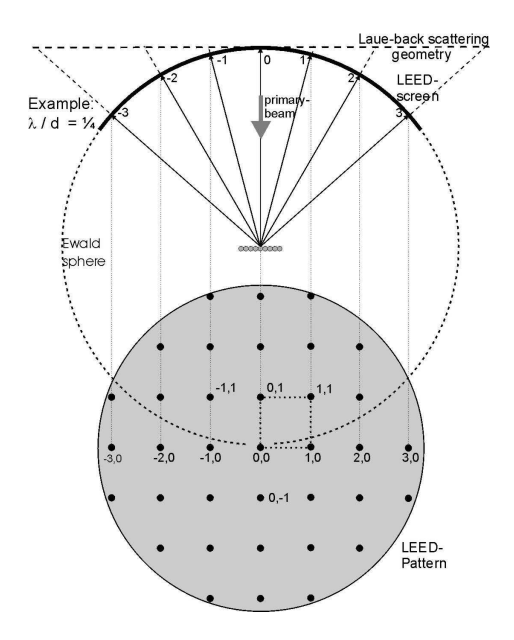
- plot reciprocal lattice (rods)
- plot direction of incident beam (s₀) towards (00) spot
- go $1/\lambda$ along this direction
- make circle (sphere) with radius 1/λ
- direction from circle (sphere) center towards cut with reciprocal lattice rods gives direction of all possible diffraction spots (hk)

Usual arrangement:

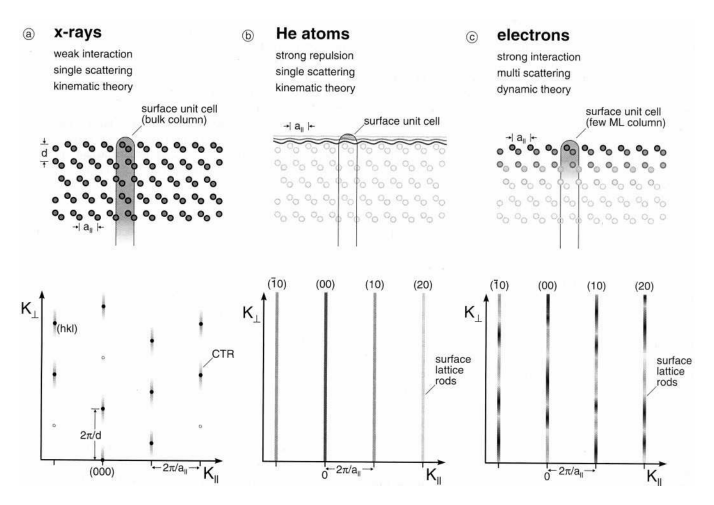
Normal incidence, symmetrical diffraction pattern







Expected diffraction pattern for (001) surface, e.g. Pt(001) (unreconstructed), E_0 =313 eV



Surface diffraction with X-rays, He-atoms and electrons. Example: diamond-type (111) surface like C, Si, Ge. The darkness of rec. latt. spots and rods symbolizes diffraction intensity

Horn-von Hoegen, fig. 2.1

LEED:

2. Simple

Kinematic theory (single scattering)
Size, shape and symmetry of surface unit cell,
Superstructures
Domains
only if long-range ordered

No information about atomic arrangement within the unit cell

3. Less simple

Kinematic theory

Deviations from long-range order:

Spot width \rightarrow domain size

Background intensity → point defect concentration

Spot splitting \rightarrow atomic steps

4. Difficult

Dynamic theory (multiple scattering)

Spot intensities $I(E_0)$ or I-V curces \rightarrow structure within unit cell

2. LEED - simple

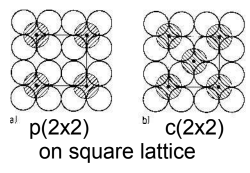
Superstructures result from:

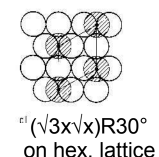
Reconstruction = rearrangement of surface atoms on clean surfaces Ordered adsorption

