

Structure of the α -Al₂O₃(0001) surface from low-energy electron diffraction: Al termination and evidence for anomalously large thermal vibrations

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We use dynamical low-energy electron diffraction (LEED) to determine the surface structure of α -Al₂O₃(0001). Sapphire surfaces are prepared in three different ways, and the diffraction results are analyzed using an exhaustive search of possible models. For all sample processing conditions, the clearly favored structure has a single Al layer termination and a large first interlayer contraction. In addition, we find that the aluminum atoms at the surface have unusually large vibrational amplitudes at room temperature, suggestive of an anharmonic vibrational mode.

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I. INTRODUCTION

Alumina (i.e., aluminum oxide and its hydrates) is widely used in aluminum production, ceramics, and catalysts, and occurs on the surface of oxidized aluminum alloys.^{1,2} Being the simplest and the only thermodynamically stable aluminum oxide,² α -Al₂O₃ is a prototype for understanding metal oxides. Because of its importance, numerous experimental³⁻⁸ and theoretical⁹⁻¹⁶ investigations of its surfaces have been performed. Nonetheless, a most basic property of its simplest clean surface, namely the structure of α -Al₂O₃(0001), remains controversial.

Compared to monoatomic materials, determining the surface structure of a compound has several additional complicating factors. First, a compound may terminate along different ideal planes, giving inequivalent surface structures. For α -Al₂O₃(0001), three different (0001)-plane terminations exist: a single Al layer (A11), an oxygen layer (O1), and a double Al layer (A12), where we denote the different surfaces by the terminating layer(s) as labeled in Fig. 1. First-principles calculations predict an A11 termination with the first interlayer spacing being greatly contracted ($\sim 85\%$) relative to the bulk.¹²⁻¹⁶ X-ray-diffraction⁶ and ion-scattering⁵ experiments concluded that the α -Al₂O₃(0001) surface is A11 terminated. However, the models considered in these investigations were limited to the ideal (0001) plane surfaces, i.e., the A11, O1, and A12 surfaces. Additionally, these experiments found a first interlayer contraction that is $\sim 35\%$ smaller than that predicted by theory. Based on their calculations of the TiO₂(110) surface, Harrison *et al.* suggested that the difference in interlayer spacings determined by zero-temperature calculations and room-temperature experiments may be explained by the existence of large, anharmonic vibrations.¹⁷ Since such vibrations cannot be accurately modeled in low-energy electron

diffraction (LEED) calculations using an isotropic Debye-Waller approximation, evaluating their existence requires the use of more complex models.¹⁸⁻²⁰ Static disorder may also be present as pointed out by Gloege *et al.*²¹ in a surface X-ray diffraction (SXRD) study of the equivalent α -Cr₂O₃(0001) surface structure. Their analysis provides evidence that, at room temperature, the surface is terminated by a disordered arrangement of surface Cr⁺³, and is characterized by a 2/3 occupation probability of the top layer site (equivalent to A11) and a 1/3 occupation probability of the interstitial site between the first and second O⁻² layers. Although their results provide an explanation for the order-disorder and order-order phase transitions observed on this surface, the first interlayer distance is rather different from the values previously reported.

In addition to single-species termination, compound surfaces can potentially be phase-separated, i.e., consist of a thermodynamic equilibrium of domains having different stoichiometry or structure.^{12,13,15,22} For example, calculations suggested that under typical experimental conditions, the (0001) surface of the isostructural phase α -Fe₂O₃ is covered by two distinct domains, one terminated by Fe and one by O.²² Experimentally, the surface has been reported to consist either of two domains with different structure,²² or, inconsistently, as exclusively terminated by oxygen.²³ For α -Al₂O₃(0001), the three ideal bulk terminations have different stoichiometries at the surface. Therefore, their surface energies depend differently on the oxygen (alumina) chemical potential.^{12,13,15,16} Because the single-Al layer (A11) surface has the same stoichiometry as the bulk, its energy is independent of the aluminum or oxygen chemical potential, unlike the nonstoichiometric surfaces terminated by an oxygen layer or a double Al layer. This raises the possibility that the lowest-energy state, for a given chemical potential, is actually a phase-separated mixture of two different surface terminations. Therefore, the existence of a phase-separated

