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LETTER TO THE EDITOR

Oscillatory electron–phonon coupling in ultra-thin silver films on V(100)

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Received 9 June 2000

Abstract. The temperature dependence of peak widths in high-resolution angle-resolved photoelectron spectroscopy from quantum well states in ultra-thin Ag films on V(100) has been used to determine the electron–phonon coupling constant, $\lambda$, for films of thickness 1–8 layers. A strong oscillatory variation in coupling strength is observed as a function of film thickness, peaking at a two layer film for which $\lambda \approx 1.0$. A simple theory incorporating interaction of the photohole with the thermal vibrations of the potential step at the adlayer–vacuum interface is shown to reproduce the main features of these results.

The physical properties of metal surfaces and thin metallic films are often different from those of the bulk. In general, reduced dimensionality and a higher degree of localization enhance correlation in electronic systems. In this regard quantum-well (QW) states in thin films provide a testing ground of such effects, because the degree of localization can be varied by changing the film thickness. While a significant body of previous studies has explored the one-electron behaviour of quantum-well states, there have been no comparable studies on the influence of the changing degree of localization in ultra-thin quantum wells on the many-electron properties. Here we present data which provide a clear manifestation of essentially new phenomena in the electron–phonon coupling in such systems.

Specifically, we report measurements of the electron–phonon coupling in ultra-thin silver films deposited on a V(100) surface obtained from the temperature dependence of the widths of angle-resolved photoelectron spectroscopy (ARPES) peaks from QW states in the films. An exceptionally strong oscillatory change in the coupling strength is observed, as a function of the thickness of the silver film, with a pronounced maximum associated with the QW state corresponding to a 2 ML (monolayer) film. We show that this observed variation of the coupling strength can be explained in terms of the interaction of the photohole with the thermal vibrations of the potential step at the adlayer–vacuum interface. The oscillatory behaviour reflects mainly the variable degree of localization of the QW states which, through the amplitude of the wave-function at the surface-vacuum interface, influences the coupling to the surface vibrational mode. The increased electron–phonon coupling strength of these films, which at certain thicknesses substantially exceeds the value for bulk Ag, could have important consequences for the possibility of tailoring ultra-thin structures to have particular characteristics. For example, an increased superconducting transition

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0953-8984/00/280477+06$30.00 © 2000 IOP Publishing Ltd L477
temperature is one possibility, experimentally observed in number of systems [1], although strong electron–phonon coupling itself is not sufficient to generate a high superconducting transition temperature in thin film and surface states [2].

ARPES has already been shown to be particularly suitable for studying many-body interactions in quasi-two-dimensional systems, because in such systems it measures directly the photo-hole spectral function, and there have been several such studies of metallic surface states or in thin films. A study of the Mo(110) surface state [3] showed that all three interaction terms, electron–electron, electron–phonon and defect scattering, can be deduced from the temperature and binding energy dependence of the photoemission peak width. Other studies on metallic surface states on Cu [4], Be [2, 5] and Ga [6] focused on the electron–phonon coupling term alone. The electron–phonon coupling constants determined in this way for surface states can be significantly different from those measured in the bulk. For example, the value for the Be(0001) surface state is four times that found in the bulk by transport measurements. ARPES has also been used to study the electron–phonon coupling in QW states in films sufficiently thick for the results to be judged to be characteristic of the bulk metal [7, 8]; in this case some differences in the values deduced have been attributed to the momentum-resolved character of an ARUPS measurement as opposed to a directionally-averaged transport measurement. There has been just one previous study of electron–phonon coupling in a thin quantum well (of Na on Cu(111)) which we will discuss in the context of our own results [9].

The experiments reported here were carried out at the National Synchrotron Light Source using undulator beamline U13UB which provides photon energies in the range between 12 and 23 eV. The electron energy analyser was a Scienta SES-200 which collects simultaneously a large energy and angular window (∼12°) of the photoelectrons. This reduces the time needed for data acquisition and ensures that a wide range of the states in $k$-space are recorded under identical conditions of temperature and surface cleanliness. The combined instrumental energy resolution could be set to a value in the range 8–25 meV, small enough to make no significant contribution to the measured photoemission peak width from the QW states. The angular resolution was ∼0.2°. The base pressure in the experimental chamber was 4 × 10⁻⁹ Pa. Detailed characterization of the epitaxial growth of silver on V(100) has been presented in [10, 11] while [12–15] describe in detail the QW state properties in this system. Films of 1 ML and 2 ML are stable up to 800 K [11], while thicker films are stable only up to 350 K.