Structure examples

Overlayer structures

Ertl/Küppers fig. 9.2, p.204





Superstructure nomenclature

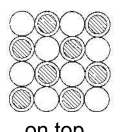
Wood: Simplest in most cases p or $c(n\times m)R\vartheta^{\circ}$ unit cell vector lengths $b_1 = n a_1$ $b_2 = m a_2$ rotation ∂ p=primitive, c=centered Matrix notation (Park and Madden) more general

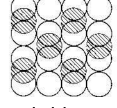
$$\begin{array}{lll}
 m_{11} & m_{12} & \mathbf{b_1} = m_{11} \mathbf{a_1} + m_{12} \mathbf{a_2} \\
 m_{12} & m_{22} & \mathbf{b_2} = m_{12} \mathbf{a_1} + m_{22} \mathbf{a_2}
 \end{array}$$

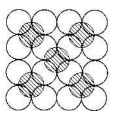
Wood	(2×2) [ϑ=0 is omitted]	(√3×√3)R30°
Matrix	2 0 0 2	1 1 2 -1

Three possible arrangements yielding c(2x2) structures. Note: different symmetry!

Ertl/Küppers fig. 9.6, p.208



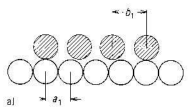




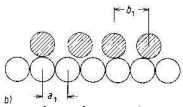
on top

bridge

4-fold hollow

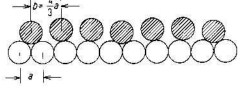


coincidence latt. commensurate

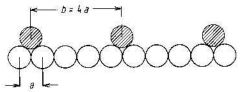


incoherent, incommensurate

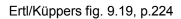
Ertl/Küppers fig. 9.3, p.205

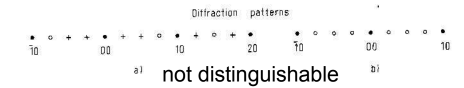


coincidence latt.



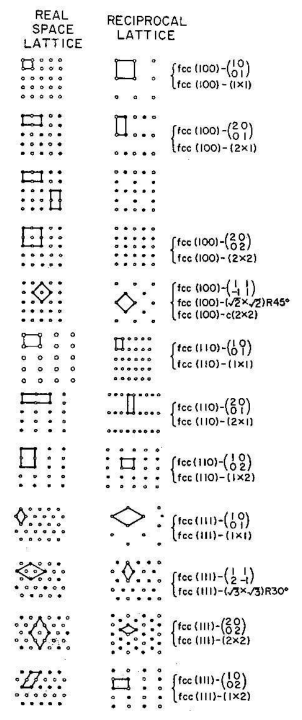
simple overlayer struct.





Real and reciprocal space lattices

Van Hove et al. fig. 3.5, p.55



Superstructures, example 1

GaAs(001) clean, different preparations

As(31)/Ga(55)
Auger peak height ratios:

c(8x2) 1.74 (4x6) 1.77 c(6x4) 1.92 (1x6) 2.12 c(2x8) 2.25 c(4x4) 2.7

Information from patterns:

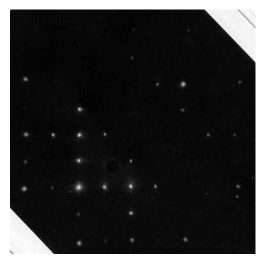
- symmetry of unit cell
- size and shape of surface unit cell
- sharpness of spots
 - \rightarrow domain size
- background intensity
 - → concentration of point defects

