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Step effects on diffusion near a substrate reconstructive phase transition

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Step effects on diffusion near a substrate reconstructive phase transition: H on W(100)


I. INTRODUCTION

Surface diffusion plays an important role in many interesting surface processes, such as adsorption, catalysis, and film growth. Changes in the structure of a substrate may have a considerable effect on surface diffusion. For example, surface diffusion near an adsorbate or substrate phase transition is expected to have anomalous non-Arrhenius temperature dependence based on theoretical grounds. 1–3 Experimentally, such behavior has often been observed near adsorbate layer phase transitions. 4–6 However, the underlying physics for such observations is rarely understood, mostly due to the lack of knowledge of the nature of the relevant phase transitions. In contrast, the nature of the substrate reconstructive phase transition of the W(100) surface is well understood. 7–16 It provides an ideal model system to investigate the effect of the substrate phase transition on surface diffusion. As reported in a recent Letter, 17 the H diffusion anomaly near the reversible phase transition of W(100) from a (1 × 1) disordered phase at high temperature to a c(2 × 2) phase at low temperature was clearly demonstrated by low-energy electron diffraction (LEED) pattern experiments. The W(100) surface has been determined to have a fourfold symmetry. 18

Surface steps, even at low densities, can have a considerable effect on the surface diffusion process. 19 This is another example showing that changes in substrate (an increase in step density) can have significant effects on surface diffusion. In a recent study of CO diffusion on vicinal Pt(111) surfaces, 20–22 Ma and co-workers observed new and very interesting diffusion behavior related to steps. For diffusion perpendicular to steps, the diffusion was impeded by an extra step trapping potential well rather than an extra step barrier (Schwoebel-Ehrlich barrier). 20 For diffusion parallel to steps, in addition to the channels over the terrace and along the step edges, a new channel was observed that enhanced diffusion by about one order of magnitude. 21 Step effects, which are unavoidable in any macroscopic measurement that probes an area containing a large number of steps, can sometimes mask the true surface diffusion process such as intrinsic terrace diffusion. In this paper, we therefore extend our earlier study of the H diffusion anomaly near the W(100) substrate phase transition 17 to the stepped surface, in part to investigate possible step effects on the anomalous behavior of D near the phase transition.

The W(100) (1 × 1) surface has a fourfold symmetry and the diffusion of H on it is therefore expected to be isotropic. However, the reconstructed c(2 × 2) structure has a twofold symmetry and diffusion of H on it is expected to be anisotropic. For the clean surface and the surface with very low H coverage [less than 0.12 monolayer (ML)], the structure of the reconstruction at low temperature was originally deduced by symmetry arguments to consist of atomic displacements along the diagonal (110) surface directions to form zigzag chains. 9,14 The displacement magnitude of the top-layer atoms on the clean surface has been determined to be 0.22 Å. 15,22 The low-energy electron diffraction (LEED) pattern of the c(2 × 2) structure shows fourfold symmetry due to contributions from two orthogonal degenerate domains whose symmetries are only twofold (p2mg). When the hydrogen coverage exceeds about 0.12 ML, a change of the c(2 × 2) structure symmetry from p2mg of the clean surface to c2mm is induced. 10,13,14,23 From this change, it was deduced that hydrogen causes W atom displacements to switch from along the (110) surface directions to along the (100) surface directions, 10,14 to form a dimerlike structure. Again, there are two types of domains that are related by a rotation of 90° about the surface normal. They are equivalent and coexist with about equal probability on a flat W(100) surface. The degeneracy of the two types of domains possibly renders the diffusion isotropic when averaged over a macroscopic measurement length scale of micrometers, which is much...
larger than the microscopic domain size. However, the degeneracy of the two types of domains can be lifted by properly orienting and spacing surface steps. For example, a domain ratio of about 30:1 was reported for the \( p2mg \) structure on the clean surface as a result of miscutting the surface along the [110] direction by 3.25°, which resulted in a terrace width about 28 Å. The atomic displacements in the \( p2mg \) reconstructed structure are preferentially perpendicular to the step edges for this miscut. For the \( c2mm \) structure, miscutting a surface along the [100] direction, which results in steps along the surface [010] direction, can break the degeneracy and results in preferential domains with displacements of the W atoms along the step edge. A second purpose of our experiment will be to study the H diffusion anisotropy on single-type domains that are obtained on a stepped W(100) surface.

