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Kurnosikov, O.; Nietsch, J.H.; Sicot, M.V.; Swagten, H.J.M.; Koopmans, B.

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Long-Range Electron Interferences at a Metal Surface Induced by Buried Nanocavities


Department of Applied Physics, cNM, Eindhoven University of Technology, 5600MB Eindhoven, The Netherlands

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Apparent $c(2 \times 2)$ superstructures within the narrow beams of an interference pattern spreading in the ⟨100⟩ directions at the surface of Cu(001) are observed by scanning tunneling microscopy. These features are induced by electron scattering from Ar- and Ne-filled subsurface nanocavities. The beams originate from electron anisotropy resulting in focusing of bulk electrons. We developed a model providing a good agreement between simulations and experiments. Particularly, a simple explanation of the angular distribution for the interference pattern and the period in the superstructure is found.

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Electron interference at a surface is well-known phenomenon and can be induced by electron scattering at surface defects [1] or by subsurface scattering at interfaces [2]. Interestingly, nanosized reflectors can be formed by aggregated noble-gas nanocavities [3], which have been studied in different metals and semiconductors for their formation and growth [4], and for surface and bulk structural changes [5,6]. However, only recently Ar-filled nanocavities were studied with scanning tunneling microscopy (STM) aiming particularly at the quantum well (QW) states formed between surface and nanocavity [7,8]. It was demonstrated that the area of modified surface conductance of Al(111) generally corresponds to the shape of the upper facet of the subsurface object [8]. For Ar-nanocavities in Cu, it was shown [9] that the spatial distribution of the conductance in the area right above the upper cavity facet is much more complicated and additionally determined by the band structure anisotropy of the host material and the related focusing of hot electrons [10]. In the paper of Kurnosikov et al. [9], also preliminary data were shown for conductance modifications away from the center of the nanocavity for specific crystallographic directions. Although this was tentatively assigned to the influence of electron anisotropy, the authors have stressed the lack of a consistent description and analysis of the experimental STM image, which did not allow for a conclusive physical understanding at that time.

Inspired by these initial observations, in this Letter we entirely focus on long-range electron interferences due to subsurface nanocavities when using STM or STS to address the Cu(001) surface conductance away from the area right above a nano-object. We have observed multiple long-range narrow interference beams in the ⟨100⟩ directions superimposed on the atomic structure of the substrate, by which an apparent $c(2 \times 2)$ superstructure within the beams is revealed. The slight difference between the in-plane lattice constant and oscillation period related to the planar component of the ⟨101⟩ wave vector, leads to a moiré pattern that is additionally modified with a periodic phase shift of the apparent surface arrangements. By taking into account the atomic surface periodicity in our previously developed model dealing with injection, propagation, reflection, and interference of electrons, as well as the anisotropy of electronic properties in Cu [9], an excellent agreement between the experimental and simulated STM images is obtained. Our analysis gives a clear and consistent explanation of the origin of long-range interference beams and the apparent $c(2 \times 2)$ superstructure, and is further strengthened by the observation of similar patterns when using Ne-filled cavities instead of the exclusive use of Ar so far [7–9].

To implant noble gas and form subsurface nanocavities, the Cu(001) sample was bombarded by Ar or Ne with an energy of 5 keV and current density of about 30–80 μA/cm² during 30 min. After the implantation, the sample was annealed at 1000 K for 5 min. The STM study was done at 77 K in an ultrahigh vacuum (UHV) “Omicron” STM. The x-ray photoelectron spectroscopy (XPS) analysis made after implantation without annealing reveals the $2p_{1/2}-2p_{3/2}$ double peak of argon (Fig. 1). The

![FIG. 1 (color online). XPS spectra of the Cu sample after implantation of Ar without annealing (1) and annealing at (2) 1000 K, (3) 1050 K, and at (4) 1075 K. The positions of Ar $2p_{1/2}$ and $2p_{3/2}$ maxima of binding energy are marked by the vertical bars. For better visibility the curves 2 and 3 are shifted down by 30 and 60 counts, respectively. The corresponding structure is shown in the inset.](https://example.com/figure1.png)
binding energy corresponds to the values reported elsewhere [11], and it is shifted with respect to the binding energy of free argon, indicating that Ar is implanted in Cu. The shift of the $2p_{1/2}$-$2p_{3/2}$ peaks back to higher binding energy together with a reduction of their amplitude is observed after annealing at 1000 and 1050 K for 5 min, showing the formation and growth of Ar-filled nanocavities [12] as well as the loss of argon. After annealing at 1075 K for 20 min the argon peaks vanish. Although not specifically tested in great detail, similar processes should take place when implanting Ne [13]. This shows that thermal treatments allow us to control size and density of the subsurface nanocavities formed by different gases.