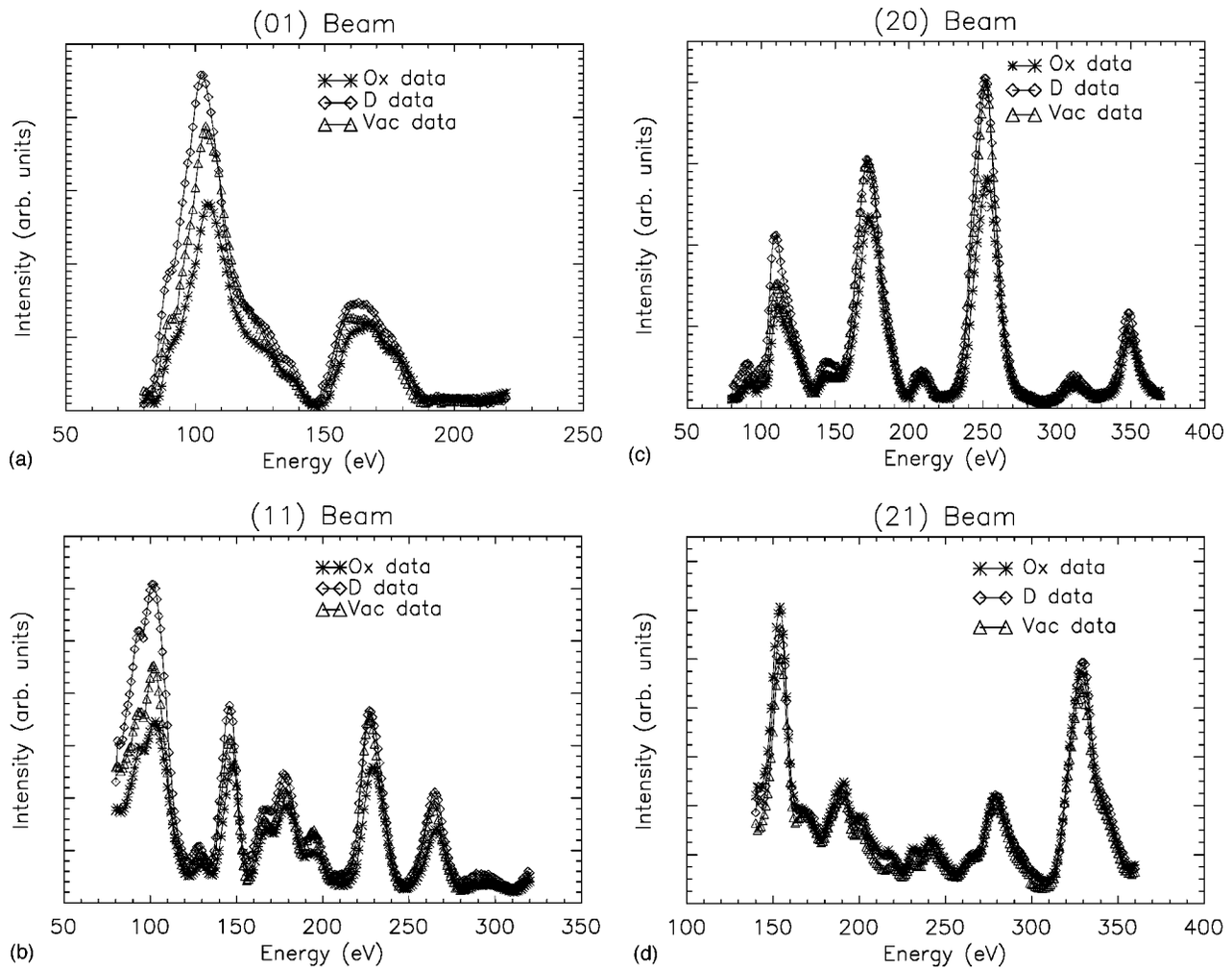


FIG. 2. Representative experimental LEED $I(V)$ curves for the three different sample preparation procedures of the α -Al₂O₃(0001) surface. The (i,j) notation gives the index of the diffraction spots.

moved at 650 °C using an atomic deuterium beam created by a commercial, neutralized, rf plasma discharge.³⁰ To investigate the sensitivity of the surface structure to processing conditions, we finished the processing in three very different ways: (1) turn off the atomic deuterium beam and cool from 650 °C in vacuum (“Vac” data), (2) cool to 200 °C before turning off the atomic deuterium beam (“D” data), and (3) turn off the atomic deuterium beam, heat for 5 min in 5×10^{-5} -Torr O₂, and then cool in vacuum (“Ox” data). All three procedures produced bright, sharp, 1×1 LEED patterns with a clear threefold symmetry.