The organization of the paper is as follows. We will first describe how the experiment is performed. Then, the results of H diffusion on a stepped W(100) surface together with those for a flat surface will be presented. The results on the flat surface have been published in a recent Letter and are included here for comparison. Diffusion both perpendicular and parallel to steps will be covered. In the Discussion section, we will discuss the step effect on the diffusion anomaly near the W(100) reconstruction phase transition, followed by a discussion of the diffusion across and along steps. The anisotropy effect of H diffusion on a single-domain phase will also be briefly discussed. The experimental results show that the step effects on both the diffusion anomaly and diffusion anisotropy are small. This nontrivial observation by itself is already interesting and will stimulate further theoretical study.

II. EXPERIMENTAL DETAILS

The experiments were carried out in a stainless steel UHV system with a base pressure of \( \sim 2 \times 10^{-10} \) torr. Two single-crystal W(100) samples were used: one was cut and mechanically polished to within 0.1° of the (100) plane; the other was cut 2.5° off the (100) plane in the [010] direction. A 95% W–5% Re/74% W–26% Re thermocouple was spot welded to the side of the crystals to measure the sample temperature. Each sample was first cleaned by cycles of annealing in oxygen at 1480 K and 2 \( \times 10^{-7} \) torr for 20 min, and subsequent flashing to 2000 K until Auger spectra showed no detectable surface impurities. The total time of oxygen treatment was more than 5 h. Observations of the \( c(2 \times 2) \) LEED patterns of the reconstructed surface at low temperatures also indicated that the sample was clean and well ordered. On the stepped surface, the integer LEED spots were elongated, with the amount of elongation quantitatively confirming the step density introduced by the miscut. After the sample was cleaned of bulk contaminants, the surface was routinely cleaned by high-temperature (2000-K) flashing only prior to each individual diffusion run. The sample could be cooled by liquid nitrogen (LN) from 2000 to 90 K in about 3 min. Adsorption of H on the W(100) surface was carried out at approximately 90 K by leaking H\(_2\) gas into the chamber. The H coverage was calibrated by thermal desorption spectroscopy (TDS) on the flat surface. Because the W atoms are known to be immobile at temperatures below 200 K, the sample was slowly heated to room temperature and held for about 2 min to equilibrate the surface. It was known from other studies that hydrogen adsorption is dissociative and that hydrogen atoms adsorb on the bridge sites between W atoms at all coverages. The saturation coverage of H on W(100) is two monolayers (2 \( \times 10^{15} \) adatoms/cm\(^2\)).

The diffusion coefficient \( D \) was measured using a linear optical diffraction technique, which has several advantages: (a) a wide dynamic range for measuring the diffusion coefficient, (b) a high adsorbate coverage sensitivity, and (c) an intrinsic capability for measuring diffusion anisotropy. In this method, first, an adsorbate grating is created by laser-induced thermal desorption with two interfering pulsed laser beams. In our experiment, we used a grating period of 3.9 or 6.7 \( \mu \)m to measure the diffusion coefficient \( D \) slower or faster than \( \sim 10^{-9} \) cm\(^2\)/s, respectively, and a shallow coverage modulation of about 0.03 ML to ensure the diffusion coefficient \( D \) in the relevant coverage range to be well-approximated as a constant. Once such a grating is produced, the first-order diffraction of a He-Ne laser beam with polarization modulation is used to probe the smearing of the grating induced by surface diffusion. The diffraction signal decays exponentially according to

\[
S = S(0) \exp(-t/\tau), \quad \tau = s^2/8\pi^2D.
\]

Here, \( D \) is the chemical diffusion coefficient and \( s \) is the grating period. To determine \( D \), only the decay time constant and the grating period were needed. The detailed shape of the grating would not affect \( D \). In our study, diffusion measurements were carried out between 200 and 450 K at two different H coverages (1.2 and 0.17 ML). On the stepped surface, diffusion both perpendicular and parallel to the steps was measured with the adsorbate grating appropriately oriented.