c(8x2)(4x6)66eV 110eV c(6x4)(1x6)36eV 35eV c(2x8)c(4x4) 80eV

Drathen, Ranke, Jacobi, 1978

Superstructures, example 2

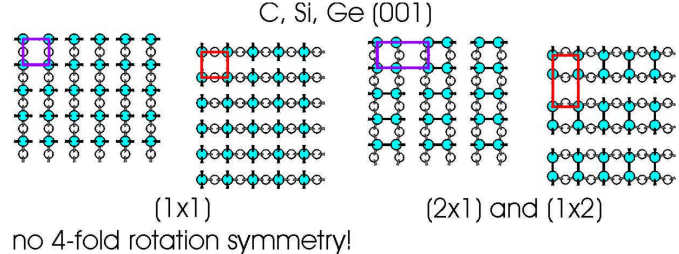
Si(001) clean

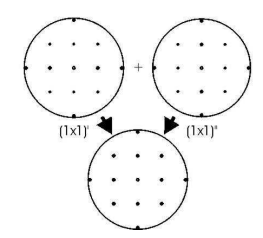


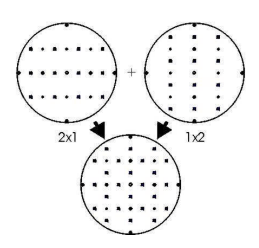
no 2x2 structure!
central spots missing

→ two-domain 2x1

Wasserfall, Ranke, 1994







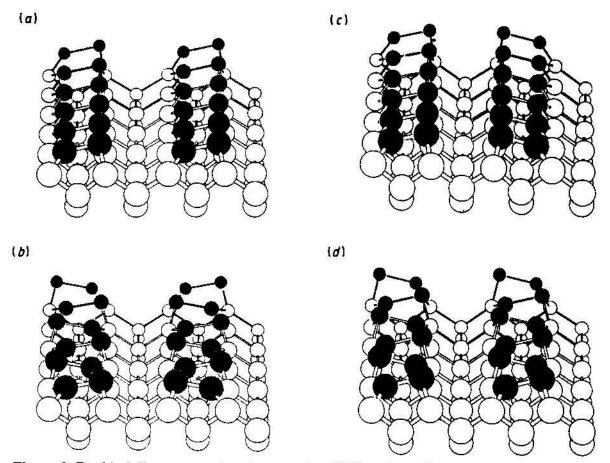
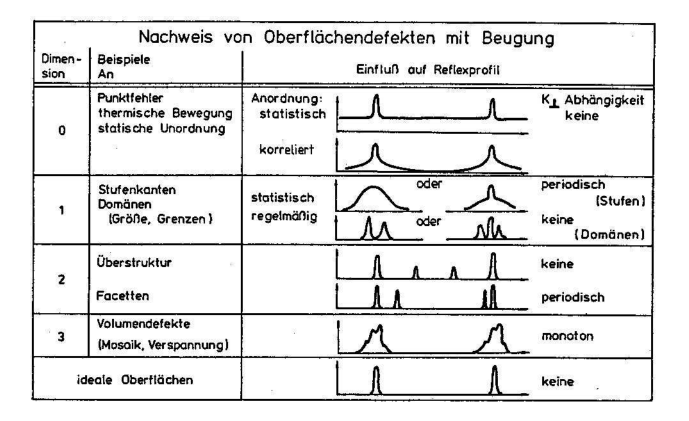


Figure 3. Buckled dimer reconstructions on the (001) surface of germanium: (a) $b(2 \times 1)$; (b) $c(4 \times 2)$; (c) $p(4 \times 1)$; (d) $p(2 \times 2)$.

Payne et al. J. Phys.: Cond. Matter 1 (1989) SB63

3. LEED - less simple

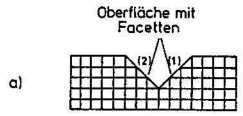
Information from spot shape (profile), background, E_0 -dependence (k_1 -dependence)

