Structural Analysis of the Fivefold Symmetric Surface of the Al70Pd21Mn9 Quasicrystal by Low Energy Electron Diffraction

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Abstract
The atomic structure of the fivefold symmetric quasicrystal surface of Al70Pd21Mn9 was investigated by means of a dynamical low energy electron-diffraction analysis. Approximations are developed to address the high structural complexity of quasicrystals, yielding average layer-by-layer structural and compositional information. A mix of several relaxed bulklike terminations is found, consistently favoring dense Al-rich outermost atomic layers. These results can be understood in terms of principles known to govern surface structures of simpler, crystalline metals.

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Over the past ten years great strides have been made in our understanding of quasicrystalline alloys. In particular, since their discovery in 1984 [1], we have gained significant insight into the bulk atomic-scale structure of icosahedral and decagonal alloys through x-ray and neutron diffraction measurements coupled with structural modeling based upon realistic atomic pair potentials [2,3]. At the same time, surface sensitive measurements such as photoemission, scanning tunneling microscopy (STM), and low energy electron diffraction (LEED) [4–7] have become possible through advances in our understanding of the metallurgy of quasicrystalline alloys, which have permitted the growth of very sizable, high quality, single grain samples [8]. While much is understood about the bulk atomic-scale structure of quasicrystals, determining their surface structure presents new and important challenges. Information about the surface structure, composition, chemistry, topology, and possible surface reconstructions is a prerequisite to understanding recent findings that quasicrystalline surfaces and coatings exhibit low surface friction and high oxidation resistance, properties important to many technological applications.

The surface structures of both icosahedral and decagonal alloys have been examined previously with both STM and LEED techniques. Most germane to the present discussion is the work of Schaub et al., who investigated the surface structure of microfacets which formed on a surface of the Al$_{70}$Pd$_{21}$Mn$_9$ icosahedral alloy, oriented close to the twofold surface, upon heating in vacuum [6,7]. The microfacets themselves grow perpendicular to a fivefold axis of the bulk alloy and exhibit fivefold symmetry in their LEED patterns. The surface topology of these facets can be described by a set of atomically flat terraces with steps of two different heights in a succession corresponding to a quasiperiodic (Fibonacci) sequence. However, the STM study suffered from limitations typical of the technique: It could not provide information about the chemical composition of the surface being imaged, nor could it clearly correlate certain STM features common on the terraces with atomic structures.

Previous LEED studies have been limited to observations and discussions of the symmetry and spacing of the diffraction spots, in order to determine whether the surface retains the quasicrystallinity of the bulk. They did not exploit the strength of dynamical LEED as a tool for obtaining atomic-scale information on surface structure and composition.

In this Letter we describe the results of a dynamical LEED analysis on the fivefold surface of an Al$_{70}$Pd$_{21}$Mn$_9$ alloy. Specifically, we find that the surface is a mixture of very similar terminations with a dense, Al-rich outermost atomic layer followed by a layer containing about 50% Al and 50% Pd. This information on composition and sub-surface structure complements that available from STM. Furthermore, it provides a basis for predicting and understanding the step heights present on these surfaces, since steps must separate the different terminations. Together, these lay the groundwork for a more quantitative understanding of the atomic-scale structure of quasicrystalline surfaces.

As with bulk x-ray and neutron diffraction, the LEED pattern for an ideal quasicrystal surface should exhibit sharp intense reflections (beams) at well-defined exit angles (Fig. 1). This leads to conventional-looking LEED IV curves, as shown in Fig. 2. To obtain such data, the sample is treated in an ultrahigh vacuum chamber equipped for Auger electron spectroscopy (AES), LEED, and other standard methods of surface analysis and
FIG. 1. LEED pattern at 85 eV incident electron energy and normal incidence. For each major set of five equivalent spots, only a single spot is indexed. The dark shadow at top and middle is the sample manipulator, which obscures some spots.

FIG. 2. Best-fit experimental and theoretical IV curves, averaging theory equally over the ten best individual terminations, including relaxations in the top four interlayer spacings of each. The intensity scales are arbitrary.

Our method of surface preparation in ultrahigh vacuum involves ion bombardment at room temperature and annealing (to temperatures in the range 870–1100 K), and is similar to that employed by Schaub et al. in the STM work [6,7]. These techniques are used until no surface contaminants (mainly oxygen) are detectable by AES, and until the fivefold pattern reaches a high quality. There is no evidence for a surface phase transformation, such as that reported for decagonal AlCuCo [12].

Like previous authors, we find that the symmetry and spacing of the LEED beams indicate that the fivefold surface of this alloy retains the bulk quasicrystallinity. IV curves were measured at normal incidence for six sets of bright diffraction spots in the energy range 70–260 eV. Each set consists of five symmetrically equivalent spots which we average, thus producing a database of six symmetry-inequivalent IV curves, indexed as shown in Figs. 1 and 2. Under a wide variety of surface preparations, a sharp fivefold diffraction pattern was obtained, and the IV curves were very similar. Thus, the IV data shown in Fig. 2 are very robust. These form the experimental basis for deducing the surface structure from diffraction theory.

Quasicrystals pose special challenges to diffraction theory, previously worked out for the cases of x rays and neutrons. In LEED, multiple scattering adds further complications: The aperiodicity of quasicrystals, and of quasicrystal surfaces in particular, implies a wide variety of local structures, and an intractable number of atoms with different scattering properties; note that the scattering properties of an atom depend in effect on its local environment in the case of multiple scattering. Thus, approximations in the LEED theory are necessary, and an appropriate LEED formalism, described in this Letter, was developed for a quasicrystalline surface region.