Figure 1 shows the photoemission spectra from 1 and 2 ML films (shown overlaid) taken at 60 K. The lineshape of the peak from the QW state is completely determined by the photo-hole self-energy $\Sigma(k, \omega)$. Depending on the rate of change of the imaginary part, the spectrum may be either Lorentzian-like (Im$\Sigma$ approximately constant), asymmetrical or even (for Im$\Sigma$ very strongly dependent on energy) 'double-peaked' [2]. In figure 1 the experimental spectra are fitted with the ‘Fermi liquid’ lineshape: $2\text{Im}\Sigma(\omega) = \Gamma_0 + 2\beta\omega^2$. The energy-independent term, $\Gamma_0$ represents the sum of impurity (or defect) scattering and phonon scattering terms (the phonon contribution being constant at a specific temperature for all energies studied here) and the quadratic term is the electron–electron contribution. Both peaks are fitted by almost the same electron–electron coupling parameter $\beta \simeq 0.04$ eV⁻¹, a value significantly larger than in bulk-like silver films [7]. These fits lead to values for the constant term $\Gamma_0$ of approximately 150 meV for the 1 ML QW state and 100 meV for the 2 ML QW state. We note, however, that the quality of these fits is not very sensitive to the relative values of $\Gamma_0$ and $\beta$, so this separation of the electron–electron coupling is not very reliable for these data. This is especially true for the 1 ML QW state for which the ARPES data clearly show a sloping background, and the exact background used evidently influences the degree of asymmetry of the resulting peak.

This ambiguity in separating the various contributions to the peak widths does not, however, extend to the electron–phonon scattering which is of primary interest for our present
purposes and which can be obtained directly from the dependence of the measured ARPES peak widths on temperature; these data are summarized in figure 2. The temperature dependence is approximately linear at high temperatures with a slope of $2\pi \lambda k_B$ where $\lambda$ is the electron–phonon coupling constant. The linear fits superimposed in the data in figure 2 show that the gradient changes strongly with the film thickness. In figure 3 we plot these experimental $\lambda$ values deduced from these linear gradients (filled circles), as a function of the Ag film thickness $d$. The most prominent feature of the $\lambda(d)$ plot is the change in the coupling strength of the QW states to phonons when the thickness of the silver film is increased from 1 ML to 2 ML; the corresponding value of $\lambda$ for the 2 ML Ag film is more than four times larger than that for the 1 ML QW state ($\lambda_{1\text{ML}} \simeq 0.23$, $\lambda_{2\text{ML}} \simeq 1.0$) and more than three times larger than the recently reported value measured for a 19 ML Ag film, believed to yield a value characteristic of bulk Ag, grown on Fe(100) [7]. Apart from the prominent maximum in $\lambda$ at 2 ML, the Ag/V(100) system shows an additional maximum of the coupling constant around 5 ML. To our knowledge, this is the first demonstration of oscillations in the electron–phonon coupling strength with the thickness of the metallic film. The only relevant prior measurements are from 1 ML and 2 ML QW states in the Na/Cu(111) system [9]; these showed a small difference for the two layers, while both QW states showed stronger coupling than the corresponding bulk value.

The $\lambda$ value for the intrinsic surface state on clean V(100) ($\lambda_V \simeq 1.45$) is also shown in figure 3. It is derived from the mass enhancement of the surface state (band dispersion) near the Fermi level in the same manner as in the earlier surface state studies of Mo(110) [3] and Be(0001) [2, 5]. This value is one of the highest measured by ARPES so far and is significantly larger than the bulk value for V of 0.8 [16]. However, this large electron–phonon coupling constant of the vanadium substrate cannot explain the maximum in the coupling constants for the thin film QW states at 2 ML. If the large $\lambda_{2\text{ML}}$ value were solely due to the interaction of the silver overlayer with the electronic and phonon system of the underlying vanadium substrate, we would expect the coupling in the 1 ML film to be even stronger. Notice too,
Figure 2. The peak widths of QW state photoemission peaks plotted as a function of substrate temperature. From the linear fit of each set of data the corresponding electron phonon coupling constant ($\lambda$) was calculated and is shown in brackets. For the sake of clarity not all sets of data are shown.

Figure 3. The electron phonon coupling constant ($\lambda$) values obtained from figure 2 shown as a function of silver film thickness (solid circles). Calculated values of $\lambda$ assuming an effective mass $m^* = 1$ for all QWS are shown as open squares; calculated values including experimentally-determined values of the effective mass for each QW state are shown as open triangles. The experimental value of $\lambda$ for the V(100) surface state is shown at zero film thickness.
that the earlier study of the Na/Cu(111) system found electron–phonon coupling of the QW states for the 1 ML and 2 ML Na films to be found significantly larger than the bulk coupling constants of both Cu and Na, while the coupling constant for the surface state on Cu(111) is essentially identical to that of bulk Cu [4]. These results were taken to imply that the increased electron–phonon coupling strength has its origin in the low dimensionality of the films, rather than in any interaction with the substrate [9].