The LEED data were acquired with the sample at room temperature using a high-sensitivity CCD camera and an automated data acquisition system. Nine inequivalent beams were recorded at normal incidence in the energy range of 80 to 370 eV (total range 2080 eV). After subtracting the background, defined by the average intensity in the pixels surrounding the region of integration for each beam, equivalent beams were averaged. The spectra were scaled to the incident electron current, which was set low enough to prevent nonlinear charging effects ($\sim 0.3 \mu A$). Because of the high quality of the data, no mathematical smoothing was required

prior to the analysis. Representative experimental LEED $I(V)$ curves (diffracted intensity as a function of electron energy) for the three different sample preparation procedures are shown in Fig. 2. Although the three sets are closely similar, we perform independent structural analyses using each data set individually.

III. MODEL DESCRIPTIONS

While the corundum structure of bulk α -Al₂O₃ has a rhombohedral symmetry, the atomic positions are usually given in terms of an hexagonal unit cell (Fig. 1).⁴ This unit cell can be viewed as a sequence of 12 Al layers, which are translationally equivalent to each other, and six O layers, with the O atoms in positions close to those of an hcp lattice. For the six O layers, alternate layers are translationally equivalent, and sequential layers are equivalent only after a translation and mirroring through a plane perpendicular to the surface. Any of these 18 layers may serve as a surface termination, and each of these surfaces has $p3$ symmetry, i.e., threefold rotational axes through the Al atoms and no mirror planes. However, while a surface that terminates in

TABLE I. List of the 21 different models within six different model classes considered in this study of the α -Al₂O₃(0001) surface. Each model is numbered and given a descriptive notation. When the model considers more than one domain, the experimental data were fit by varying the fractions of the domains, as suggested in the notation.

Model class	Model
Single-species bulk termination	1) Al1 2) Al2 3) Al3 4) O1 5) O2 6) Al1-O \equiv "hydroxyl"
Single-species termination with stacking fault	7) Terminated by Al1 on the <i>b</i> site 8) Terminated by Al1 on the <i>a</i> site 9) Terminated by O1 on the <i>C</i> site 10) O1 on the <i>C</i> site, terminated by Al1 above "open" sites 11) O1 on the <i>C</i> site, terminated by Al1 above O2 sites
Single-species termination with diffractationally inequivalent domains	12) Al1 + Al3 \equiv x Al1 + (1- x)Al3 13) O1 + O2 \equiv x O1 + (1- x)O2 14) Alm \equiv x Al1 + (1- x)Al3 same relative positions each domain 15) Om \equiv x O1 + (1- x)O2 same relative position each domain
Mixed-species termination	16) Al1 + O1 \equiv x Al1 + (1- x)O1 17) Al1 + O2 \equiv x Al1 + (1- x)O2 18) O1 + Al3 \equiv x O1 + (1- x)Al3 19) O1 + Al2 \equiv x O1 + (1- x)Al2
Mixed-species termination with diffractationally inequivalent domains	20) (Alm) + (Om) \equiv x (Al1 + Al3) + (1- x)(O1 + O2) same relative positions each domain
Split position	21) Al1-split \equiv Al1 + Al1
Disorder model	22) Al1 disorder \equiv x Al1 + (1- x)Al in interstitial state

layers Al1-O1-... is energetically equivalent to one that terminates in layers Al3-O2-..., through a symmetry transformation, the mirror-symmetry relationship of the adjacent O layers results in these terminations being inequivalent from the point of view of diffraction. That is, separate regions of Al1 termination and Al3 termination could coexist as energetically equivalent but diffractationally inequivalent terraces, and these must be averaged over to correctly model a terraced surface.