As a starting point for the LEED analysis, we used the bulk structure of icosahedral AlPdMn determined by x-ray and neutron diffraction [3] and produced first a large set of possible ideal bulklike fivefold surface terminations bounded by a box of $100 \times 100 \times 50 \, \text{Å}$. As a result of quasiperiodicity, all of these terminations are, in principle, different. Many of them have, however, very similar chemical compositions and geometry so that only a finite number of qualitatively different terminations has to be considered. Each of these surfaces is represented as individual layers (parallel to the surface) of coplanar atoms with a particular composition and density. The average $t$-matrix approximation [13] is applied to each layer with its own bulklike composition to generate a unique atom with average scattering properties. The layer density also scales the contribution from each layer.

To account for multiple scattering, we use the giant matrix approach in the framework of the Beeby matrix inversion [14]. We assume that the scattering properties of all atoms in a particular layer parallel to the surface are equal. We describe the average neighborhood of an average atom in a given layer by a kind of radial distribution...
function: A representative number of neighbor atoms is distributed uniformly on rings, with proper polar angles but without regard to proper azimuthal angles. This local “cluster” is summed over in the sense of the lattice sum familiar in the $X$ matrix of LEED theory [14]. The $X$ matrix, which describes the propagation of the electron wave field within and between layers, is thereby assumed to be independent of the azimuthal angle; this works best at normal incidence, as used in the experiment. Inversion of the matrix $1 - X$ then introduces multiple scattering paths to infinite order. Checks on the sufficiency of these approximations were performed by also using more representative structural models. These checks will be presented elsewhere, together with further details about phase shifts, Debye temperatures, muffin-tin zero, and inelastic damping, which are standard. The Pendry $R$ factor [15] was used to evaluate the fit to experiment.

In analyzing the experimental LEED data, we first assumed single terminations of the bulk, i.e., no mix of terminations due to possible steps. The top two interlayer spacings were optimized for each termination within a cube of 50 Å depth, yielding $R$ factors varying from about 0.42 to 0.85 for the different terminations. The better fitting terminations share a common feature: a pair of closely spaced dense layers with an interlayer spacing of about 0.4 Å. The top one of these two layers is composed primarily of Al, mixed with about 10% Mn, while the second one contains about 50% Al and 50% Pd and only small amounts of Mn (Fig. 3). In line with the STM observations of Schaub et al. [6,7], many of these terminations are separated by steps with heights of 4.08 or 6.60 Å. Low energy ion scattering data also support the composition of the favored layers [16].

Next, the ten best terminations were averaged together with equal weights to model the case of equal-sized terraces, each having one of the different terminations. By an automated optimization after the intensity mixing, we got information about the average interlayer spacings and could reduce the $R$ factor to 0.31, which is comparable to that found in many other successful structure determinations, particularly for complex surfaces. The topmost interlayer spacing of 0.38 Å is reduced by 0.1 Å from the bulk value of 0.48 Å, which trend is consistent with clean single-crystal results of comparable local buckled structures (e.g., stepped fcc surfaces); the interlayer spacings of deeper layers are almost bulklike. Note that the two topmost layers are so close that they might be considered a single, buckled layer, in which case the lateral density of this buckled layer is 0.135 atoms/Å$^2$, close to the value of 0.141 atoms/Å$^2$ for the close-packed Al(111) fcc surface. This high density is probably one factor which favors the terminations we find for the quasicrystal, since for metals, the closest-packed face generally has the lowest surface free energy.

The Al-rich terminations found here also correlate with factors known to govern surface compositions of crystalline alloys. At those simpler surfaces, one finds

![Fig. 3. One of the favored terminations of Al$_{70}$Pd$_{21}$Mn$_9$: (a) The two topmost planar bulk layers perpendicular to the fivefold axis; many Mn atoms in the top layer are surrounded by pentagons in the second layer, one of which is indicated at the lower right. In (b), the layers are drawn at their respective depths $z$, as bars with thickness proportional to the atomic density in each layer, together with their approximate composition; parentheses indicate minority species.](image-url)
preferential segregation by components with the lower surface free energy, larger atomic volume and more positive heat of solution [17]. (Surface free energy typically is the dominant factor.) The Al-rich terminations found here are expected if these factors are taken into account.

As shown in Fig. 3(a), many of the Mn atoms in the topmost layer of the favored terminations are surrounded by five atoms (Al or Pd) in the second layer. The orientation of these Al/Pd “pentagons” in the second layer is the same for all terminations, and by connecting their edges with straight lines, one obtains a Fibonacci pentagrid. Interestingly, in the STM study [6], fivefold symmetric holes with very similar quasiperiodic long range order were found; one may speculate that these holes can be identified with Mn atoms, surrounded by five atoms in the second layer since no other features fit the STM images better.

In summary, the atomic structure of the fivefold symmetric quasicrystal surface of Al$_{70}$Pd$_{21}$Mn$_9$ has been investigated by means of a dynamical LEED analysis. Approximations were developed to make the structure of an aperiodic, quasicrystalline surface region accessible to LEED. A mix of several relaxed, bulklike terminations is found with a dense Al-rich layer on top followed by a layer with a composition of about 50% Al and 50% Pd. The interlayer spacing between these two topmost layers is contracted by 0.1 to 0.38 Å, and the lateral density of the two topmost layers taken together is similar to that of an Al(111) surface.

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