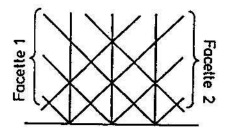


Henzler, Göpel Abb. 3.8.10, p.176

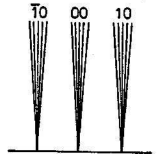
Facets and mosaic

Henzler, Göpel Abb. 3.8.4, p.167



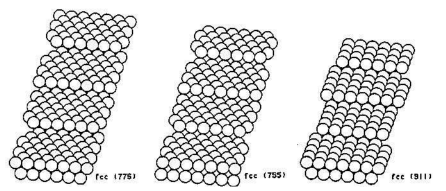


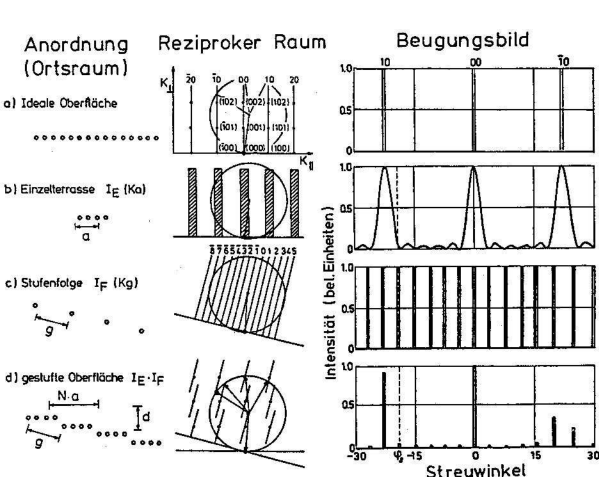




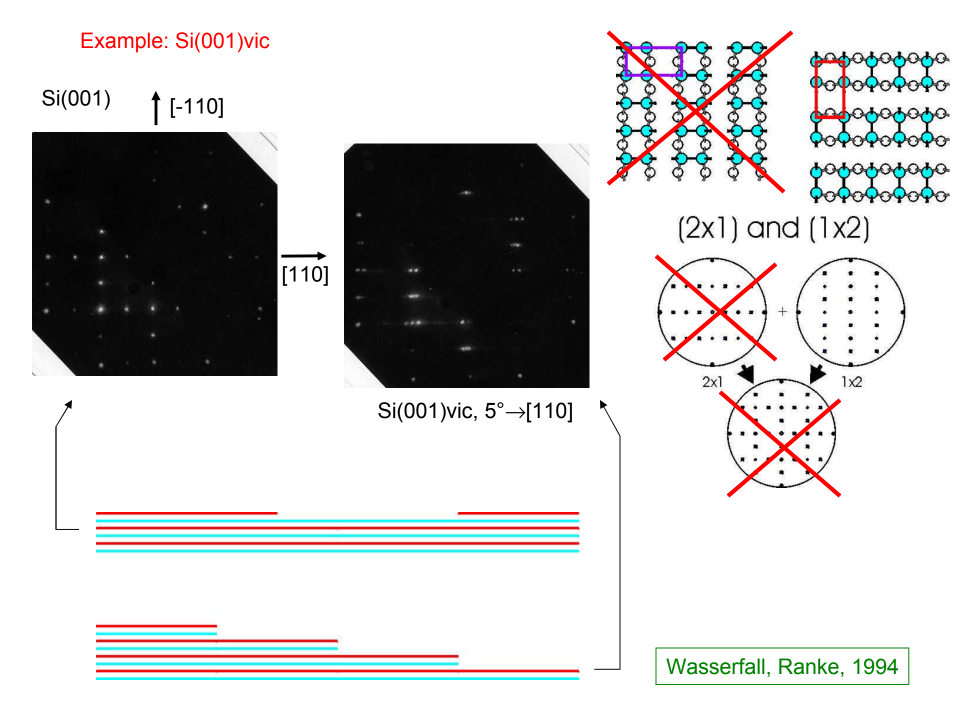
Regular atomic steps

Van Hove et al., fig. 3.6, p.58





Henzler, Göpel, fig. 3.8.3, p.165



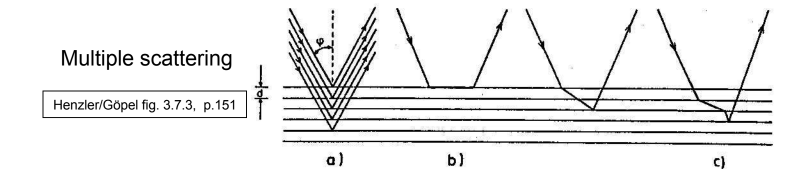
4. LEED - difficult

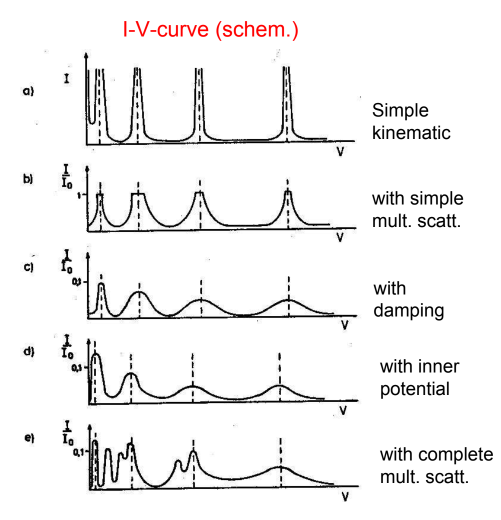
Spot intensities contain information on structure within the unit cell

