THE STRUCTURE OF ATOMIC NITROGEN ADSORBED ON Fe(100)

R. IMBIHL, R.J. BEHM and G. ERTL

and

W. MORITZ
Institut für Kristallographie und Mineralogie, Universität München, D-8000 München 2, Fed. Rep. of Germany

Received 16 July 1982; accepted for publication 2 September 1982

Nitrogen atoms adsorbed on a Fe(100) surface cause the formation of an ordered c(2\times2) overlayer with coverage 0.5. A structure analysis was performed by comparing experimental LEED I-V spectra with the results of multiple scattering model calculations. The N atoms were found to occupy fourfold hollow sites, with their plane 0.27 Å above the plane of the surface Fe atoms. In addition, nitrogen adsorption causes an expansion of the two topmost Fe layers by 10% ( = 0.14 Å). The minimum r-factor for this structure analysis is about 0.2 for a total of 16 beams. The resulting atomic arrangement is similar to that in the (002) plane of bulk Fe₄N, thus supporting the view of a "surface nitride" and providing a consistent picture of the structural and bonding properties of this surface phase.

1. Introduction

Catalytic synthesis of ammonia over iron catalyst proceeds through atomically adsorbed nitrogen (whose formation is the rate-limiting step) [1], and therefore knowledge on the surface structure of this species is of importance for understanding the microscopic reaction steps. Previous LEED investigations [2–4] revealed that on Fe(100) a simple c(2 × 2) structure is formed, while with Fe(110) or (111) surfaces a series of complicated diffraction patterns was observed which strongly suggested the occurrence of surface reconstruction.

It was proposed [3,4] that these surface phases (which were called “surface nitrides”) exhibit structures which closely resemble those of the bulk nitride Fe₄N, even if the latter cannot be formed for thermodynamic reasons. According to this concept the Fe(100)-c(2 × 2)-N structure should be similar to the (002) plane of Fe₄N, while with Fe(110) and (111) no low-index planes of...
Fe₄N [5] are compatible without pronounced displacements of iron atoms, thus causing reconstruction.

The present work concerns a structural analysis of the Fe(100)-c(2 × 2)-N surface phase on the basis of LEED intensity/voltage (I–V) data which will confirm the proposed picture.

2. Experimental

The experiments were performed with a standard UHV system (base pressure below 2 × 10⁻¹⁰ Torr) which was equipped with a four-grid LEED optics serving also as retarding field analyser for AES. I–V spectra were recorded with a moveable Faraday cup (aperture 14°). A system of Helmholtz coils controlled by a magnetic field probe (Förster) was used in order to suppress magnetic stray fields to < 1 mOe.

The sample was cut from a single crystal rod (purity 99.99%, Metals Research) and oriented to within 1/2°. The surface impurities (mainly C, O and S) were removed by prolonged oxidation and sputter/annealing cycles as described earlier [2]. The cleanliness was checked by AES. During the LEED measurements the peak ratios of the C(270 eV) and O(512 eV) signals over the Fe(650 eV) signal could always be kept below 0.05, indicating impurity concentrations of less than a few percent of a monolayer.

The ordered c(2 × 2) overlayer of chemisorbed nitrogen atoms was produced by thermal decomposition of NH₃ at 670 K [6] which needs considerably lower gas exposures as if N₂ is used [2]. This procedure leaves only nitrogen on the surface since hydrogen desorbs at much lower temperature [6]. The onset of bulk-Fe₄N formation occurs, on the other hand, at considerably higher exposure than applied here [7]. Nevertheless part of the nitrogen atoms formed diffuse into the bulk as demonstrated previously by means of isotope exchange experiments [3].

The c(2 × 2) structure was characterised by an Auger intensity ratio \( \gamma_N = N(380 \text{ eV})/\text{Fe(650 eV)} = 0.6. \) The half-order LEED spots were rather sharp and exhibited intensities of about 30% of the substrate lattice spots.