Before presenting the main result we would first like to ensure that similar electronic features at the surface are induced by implantation of Ar or Ne and annealing at 1050 K. The STS maps reveal a very pronounced variation of the surface conductance above the nanocavities depending on the applied bias voltage $V_b$ (Fig. 2). The characteristic size of the spots corresponding to the perturbed conductance is only a few nanometers. Remarkably, the spots in the STS images are strongly anisotropic (Fig. 2) and differential conductance in the center of the spots reveals an oscillation when sweeping the energy (not shown). This points out that the formation of standing electron waves and localized QWs between surface and cavity interface, is a robust and reproducible phenomenon not only restricted to Ar as previously reported [9], but also present for other noble gases such as Ne.

Now the presence of embedded nanocavities and QW formation at the surface is demonstrated, we can focus on the long-range interferencelike beams. These are clearly visible in the STM images at high resolution (Figs. 3 and 4) obtained by mapping of uncompensated tunnel current. This regime is almost the constant height mode of STM, obtained by mapping of uncompensated tunnel current.

Visible in the STM images at high resolution (Figs. 3 and 4) are the long-range interferencelike beams. These are clearly present for other noble gases such as Ne. Not only restricted to Ar as previously reported [9], but also present for other noble gases such as Ne.

Later on, we will come back to this surprising behavior. (vi) Usually, four beams are present in STM images. However, very often we observe only one beam [Fig. 4(a)], two opposite [Fig. 4(b)] or two adjacent [Fig. 4(c)] beams, and sometimes three beams. Even if we observe all four beams [Figs. 3(a) and 3(b)], their length and contrast can be unequal. Remarkably, the number of observed beams is generally the same for several nanocavities within one scan of the surface. However, when the tip state accidently changes in time, we observed the appearance/disappearance of extra beams while keeping the atomic resolution of the surface.

![FIG. 2 (color online). Two couples of differential conductance maps of the same Cu surfaces after implantation of Ar (a), (b), 60 × 60 nm², or Ne (c), (d), 15 × 15 nm², and annealing at 1050 K. The bias voltages $V_b$ of (a) 400 mV, (b) 600 mV, (c) 500 mV, and (d) 800 mV were applied at the STS scanning with tunnel current 5 nA stabilized by the closed feedback loop.](image)

![FIG. 3 (color online). (a), (b) and (c) STM images of Cu surface above the nanocavities with atomic resolution: (a) typical interference pattern above a single Ar-filled nanocavity, 15 × 15 nm², $V_b = 46$ mV, $R_t = 2$ MΩ; (b) several Ar-filled nanocavities induce overlapping beams of interference fringes, 18 × 18 nm², $V_b = 7$ mV, $R_t = 0.05$ MΩ; (c) a $c(2 \times 2)$ superstructure within the interference beam near a Ne-filled nanocavity, 12 × 12 nm², $V_b = 7$ mV, $R_t = 0.1$ MΩ; (d) schematic drawing similar to (c) (scale not corresponding) illustrating the shift of the rows within the superstructure.](image)
face. All these facts prove that the observed \(c(2 \times 2)\) superstructure within the beams is not intrinsically inherent to the atomic arrangement, but an electronic effect, where the tip state is crucial. In fact, the injection in the bulk by a STM tip can strongly relate to the angular distribution of electron \(k\) vectors at the tip apex. Although only the component perpendicular to the surface \(k_\perp\) can be taken into account by a simplest approach for the tunneling process in STM, \(k_\parallel\), which is conserved for elastic tunneling, should determine the \(k\) vector of electrons after injection [14]. If \(k_\parallel\) at the tip end is nonuniform, for example, due to its particular shape and atomic structure, the injection in the bulk should not be characterized with axial symmetry.

If the injection is axial-symmetric, the specific shape of the observed pattern is obviously determined by the inhomogeneous angular distribution of electrons coming back to the surface after reflection from the nanocavity. There are two reasons for this: the bulk anisotropy of the metal and the shape of the subsurface reflector. Our model based on the approach developed earlier [9] takes these two factors into account. In order to describe the features obtained in the present experiments, we modified the model by including the periodical position-dependent tunneling probability corresponding to the atomic structure of surface.