The simplest models used to analyze our LEED $I(V)$ data were the ideal planar cleavages: models Al1, O1, and Al2 (see Table I). A closely related, but nonideal, model consists of an O atom on top of each surface Al atom of the Al1 model. Since the scattering power of hydrogen is small enough to be neglected, this model represents a type of

water- or hydroxyl-covered surface. The next level of complexity involves surfaces that terminate in a single species (i.e., Al atoms or O atoms) but contain diffractationally inequivalent domains. That is, in these models, the surface consists of the two distinct terrace types (separated by a $c/6$ length along the c axis). In all of these models (Al1 + Al3, O1 + O2, Alm, and Om), the fractional coverage of each terrace type was treated as a fitting parameter. For the Alm and Om models, the atoms in the "mirrored" (i.e., $c/6$ separated) domains were constrained to have the same relative positions (i.e., mirrored domains were kept identical). In the Al1 + Al3 and O1 + O2 models, the atoms in the diffractationally inequivalent terraces (i.e., Al1 and Al3; O1 and O2) were not constrained to have the same relative positions.

We also considered mixed-species models, i.e., surfaces

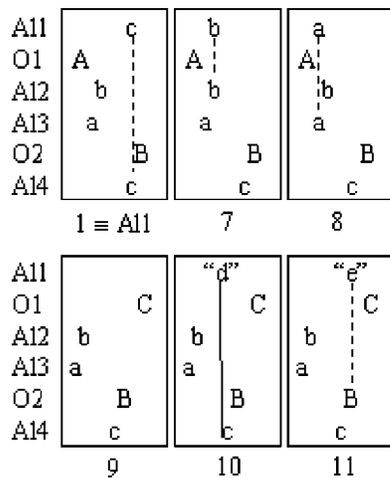


FIG. 3. Schematic illustrations showing the stacking sequence perpendicular to the surface of five stacking-fault models consistent with the observed $p3$ symmetry. The dashed lines connect atoms that lie on top of each other. Model 1 (the “A11” model, upper left) maintains the bulk stacking (see Fig. 1). In models 7 and 8, the topmost Al layer is shifted to lie above the Al atoms in the A12 and A13 layers, respectively. Models 9–11 consider a stacking fault such that the O1 layer is shifted to give an fcc stacking (ABC) to the topmost three O layers. In model 10, the A11 layer sits in the three-fold hollow sites below which no atoms in the bulk structure occur. In model 11, the A11 layer sits in the three-fold hollow sites above the O2 layer.

having regions terminated by oxygen atoms and regions terminated by aluminum atoms. In four of these models (A11 + O1, A11 + O2, O1 + A13, and O1 + A12), the fractional coverage of the aluminum and oxygen-terminated domains, and the atomic positions on each domain were independently varied. Finally, we considered a mixed-species model that also had diffractationally inequivalent steps, (Alm) + (Om). The surface fraction occupied by the two inequivalent Al-terminated domains (A11 and A13) and the fraction occupied by the two inequivalent O-terminated domains (O1 and O2) were varied. However, to limit the number of adjustable parameters in this “mixed-mirrored surface,” the atomic positions within both O-terminated domains and within both Al-terminated domains were constrained to be the same and the two O-terminated domains had equal abundance, as did the two Al-terminated domains (i.e., A11:A13 = O1:O2 = 50:50).

All of the previously discussed models are derived by cleaving the bulk structure along appropriate plane(s). However, it is possible that the surface differs from the usual bulk stacking sequence yet has the observed surface symmetry. Therefore, we considered five models that have stacking faults in the topmost one or two layers. Figure 1 shows the stacking of the Al and O layers with the usual notation associated with close-packed structures. The oxygen sublattice follows hcp-type packing (ABAB . . .) while the Al sublattice follows fcc-type packing (cbacba . . .). As illustrated in Fig. 3, the two simplest stacking-fault structures involve rigidly shifting the top Al layer in the A11 model from the c site (which is not occupied by the two Al layers located

between layers O1 and O2) to either the a site or the b site (both of which are occupied by Al atoms located between layers O1 and O2). The second type of stacking fault involves shifting the O1 layer from the B site to the C site, along with removing the distortion of the layer such that it has perfect hexagonal symmetry. The upper three oxygen layers then have fcc-type stacking, as in the cubic phase γ -Al₂O₃. In addition, the aluminum atoms in the A12 and A13 layers occupy tetrahedral sites, unlike the exclusive octahedral occupancy of α -Al₂O₃. Three different structures follow from the O stacking fault (Fig. 3), namely, terminating in the O1 layer (i.e., the A11 layer is absent, model 9), placing the A11 layer in the threefold hollow sites directly below which there are no atoms (model 10), and placing the A11 layer in the three-fold hollow sites directly above the O atoms of the O2 layer (model 11).