III. RESULTS

The diffusion coefficients at 1.2-ML H coverage on flat and stepped surfaces are depicted in the Arrhenius plot in Fig. 1. On the flat surface, the diffusion coefficient can be fitted by the simple Arrhenius law \( D = D_0 \exp(-E_{\text{diff}}/k_B T) \) over the entire temperature range, with a prefactor \( D_0 \sim 10^{-5(2.2\pm0.2)} \) cm\(^2\)/s and a diffusion activation energy \( E_{\text{diff}} \sim 10.5\pm0.3 \) kcal/mol. On the highly stepped surface, it is seen that the steps, even at a high density of 1 step/12 terrace rows, do not affect the H diffusion to a large extent. Within the experimental error, diffusion parallel to steps is essentially the same as that on the flat surface. Diffusion perpendicular to steps is slightly slower than that on the flat surface, perhaps reflecting a somewhat larger diffusion barrier across the steps, as will be discussed later.

The diffusion data at 0.17-ML H coverage on the flat and stepped W(100) surfaces are shown in Fig. 2. \( D_0 \sim 10^{-6(1.2\pm0.4)} \) cm\(^2\)/s and \( E_{\text{diff}} \sim 11.7\pm0.5 \) kcal/mol are obtained from fitting the diffusion data on the flat surface below 325 K. On both the flat and stepped surfaces, the diffusion
coefficient appears to have a pronounced dip around 355 K. The dip on the flat surface was found to correlate with $T_C$ determined from LEED measurements. Within experimental error, the behavior of the dip is identical on the flat and stepped surfaces. While the diffusion coefficient parallel to steps is basically the same as that on the flat surface, the diffusion coefficient perpendicular to steps deviates somewhat from that on the flat surface and is most likely related to the Schwoebel-Ehrlich barrier effect.

For completeness, the diffusion data at 0.08- and 0.31-ML H coverages on the flat W(100) surface (the former were reported in our previous Letter\(^{17}\)), are included in Fig. 3. The behavior of $D$ is similar to that for 0.17 ML, except that the dip of $D$ occurs at a temperature near 290 K at 0.08 ML and near 365 K at 0.31 ML, corresponding to the different $T_C$ at these coverages. The diffusion data in the low-temperature range can be fitted by $D = D_0 \exp(-E_{\text{diff}}/k_B T)$, $D_0 \sim 10^{-1.9 \pm 1.2} \text{ cm}^2/\text{s}$ and $E_{\text{diff}} \sim 10.8 \pm 1.4 \text{ kcal/mol}$ at 0.08 ML and $D_0 \sim 10^{-2.8 \pm 0.6} \text{ cm}^2/\text{s}$ and $E_{\text{diff}} \sim 9.9 \pm 0.5 \text{ kcal/mol}$ at 0.31 ML.

### IV. DISCUSSION

#### A. Phase transition effect

While the step density on the flat surface could be higher that determined by the miscut ($<0.1^\circ$) due to a possible hill-and-valley morphology developed through repetitive high temperature annealing, the size of the LEED spots on the flat surface remains limited by instrument resolution and the corresponding step density cannot be deduced. On the stepped surface, however, the integer LEED spots are clearly elongated, with the amount of elongation consistent with a terrace size of $\sim 36$ Å. The size of the LEED spot in the orthogonal direction is the same as that on the flat surface, with an instrument-limited dimension at least three times smaller than that in the elongated direction. Thus, the step density on the stepped surface must at least be three times higher than that on the flat surface, limited by the ability of measuring the true step density on the latter. This independent evidence on different step densities for the flat and stepped surfaces should validate our following discussion on the step effects.