Our model uses the anisotropic band structure of copper \(E(k)\) [15]. Moreover, an electron wave can additionally favor only a few specific directions when traveling over a long distance, a so-called focusing effect already studied for other systems [10,16–18], intimately relates to specific features of the band structure. In our experiment, the beams of interference fringes appear always along the \((100)\) directions indicating the concentration of coherent electron waves due to this effect. The shape of the subsurface reflector can also determine particular angles of backscattering and contribute to the formation of the long-range beams especially if it contains flat planes. Several experimental reports [7,8,19] prove the presence of flat facets forming nanocavities in single-crystalline solids. In our model, the interface with the nanocavity is built by three types of differently oriented facets \(\{001\}, \{110\}\) and \(\{111\}\) similarly to a Wulff construction [20].

Results of our calculation and comparison with the experimental data give a typical depth of the nanocavity of 3–15 nm, while the size of the upper facet is estimated to be a few nanometers. The simulation of the conductance map is presented in Fig. 5(a). The simulation shows long radial beams in the \((100)\) directions [Fig. 5(a)] very similar to those observed in the experiment [Fig. 3]. The simulation [Fig. 5(a)] also shows the rounded rectangular-like feature around the center. Additionally, it shows the structure just above the nanocavity which is ill resolved in the experimental pattern with high resolution (Figs. 3 and 4) because of the feedback operation, but appears on the experimental images with low resolution in the STS regime [Figs. 2(a) and 2(b)]. We address readers for these details to our previous publication [9].
The origin of beams in the $\langle 100 \rangle$ directions is the concentration of electron waves along the $\langle 101 \rangle$ directions due to the focusing effect in copper, which is shown schematically in Fig. 5(b). The same directions correspond to the $\{101\}$ facets of the nanocavity effectively reflecting the electrons. The length of the beams can depend on several factors. The fundamental limit is the electron mean free path which can reach hundreds of nanometers at low temperature. But there are other factors that are specifically related to the configuration of the system, such as size of the $\{101\}$ facets reflecting electrons, energy spread of the injected electrons, and the signal-to-noise ratio in the STM system. Additionally, a particular tip state can reduce injection of electrons in a direction favorable for focusing.

The $c(2 \times 2)$ superstructure within the beams appears as a superposition of the atomic arrangement and the interference fringes at Cu(001) surface schematically shown in Fig. 5(c). The formation of the $c(2 \times 2)$ superstructure is due to the accidental coincidence of the Cu unit cell size $a$ with the periodicity of interference fringes $d$ at the surface which is determined by the planar component of $k$ vector of electrons propagating in the $\langle 101 \rangle$ directions [Fig. 5(b)]: $d = \pi \sqrt{2}/k_{\langle 101 \rangle}$. This gives the opportunity to make a rough estimation of $k_{\langle 101 \rangle}$ from our experiment: $k_{\langle 101 \rangle} \approx 1.24 \times 10^{10} \text{m}^{-1}$, which is close to the tabulated value $k_F = 1.36 \times 10^{10} \text{m}^{-1}$ within an isotropic approximation [21]. Additionally, the slight difference between the lattice parameter $a$ and the period $d$ leads to the formation of a moiré pattern with a period $2L$, where $L$ is determined as $L = ad/(2[a - d])$. As a result, the modulation of the contrast within the beam, and in the shift of arrangement of rows of superstructure indicated by arrows in Figs. 5(a) and 5(c), is formed each half of period $L$. On the one hand, the value $L = 3 \text{ nm}$ estimated in Fig. 3(c) allows us to correct the $k_{\langle 101 \rangle}$ presented above to get more precise value, but on the other hand, we realize that this correction can be done only within a simple approach considering the unperturbed band structure of copper. This perturbation should be due to extra stresses in the vicinity of the nanocavity and leads to a slight variation of both the value of $k$ vector corresponding to the focusing direction, and the focusing direction itself. Although we still can define a planar component of the $k$ vector at the surface from experiment, the precise determination of the complete $k$ vector corresponding to focusing desires additional studies.

For other materials, if a similar coincidence of lattice parameter and wave vector of reflected electrons is achieved, a similar superstructure can also be observed although other directions of focusing can be considered.

In summary, we have studied the variation of differential conductance of the Cu(001) surface above buried nanocavities filled by Ar or Ne. Electron focusing plays a crucial role in formation of the interference beams in the $\langle 100 \rangle$ directions. Superposition of the interference fringes with the surface atomic structure results in a $(2 \times 2)$ superstructure, all in perfect agreement with results of our electron interference calculations.

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