We have also considered the possibility that the surface Al atoms in the theoretically favored A11 model have anisotropic or unusual vibrations that cannot be correctly described using the isotropic Debye-Waller factor to which the standard LEED calculations are limited. If this is the case, the A11 model will result in a poor fit to the data even if the surface is in fact terminated by a single Al layer. To investigate this possibility, we modeled the surface with the well-established “split-position” technique,^{18–20} by constructing an equal mixture of two identical Al-terminated domains in which the topmost interlayer spacing is allowed to relax independently. (This model allows the split atoms to be half the time in one position and half the time in the other position, thereby simply representing a large vibrational amplitude perpendicular to the surface. LEED is less sensitive to vibrations parallel to the surface, so that it is not useful to try to refine that aspect further.) If the vibrational amplitude perpendicular to the surface is indeed too large to be correctly modeled by an isotropic Debye-Waller factor, the spacing between the split surface atoms in the two domains will increase, while the position of all other atoms in the model will be very similar.

Finally, we have considered a model which takes into account the existence of static disorder as proposed by Gloege *et al.*²¹ In this model the surface is terminated by an A11 plane but some of the Al atoms are located in an interstitial site between the O1 and O2 planes.

IV. LEED CALCULATIONAL TECHNIQUE

The LEED analysis applied the familiar method of symmetrized automated tensor LEED,³¹ which has been used, for example, to study the complex oxide Fe₃O₄(111).^{32,33} Although α -Al₂O₃ is an ionic compound, neutral scattering phase shifts were used. It is well known that the structural fit depends very little on those nonstructural parameters, provided their values are reasonable. This point was explicitly checked by Barbieri *et al.*³² in a surface structural analysis of Fe₃O₄(111). To take into account the difference in the ionic radii, we assumed that the oxygen muffin-tin radius (r_{muf}^O) was twice the Al muffin-tin radius ($r_{muf}^O = 2r_{muf}^{Al}$). The muffin-tin potential and the phase shifts were calculated using the Barbieri/Van Hove Phase Shift Package.³⁴ In particu-