$$I \sim |F|^2 \cdot |G|^2$$

|G|² = structure factor or lattice factor
contains shape and arrangement of repeat units (unit cells)
yields reciprocal lattice
determines location and shape of spots,
kinematic theory

|F|² = structure factor or form factor contains contribution from all atoms within the repeat unit, includes multiple scattering, in-depth attenuation, dynamic theory





Henzler/Göpel, fig. 3.7.4, p.152

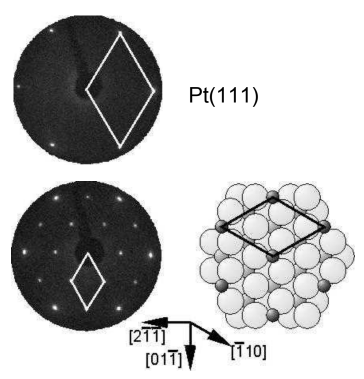
Dynamic LEED analysis: No direct deduction of structure from I-V-curves:

Guess structure model calculate I-V-curves compare with measured curves modify model check if improval if yes: proceed modifying in this direction if no: modify in another direction or guess new model

Disadvantage:
Only for ordered structures
Much computer time

But:

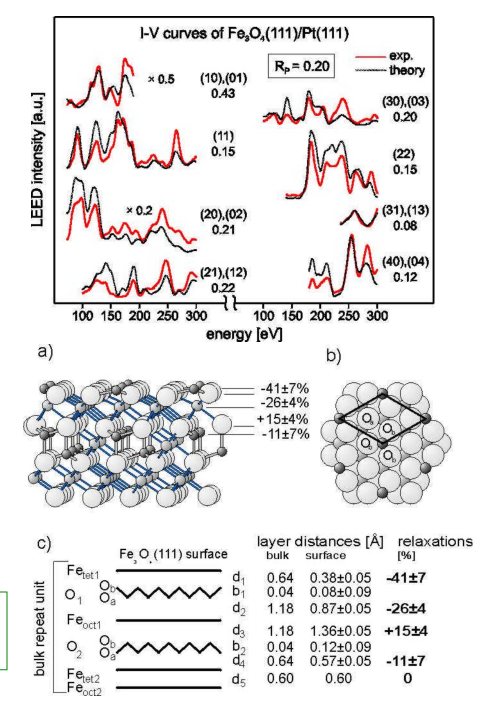
One of very few methods for structure analysis of first few atomic layers (~1 nm)



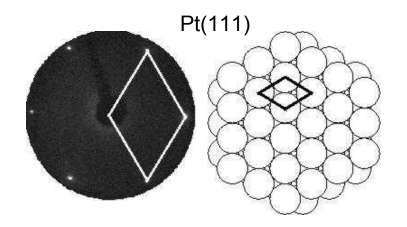
Fe₃O₄(111), (inverse spinel) 10 nm thick on Pt(111)

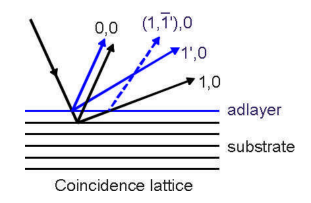
LEED-I-V analysis is one of very few reliable surface structure analysis methods!

Michael Ritter, Werner Weiss Guido Ketteler

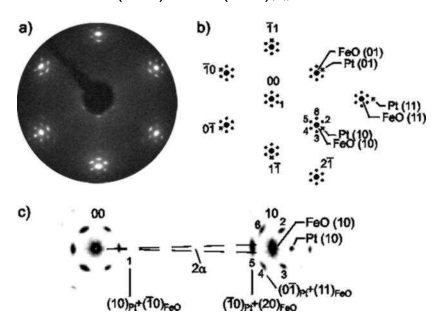


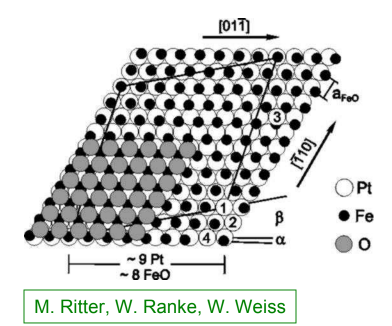
FeO/Pt(111), satellite pattern: multiple scattering, kinematic