As shown in Fig. 1, no anomalous behavior of $D$ is observed both on the flat and stepped surfaces at 1.2-ML H coverage. This is consistent with the fact that there is no substrate phase transition over the entire temperature range at this coverage.\(^{39}\) At H coverages of 0.08, 0.17, and 0.31 ML, however, an anomalous decrease of $D$ with increasing temperature is obvious as shown in Figs. 2 and 3. Our LEED...
measurements\textsuperscript{17} and the previous study of the H/W(100) phase diagram\textsuperscript{10} indicate that the dip of the diffusion coefficient occurs at a temperature that is consistent with the coverage-dependent substrate phase transition temperature $T_C$.

The origin of the anomalous diffusion behavior near the substrate phase transition temperature has been discussed in our previous Letter.\textsuperscript{17} Near the reconstructive phase transition of W(100), the vibrational modes near the critical wave vector $q_0 = (\pi/a, \pi/a)$ soften and become overdamped as $T$ approaches $T_C$. The anomalous temperature dependence of the substrate phonons leads to critical singularities in the dynamic structure factor $S(q, \omega)$ near $T_C$ and ultimately to an anomaly in the temperature dependence of the friction acting on the H adatoms. Near $T_C$, $S(q, \omega)$ obeys the scaling form $\xi_c^{x-\eta} g_{\gamma}(q - q_0, \epsilon, \omega \xi_c^z)$, where $g_{\gamma}$ is a scaling function, $\xi_c \propto [T/T_C - 1]^{-\nu}$ is the divergent correlation length, $\gamma$ is the susceptibility exponent, and $z$ is the dynamical critical exponent.\textsuperscript{3} In the Markovian limit of instantaneous damping, the frictional damping $\eta$ is simply determined by the average of $S(q, \omega = 0)$ over a range of $q$ determined by the coupling potential of the H adatom to the substrate.\textsuperscript{1,3} This leads then to a $\eta$ diverging as $[T - T_C]^{-\nu}$ with $x = \nu(z - d) + \gamma$ (in $d$ dimensions) and $D \sim \eta^{-1}$ vanishing as $T$ approaches $T_C$.

The simple argument presented above only applies when $T$ is not too close to $T_C$. Two additional factors need to be considered, which eliminate the infinite divergence of the frictional damping and result in a finite dip in $D$ instead of it vanishing at $T_C$. The first is the breakdown of the Markovian approximation that the frictional damping is proportional to the instantaneous velocity.\textsuperscript{40,41} As one approaches $T_C$, the dynamic structure factor $S(q, \omega)$ gets narrower and stronger and develops into a “central peak” structure.\textsuperscript{3} When the width of this central peak gets narrower than the characteristic frequency for the motion of the H adatom, the time scale of the relevant substrate excitations is longer than the time scale of the motion of the hydrogen adatom. At this point, the instantaneous damping picture breaks down and the divergence in the frictional damping is cut off. The second factor that suppresses divergence is sample imperfection such as finite size or a finite step density. Since the phases of the reconstruction for the domains on each terrace are uncorrelated, divergence of the correlation length $\xi$ is cut off when it reaches the average size of a reconstructed domain.\textsuperscript{42} Thus the finite frequency of the H adatom acts as a temporal cutoff, while the finite size of the reconstructed domain acts as a spatial cutoff that can be described by appropriate finite-size scaling theory.\textsuperscript{3,41} The rounding of the anomaly and the elimination of the true divergence of the friction are due to whichever effect that first becomes dominant as one approaches $T_C$. For the “flat” surface, the terrace size is at least 100 Å and it is most likely that the non-Markovian effect, i.e., the temporal cutoff, is responsible for the rounding of the anomaly. On the stepped surface, the average size of the reconstructed domains is even smaller than the average terrace width since experimentally, there is evidence that the W(100) reconstruction may be inhibited near step edges at low temperatures.\textsuperscript{13,24,25,27,43,44} The degree of the inhibition is described by the inhibition range. The most reliable measurements of this quantity obtained from LEED spot profile analysis\textsuperscript{24,25,27} indicate that both the clean and H-induced structures are inhibited over a range that does not exceed 8 Å in the proximity of steps along the surface [110] or [010] directions. The 2.5° miscut of our sample implies a terrace width of only about 36 Å. Thus the size of the reconstructed domain is of order of 28 Å. The observation of $\frac{1}{2T_C}$ LEED spots and the dip in the plot of $D$ vs $1/T$ at 0.17 ML (Fig. 2) indicates that the size of the 2.5° miscut W(100) substrate still undergoes a reconstructive phase transition. Since there is little difference in the dip of $D$ between that on the flat surface and that on the stepped surface, this implies that on the stepped surface, either the temporal cutoff is still responsible for the rounding of the anomaly, or else the temporal and spatial cutoff occur at a similar temperature region around $T_C$. At this point, we do not have enough information on the detailed parameters of the system to distinguish between these two scenarios.