The LEED measurements were performed after brief heating of the sample to 700 K (in order to remove adsorbed residual NH₃, etc.) and cooling down to room temperature. I–V curves were recorded in steps of 1 eV between 40 and 200 eV for a total of seventeen beams, namely 9 integral-order and 8 fractional-order beams at three different angles of incidence (θ = 0, 6 and 11.6°). No correction of the energy scale was made for the difference of the work function of the sample (5.2 eV) and the tungsten filament of the LEED gun (4.5 eV) in view of the general uncertainty (~ 1 eV) of the voltage scale.

Beam profile measurements revealed that at 56 eV only about 50% of the integral intensity of a beam is collected by the Faraday cup with its small
aperture (≈ 1°). Since with increasing energy the diffraction spot diameter decreases (as long as it is not limited by the instrumental function) also an increasing fraction of the integral intensity is recorded. As a consequence the relative intensities are somewhat too high at higher electron energies. It was, however, estimated, that this effect is only of very minor influence on the $r$-factor determination, and therefore no correction was made.

3. Calculation procedures

A Hartree–Fock–Slater selfconsistent potential was used for Fe in analogy with previous work [8–10]. The nitrogen potential was constructed from a superposition of free atom potentials in a bcc unit cell [11]. LEED intensity calculations were performed using the layer-doubling method [12] for multiple scattering between bulk layers, while the small distance between the N layer and the topmost Fe layer required a solution by means of matrix inversion in angular momentum representation within the first two layers. Up to 28 symmetrized beams at normal incidence were involved in the layer-doubling scheme for the models with an adsorption site of fourfold symmetry. A corresponding large number of beams was used for the bridge-site models with p2mm symmetry as well as for oblique incidence. The number of phase shifts was reduced to seven, since the calculation for the first layer with three atoms per unit cell required rather long computing times. It was checked that seven phase shifts were sufficient for the energy range up to 200 eV; calculations with eight phase shifts yielded only minor differences.

The real part of the inner potential was assumed to be independent of energy. Its optimum value found was $V_0 = 9.5$ eV, whereby no correction for the work function difference is taken into account. The imaginary part of the inner potential was set equal to $0.85E^{1/3}$ [eV], which yielded better agreement between theory and experiment than the smaller value of approximately $0.7E^{1/3}$ [eV] as used by Legg et al. [8] in a study of the clean Fe(100) surface. No further efforts were made to optimize this as well as other non-structural parameters. The bulk Debye temperature is $\theta_D = 467$ K [13] and for the surface layer a value of 400 K was assumed in order to allow a slightly larger mean square vibrational amplitude for the topmost atoms. Variation of the surface Debye temperature was of only small effect on the calculated $I$–$V$ spectra, and the results were not significant enough in order to draw any firm conclusion on the actual mean square vibrational amplitudes.

4. Results

Before starting the analysis of the overlayer structure, data for the clean Fe(100) surface were investigated. Both the experimental and the calculated
results were found to be in excellent agreement with those reported by Legg et al. [8] and are therefore not repeated here. The topmost interlayer spacing is slightly reduced (by about 1.5%) with respect to the bulk value: A minimum averaged reliability factor $r = 0.15$ was determined for four beams, using the $r$-factor program by Zanazzi and Jona [14].

The formation of the N overlayer causes not only the appearance of new fractional-order beams, but also affects the $I-V$ spectra from the integral-order beams. As an example fig. 1 compares the $I-V$ data of the (00) beam at $11.6^\circ$ angle of incidence for both the clean (b) and nitrogen-covered (a) Fe(100) surface. For this reason also the $r$-factors for the integral-order beams were included into the structural analysis.

A cubic (100) face exhibits three types of high symmetry adsorption sites, namely the fourfold hollow and "on top" sites, and the twofold bridge site. The latter requires averaging over two domains in order to retain the fourfold symmetry of the c(2 x 2) (or$\sqrt{2} \times \sqrt{2}$)R45° lattice. Calculations were performed for each of these three models. The atomic radius of N was allowed to vary between 0.5 and 1.0 Å, giving rise to corresponding changes in the spacing $d_N$ between the nitrogen and the first iron layer in steps of 0.05 Å. The Fe–Fe layer spacing in the bulk is 1.433 Å. The distance between the two topmost Fe layers $d_{Fe}$ was varied in steps of 0.03 Å between 1.35 and 1.60 Å.