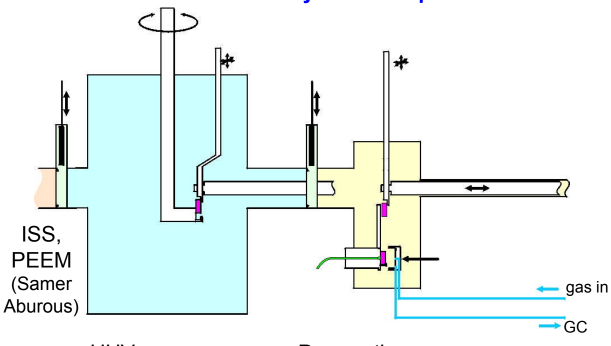


0.9 ML FeO(111) on Pt(111), "structure 1"

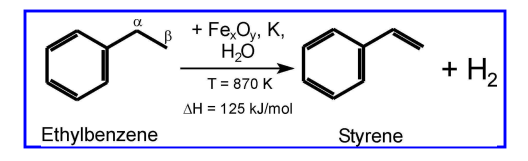


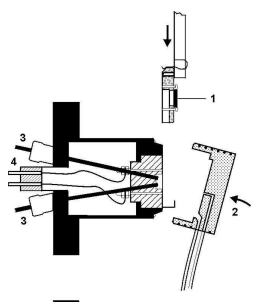


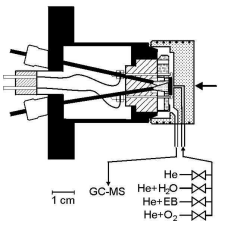
5. LEED in model catalysis - example



UHV **LEED**, AES, TDS p= 10⁻⁶ to 10⁻¹⁰ mbar Preparation reactor p=1000 to 10⁻⁶ mbar

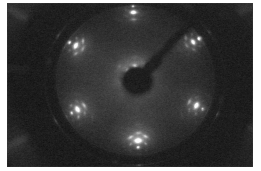




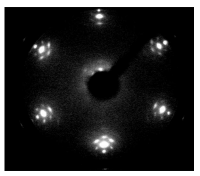


Manfred Swoboda Christian Kuhrs Werner Weiss

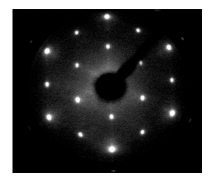
Distinguish different Fe-O-phases



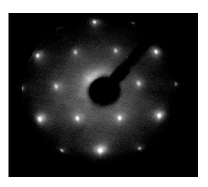
as measured



contrast enhanced



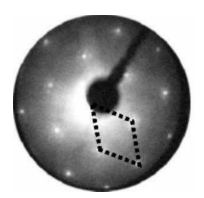
 $Fe_3O_4(111)$



 α -Fe₂O₃(0001)

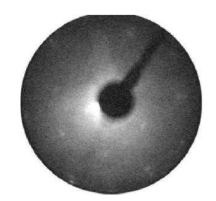
Change of order and phase during

reaction



FeO(111)/Pt(111), 1 ML

Starting surface: α -Fe₂O₃(0001) (hematite), defective



After reaction
- no long-range order

- strong C peak in AES



After mild TPO (thermal programmed oxidation)

- reordered
- no longer hematite but Fe₃O₄(111) (magnetite

Osama Shekhah



Modern Methods in Heterogeneous Catalysis Research: Theory and Experiment



6. Conclusions

For qualitative information on surface structure very simple (display LEED)

- Order
- Periodicity
- Symmetry

For quantitative information on deviations from ideal order (SPA-LEED)

- Domain size
- Antiphase domains
- atomic steps

For quantitative analysis of surface structure (dynamic I-V-curve analysis)

- Precise atomic arrangements
- Relaxations
- Reconstructions