### B. Schwoebel-Ehrlich barrier and effective coverage

It is well known that steps may affect diffusion perpendicular to steps through a Schwoebel-Ehrlich barrier\textsuperscript{45–47} and/or an additional trapping potential well at the step edges.\textsuperscript{18,20} Diffusion parallel to steps may be affected by a fast channel along the steps.\textsuperscript{18,20} Both theoretical and experimental studies have shown these effects.

Diffusion perpendicular to steps must be subject to several processes in sequence. They are diffusion over a terrace, over a step edge barrier (i.e., Schwoebel-Ehrlich barrier) to reach a trapping site, and hopping out of the trapping site to the adjacent terrace site. The diffusion time for these sequential processes is additive. Based on the lattice gas model with nearest-neighbor hopping, Merikoski and Ying have analyzed these processes in detail.\textsuperscript{18} For a Langmuir gas with no adsorbate-adsorbate interaction except site blocking, the overall diffusion coefficient perpendicular to steps $D_s$ in the limit of no additional trapping barrier ($\Delta E_B \sim 0$) at the step sites is given by\textsuperscript{18}

$$\frac{1}{D_s} = \frac{L - 2}{L} \frac{1}{\nu_v a^2 e^{-E_s/k_B T}} + \frac{1}{L} \frac{1}{\nu_{eff} a^2 e^{-\left(E_t + \Delta E_s\right)/k_B T}}, \quad (2)$$

where

$$\nu_{eff} = \frac{\nu_v}{1 + \left(\nu_v/\nu_B\right) e^{-\Delta E_s/k_B T}},$$

$L$ is the terrace width in units of the lattice constant $a$, $\nu_v$ and $E_s$ are the attempt frequency and activation energy on a terrace, respectively, $\nu_v$ and $\nu_B$ are the attempt frequencies over the Schwoebel-Ehrlich barrier and out of the trapping well, respectively, and $\Delta E_s$ is the height of the Schwoebel-Ehrlich barrier. Thus, $D_s = \nu_{eff} a^2 e^{-\left(E_t + \Delta E_s\right)/k_B T}$ is the diffusion coefficient and $D_s = \nu_v a^2 e^{-E_s/k_B T}$ is the diffusion coefficient over steps.