In order to understand the origin of the increased and oscillatory thickness-dependent electron–phonon coupling in our Ag on V(100) QW states, we have performed calculations for a simple model [17] in which the main contribution to the hole lifetime is assumed to be the interaction of the photo-hole with the oscillations of the potential step at the interface between the vacuum and the silver film due to thermal vibrations of the surface atoms. It can be shown that the phonon-induced lifetime $\tau_g$ of the photo-hole created in the QW band $g$, can be written as:

$$\frac{1}{\tau_g} = \frac{m_{x,y}}{M \frac{A}{\hbar^2}} [T_{g,g}]^2 (n(\omega_0) + 1) + \sum_{f<g} |T_{f,g}|^2 (2n(\omega_0) + 1) \right].$$

(1)

Here, $f$ and $g$ are discrete indices identifying the specific QW state(s) involved in the transitions. $A_c$ is the area of the surface unit mesh of the silver film, $M$ is the atomic mass of silver and $n(\omega_0)$ is the Bose–Einstein distribution. Equation (1) is derived assuming that the perpendicular surface vibrational mode is characterized by the Einstein frequency $\omega_0$ and is completely localized at the topmost layer. We have assumed $\omega_0$ to be independent of film thickness and taken the dispersion of the QW states to be isotropic and parabolic parallel to the surface plane with a mean effective electron mass, $m_{x,y}$. The effective transition matrix element, $T_{f,g}$, is given by

$$|T_{f,g}|^2 = V_R^2 |\Psi_f(z=0)\Psi_g(z=0)|^2$$

(2)

where $\Psi_f(z=0)$ and $\Psi_g(z=0)$ are the wave functions of the QW states between which the transition takes place, at the position of the potential step between the film and the vacuum $(z=0)$. $V_R$ is the height of this step at the film-vacuum interface. Figure 3 shows values of the electron–phonon coupling constant calculated from the slope of the $\hbar/\tau$ function in the high temperature range. The open squares correspond to values of $\lambda$ calculated assuming that the effective mass of all the QW states around the centre of the Brillouin zone is equal to one, while the open triangles show values of $\lambda$ obtained when the experimentally measured effective masses for each QW state are used.

These theoretical results show the oscillation of the coupling constant to be present, even if free-electron like dispersion of all the QW states is assumed. In this case the changes in the coupling appear to arise mainly from the different localization of the states, and specifically the amplitude of the associated wave-functions at the surface barrier. These amplitudes are influenced by the QW state binding energy, the more shallowly-bound states extending further into the vacuum, and by the degree of localization as determined by the film thickness, thicker films having more extended states which (when normalized) have lower amplitudes at the surface. For example the largest binding energy (approximately 1.5 eV) QW states are those in the 1 ML and 4 ML films which show weak electron–phonon coupling. The lowest binding energy state (0.2 eV) is that in the 6 ML film, but while this shows a higher coupling, the deeper (0.7 eV) but more localized state in the 5 ML film has slightly stronger coupling. The strongest coupling is found for the much more localized state in the 2 ML film with a comparable (0.6 eV) binding energy. There is also some influence on the coupling in this simple theoretical model associated with differences in the available phase space for the photo-hole decay. In particular, in the case of a 1 ML Ag film only a single QW state exists so only intra-band transitions are
possible, while for the thicker films, both, inter- and intra-band transitions can occur. However, calculations allowing only intra-band transitions in all films produce results similar to those shown in figure 3, indicating that this phase space consideration is secondary.

Of course, the model presented here is very simple. Being based only on a surface-related process, its contribution to the electron–phonon coupling in the QW states vanishes as the number of layers goes to infinity; no account is taken of the interaction with the bulk phonons. Nevertheless, this very simple version of the calculations reproduces the main qualitative features of the experimental data, and thus appears to identify the key underlying physics. In this regard we note that much better quantitative agreement is obtained when the measured effective mass for each QW state is included in the calculations.

In conclusion, we have shown that the electron–phonon coupling constant for silver films of only a few ML thickness oscillates strongly as the thickness is varied. A particularly high value of this constant, $\lambda$, is found for the 2 ML film. The oscillation and the high values of electron–phonon coupling constant found in the silver films deposited on V(100) have been explained in terms of the hole interacting with thermally-induced oscillations of the potential step at the adlayer–vacuum interface, and the variations in coupling of the QW states to this vibrational mode as the degree of localization (including that determined by the QW state binding energy) varies with film thickness.

The authors acknowledge the support the British Council and the Ministry of Science of Croatia (ALIS project, grant Zag/984/CRO/042 to the Zagreb and Warwick groups), the Ministry of Science of Croatia (grant 003501108 to the Zagreb group), the Engineering and Physical Sciences Research Council (Warwick group) and the US Department of Energy Contract No DE-AC02-98CH10886 (BNL group)

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