Fig. 2 shows a set of experimental data taken at normal incidence together with representative theoretical spectra calculated for the three models indicated. It becomes evident that the best agreement is obtained for the fourfold hollow sites structure as reproduced in fig. 3a. Some calculations were also

![Fig. 1. Intensity–voltage spectra for the (00) beam of the adsorbate covered (a) and clean (b) Fe(100) surface. Angle of incidence $\theta = 11.6^\circ$.]
performed for the "underlayer" model (i.e. where the N atoms are located below the topmost layer of metal atoms) as derived for the N/Ti(0001) system [15] but the results gave no evidence for such a structure in the present case.

Quantitative structure analysis was now performed for the fourfold hollow site model of fig. 3a. Figs. 4 and 5 compare representative theoretical spectra calculated for varying $d_N$ and $d_{Fe}$, respectively, with the corresponding experimental data. Only those results are reproduced which revealed to be sensitive to small variations of the structural parameters. It turned out that $d_N$ affected

---

Fig. 2. $I-V$ spectra calculated for N atoms in the three high-symmetry adsorption sites are compared with the experimental data (normal incidence, $\theta = 0^\circ$).
mainly the fractional-order beams, but that the integral-order beams, on the other hand, were essentially influenced by variations of $d_{Fe}$.

The best quantitative agreement between experiment and theory was sought by using the $r$-factor concept introduced by Zanazzi and Jona [14]. Variation of both $d_N$ and $d_{Fe}$ changes also the bond length between the N atoms and the underlying Fe atom (see fig. 3b). As a consequence the minima of the $r$-factor curves are shifted systematically to lower values of $d_{Fe}$ if $d_N$ is increased. This
Fig. 4. Comparison between experimental spectra (dotted lines) and theoretical data (full lines) calculated for various interlayer spacings $d_{Fe}$ between the topmost two planes of Fe atoms. The distance $d_{N}$ between the plane of N atoms and the first layer of Fe atoms was kept fixed at 0.25 Å.

becomes evident from fig. 6 showing data for several fractional order beams. The expansion of the Fe–Fe layer spacing at the surface becomes evident from fig. 7 which shows $r$-curves for various integral order beams as a function of $d_{Fe}$ for the optimum $d_{N}$ (≈ 0.25 Å). The minimum ranges between 1.51 and 1.57 Å.

The optimum values for the structural parameters were found, as usually, by
averaging over all available r-factor curves. The average was taken over 16 beams, namely 6 beams at normal incidence and 10 beams at $\theta = 11.6^\circ$, $\varphi = 0^\circ$. The resulting data are shown in fig. 8 as variation of $r$ with $d_{Fe}$ at different values of $d_N$. The minima are relatively flat which is mainly caused by the fact that a series of beams remains nearly unaffected by relatively large variations of the interlayer distances. The best fit between theory and experiment is obtained for $d_N$ between 0.25 and 0.30 Å, yielding an optimum value
Fig. 6. $r$-Factor averaged over 5 half-order beams as a function of $d_{Fe}$. Parameter is $d_N$.

Fig. 7. $r$-Factor for 5 integral-order beams as a function of $d_{Fe}$; $d_N = 0.25 \, \text{Å}$ was kept constant.
of $d_N = 0.27 \text{ Å}$ and $d_{Fe} = 1.54 \text{ Å}$ as sketched in fig. 3b. The distance between the topmost two Fe layers is thus increased by 0.11 Å ($= 7.7\%$) with respect to the bulk value. If compared with the clean Fe(100) surface, this spacing is expanded by 0.14 Å ($= 10\%$) under the influence of the N overlayer. The bond length between the N atom and the underlying Fe atoms is 1.83 Å, while its distance to the four neighboring Fe atoms in the topmost layer is 2.04 Å. From the spread of the $r$-factor minima for the various beams and from the fact that the minima in fig. 8 are relatively flat, it is concluded that the indicated numbers are accurate to within ±0.05 Å as becomes evident from inspection of figs. 6–8.