We fit our data for 1.2- and 0.17-ML H coverage on the stepped surface with Eq. (2) assuming the values of $D_{0,H} = \nu_v a^2$ and $E_t$ are the same as those on the flat surface. Here, we limit the data range for 0.17 ML only in the low-
temperature region (< 325 K). The fitting curves are shown in Figs. 1 and 2 as solid lines. The fitting values of $D_{0,i} = \nu_{\text{eff}} a^2$ and $\Delta E_{S,i}$ are listed in Table I. We find $\Delta E_{S}$ changes a little for different H coverages. The values determined here for H/W(100) are similar to the value $\Delta E_{S} = 2.7$ kcal/mol for H diffusion on a step Ru(001) surface determined by laser-induced thermal desorption techniques (LITD) and by assuming a negligible $\Delta E_{B}$.  

For diffusion parallel to steps, the diffusion activation energy $E_{2}$ along the step edge may be different from that on terrace, which results in a different diffusion rate along the steps. In the previous theoretical studies of Merikoski and Ying, it was shown that the diffusion coefficient parallel to the step for the condition $\Delta E_{B} \approx 0$ is given by $\nu_{\text{eff}} a^2$  

$$D_{\parallel} = \frac{L-1}{L} D_{\parallel}^0 + \frac{1}{L} D_{\parallel}^i,$$  

(3)

where $D_{\parallel} = D_{0,2} \exp(-E_{2}/k_{B}T)$ is the diffusion coefficient along the step edge. In both Figs. 1 and 2, we observe that the diffusion coefficient parallel to steps is basically the same as that on the terrace within the experimental error in the entire temperature range for 1.2- and 0.17-ML H coverages. The values determined here for H/W(100) are similar to the value $\Delta E_{S} = 2.7$ kcal/mol for H diffusion on a step Ru(001) surface determined by laser-induced thermal desorption techniques (LITD) and by assuming a negligible $\Delta E_{B}$.  

The present results are in strong contrast to CO diffusion on vicinal Pt(111) surfaces that were studied by our group previously. While diffusion of CO perpendicular to steps is impeded by a strong additional trapping potential at the step sites ($\Delta E_{B} \approx 7$ kcal/mol), diffusion parallel to steps shows an enhancement over terrace diffusion in various ways. At high temperatures, all three steps of types A, B, and AB, show the contribution to mass transport by the fast step-edge diffusion. At low temperatures, a third channel, which exists only for A- and AB-type steps, also contributes to diffusion enhancement. The enhancement of the diffusion coefficient due to steps is more than two orders of magnitude. In the present system of H/W(100), such step effects are absent or below the detection limit. The above comparison indicates that effects of the steps on the magnitude of surface diffusion depend strongly on the particular systems being studied. No universal rules about the step barriers across and along the steps can be deduced from the information available so far.

We must add a note of caution about the fits to the experimental data in Figs. 1 and 2. Aside from the magnitude of the Schwoebel-Erlich barrier and the fact that there are no fast channels parallel to the steps, the details of the fits cannot be taken too seriously because of the simplicity of the model. In particular, the existence of an inhibition range has not been taken into account explicitly. It has been shown by Lau and Ying that on a reconstructed surface, the binding energy of the adsorption sites increases monotonically with the amplitude of the reconstruction. Thus, instead of additional trapping at the step edge, the hydrogen adatoms would actually stay away from the inhibition region and only adsorb and diffuse in the reconstructed domain. Depending on the inhibition range and hence the size of the reconstructed domain, the actual H coverage on the reconstructed domain can be significantly higher than the nominal coverage determined by the exposure over the entire surface. This could explain the fact that when we examined the LEED patterns at nominal 0.17-ML coverage on the stepped surface, we found that each ($\frac{1}{2} \frac{1}{2}$) spot splits into two circular spots, perpendicular to the step directions. The splitting indicates the formation of an incommensurate structure that is known to occur on a flat surface at a coverage of about 0.3 ML, much higher than the nominal coverage of 0.17 ML averaged over the whole surface.