5. Discussion

To our knowledge so far only three papers concerning a LEED analysis of a surface containing nitrogen atoms have appeared in the literature:
With the Ti(0001) surface (exhibiting completely different structure and symmetry than the system investigated here) a \((1 \times 1)\)-N structure was analysed in terms of penetration of nitrogen atoms below the topmost layer of Ti atoms \[15\]. The resulting structure is very similar to that of bulk TiN.

As with Fe(100), adsorbed N atoms form a \(c(2 \times 2)\) structure on the Mo(100) surface which was investigated by Ignatiev et al. \[16\]. In this work again on the occupation of fourfold sites was concluded, but the agreement between theory and experiment was not very satisfactory. The estimated distance between the nitrogen and the first metal layer \(d_{N} = 1.0 \text{ Å}\) is considerably larger than that derived in the present work. The resulting Mo–N bond length of 2.45 Å appears to be unreasonably large. No expansion of the topmost two metal layers was allowed which would probably have improved the result.

Burkstrand et al. \[17\] studied the \(c(2 \times 2)\) structure formed by N atoms on a Cu(100) surface by means of an averaging technique and concluded again on the occupation of fourfold sites. The interlayer distance \(d_{N} = 1.45 \text{ Å}\) appears again to be rather large (even in spite the fact that Cu–Cu distance within the surface layer, \(d_{Cu} = 2.55 \text{ Å}\), is smaller than in the case of iron, \(d_{Fe} = 2.87 \text{ Å}\)). A reexamination of the data suggested, however, that \(d_{N}\) is more likely around \(0.8 \text{ Å}\) \[18\] which would correspond to a metal–N bond length of similar order as concluded in the present work.

As stated in the Introduction, the main motivation for the present study was the suggested similarity between the Fe(100)-\(c(2 \times 2)\) surface structure and the structure of bulk Fe\(_{4}\)N leading to the concept of "surface nitride". This assumption is nicely confirmed by the present results: Fe\(_{4}\)N crystallizes in the fcc lattice. In the (002) plane a N atom is coplanar with the four nearest Fe atoms 1.89 Å apart and is directly above another Fe atoms at the same distance. Within the (002) plane of Fe\(_{4}\)N the nitrogen atoms form also a "\(c(2 \times 2)\)"-structure. The local geometry of the adsorption site on the surface (fig. 3) differs in so far as the plane of the N atoms is 0.27 Å above the plane of the four topmost Fe atoms. The distance of the N atom to the underlying Fe atom, \(d_{1}\), is (within the limits of accuracy) identical to that in Fe\(_{4}\)N, while the distance to the four Fe atoms in the surface layer is somewhat larger (\(d_{2} = 2.04 \text{ Å}\)).

According to the structure model shown in fig. 3 and to the resulting values of \(d_{1}\) and \(d_{2}\) a substantial (if not the dominating) contribution of the bond formation arises from coupling of the N atom to the underlying Fe atom in the second atomic layer. The resulting atomic radius of the chemisorbed nitrogen atom (0.59 Å) is between the N triple bond (0.55 Å) and double bond radius (0.62 Å). The strength of the Fe–N bond (as derived from the adsorption energy) is about 140 kcal/mol \[3\] and thus again between the corresponding values for the nitrogen triple bond (225 kcal/mole) and the double bond (100 kcal/mole) \[19\]. Thus a consistent picture for the structural and bonding properties of the N/Fe(100) system is reached.
Acknowledgements

The authors are grateful to Dr. V.L. Moruzzi for providing data on the iron and nitrogen potentials. Financial support of this work was obtained from the Deutsche Forschungsgemeinschaft (SFB 128).

References