### Table I. The fitted values of the prefactor and diffusion activation energy at 1.2- and 0.17-ML H coverages.

<table>
<thead>
<tr>
<th>Coverage (ML)</th>
<th>$D_{0,i}$ (cm$^2$/s)</th>
<th>$E_{i}$ (kcal/mol)</th>
<th>$D_{0,i}$ (cm$^2$/s)</th>
<th>$\Delta E_{S}$ (kcal/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.2</td>
<td>$10^{-2.2 \pm 0.2}$</td>
<td>10.5 $\pm$ 0.3</td>
<td>$10^{-1.8 \pm 0.6}$</td>
<td>2.2 $\pm$ 0.8</td>
</tr>
<tr>
<td>0.17</td>
<td>$10^{-1.2 \pm 0.4}$</td>
<td>11.7 $\pm$ 0.5</td>
<td>$10^{-0.1 \pm 2.4}$</td>
<td>2.8 $\pm$ 3.2</td>
</tr>
</tbody>
</table>

C. Anisotropy

The adsorption of H on W(100) induces a substrate reconstruction phase transition from (1 $\times$ 1) at high temperatures to c(2 $\times$ 2) at low temperatures. The c2mm symmetry of the c(2 $\times$ 2) structure at a H coverage between $\approx 0.12$ and $\approx 0.30$ ML has a twofold rotation symmetry in contrast to a fourfold rotation symmetry for the unreconstructed W(100) surface. Thus, diffusion on the c(2 $\times$ 2) surface may be expected to be anisotropic. On a flat reconstructed W(100) surface, the orthogonal degenerate domains will be present with equal probability. Therefore, the anisotropy of surface diffusion is averaged out in a macroscopic measurement that probes many domains. In contrast, the reconstruction is preferentially oriented with atomic displacements along steps on a stepped surface, which causes an unequal population of the two orthogonal domains. A strong domain preference was also observed here. From the ($\frac{1}{2} \frac{1}{2}$) spot splitting (only two instead of four spots) on the stepped surface, it clearly shows single domain orientation. Thus, the diffusion anisotropy is expected to be observable on the stepped surface. This domain-induced diffusion anisotropy will be superimposed on the anisotropy due to steps. The diffusion anisotropy due to steps alone at 1.2-ML coverage is clearly shown in Fig. 1 and the amount of anisotropy is small. In Fig. 2, the fact that the diffusion results parallel to steps at 0.17-ML H coverage are nearly identical to the flat surface ones, which is a statistical average of diffusion over the two orthogonal domains with equal populations, indicates that the anticipated anisotropy is small. For diffusion perpendicular to steps, the diffusion data can be fitted well by Eq. (2) when we set $D_{0,i}$ and $E_{i}$ equal to the corresponding values on the flat surface, further supporting the notion that the diffusion anisotropy on the c(2 $\times$ 2) domain is small. However, this conclusion is subject to the uncertainty of the incommensurate structure which has a periodicity of $\approx 20$ Å along the steps determined from the splitting of the half order spots. How the domain wall affects diffusion anisotropy is not yet clear.
V. CONCLUSION

We have investigated the step effect on H diffusion near the W(100) reconstructive phase transition. The anomalously high diffusion coefficient \( D \) observed in our Arrhenius plot, due to the diverging friction damping near the phase transition, is not significantly affected by the steps.

From this we conclude that the reconstruction phase transition still occurs on the surface with high step densities. The divergence of the length and time scale near \( T_C \) is cut off due to either non-Markovian effects as in the case of the flat surface or the finite size of the reconstructed domain.

For diffusion parallel and perpendicular to the steps, dips of \( D \) were observed similar to that on a flat W(100) surface. The step introduces a Schwoebel-Ehrlich barrier, about 2.2 kcal/mol for 1.2-ML and 2.8 kcal/mol for 0.17-ML H coverages, to influence H diffusion perpendicular to the steps. Diffusion parallel to steps has not been affected by the steps at all. From the measurements, we also conclude that the diffusion anisotropy on the \( c(2 \times 2) \) domain is small.

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