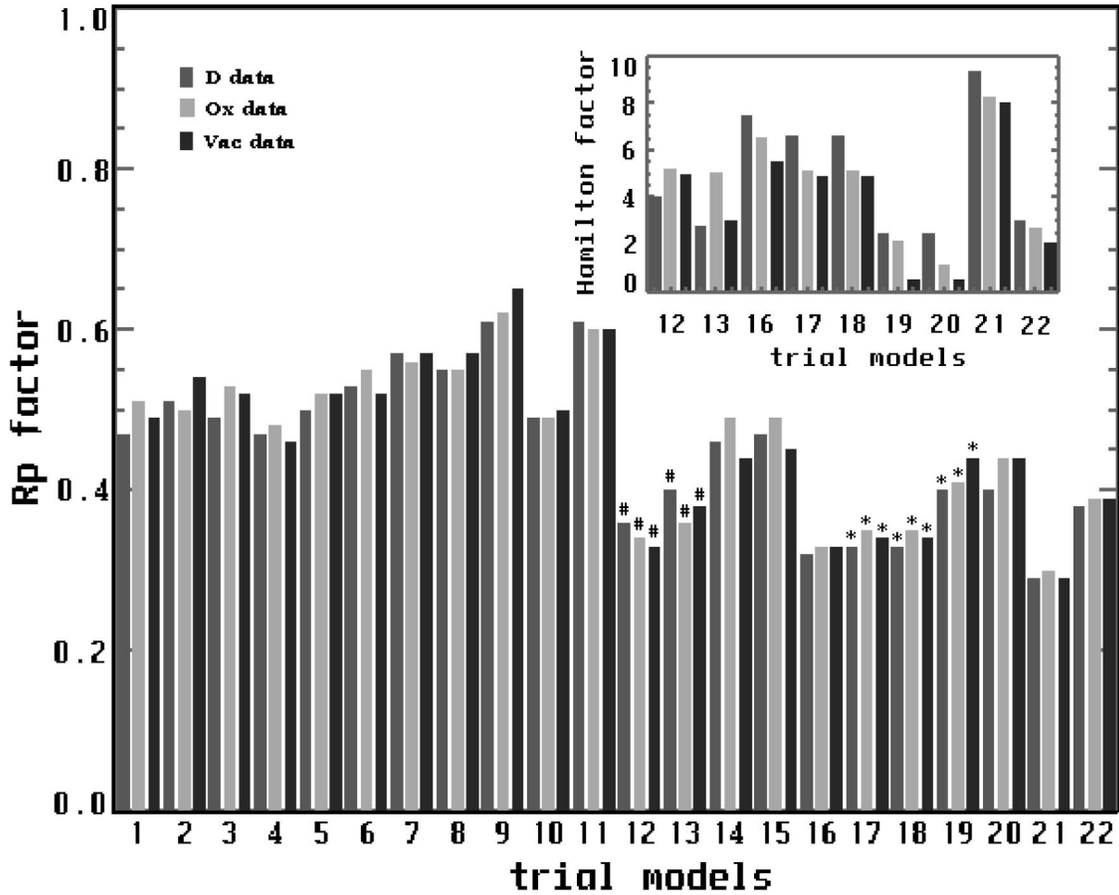


FIG. 4. Pendry R factors (R_p) for the three α - $\text{Al}_2\text{O}_3(0001)$ sample-preparation methods and the 21 models tested. The insert gives the Hamilton ratios (H_r) for the models having mixtures of two or more domain types (see Table I). The models are numbered along the horizontal axis as 1–5, ideal (0001) terminations; 6, water-covered surface; 7–11, stacking faults; 12–15, single-species mirrored surfaces; 16–20, mixed-species terminations; 21, split-position model, and 22, disorder model. The stars in the figure note models whose optimized structures are close to those of the AlI + O1 model, despite the fact that the starting configurations were very different. The pound signs in the figure note models that resulted in large, non-physical bond-lengths (e.g., a top layer expansion of 90%).

lar, a self-consistent Dirac-Fock approach was used to compute the self-consistent atomic orbitals for each element. The muffin-tin potential was then computed following Mattheiss' prescription, and the relativistic phase shifts were evaluated by numerical integration of the Dirac equation.

In all of the models we tested, the atoms were allowed to fully relax down to a depth of seven layers under the provision that they maintain the observed $p3$ symmetry. Under this constraint, atoms that lie along the axis passing through the bulk Al atoms can only relax perpendicular to the surface, while other atoms could also relax laterally. In all cases where mixed domains were considered, the calculated spectra were derived from incoherently summing over the different terraces. The value of the imaginary part of the potential was held constant at -6.0 eV for all tested models.

The goodness of the fits to the various structural models is described in terms of the Pendry R factor (R_p).³⁵ One additional criterion, known in x-ray crystallography as the Hamilton-ratio test,^{36,37} is herein introduced into LEED to deal with variable numbers of fitparameters, as occurs when comparing a model that consists of a single structure with a model that consists of more than one structure. The Hamilton

ratio helps to distinguish real improvements in a fit due to choosing a better model, from artificial improvements due only to fitting more structural parameters. As long as the structural coordinates are otherwise reasonable, a large Hamilton ratio is indicative of real improvements. Adapted to the LEED case, the Hamilton ratio is defined as

$$H_r = \frac{(R_{large}^2 - R_{small}^2)(n - p_{large})}{R_{large}^2(p_{large} - p_{small})}, \quad (1)$$

where R_{small} and R_{large} are the R factors for the same model with the smaller (p_{small}) and larger (p_{large}) numbers of fitting parameters, and n is the number of diffraction peaks, a measure of the number of independent experimental bits of information. In a LEED $I(V)$ spectrum, the width of the dominant peak is about $4|V_{oi}|$, where V_{oi} is the inner potential. In addition, the $I(V)$ curves usually contain as many peaks as can possibly be fit into the available energy range. Therefore, we assume that the number of diffraction peaks n is reasonably estimated by the total energy range divided by the peak width. In our experience this formulation is appli-

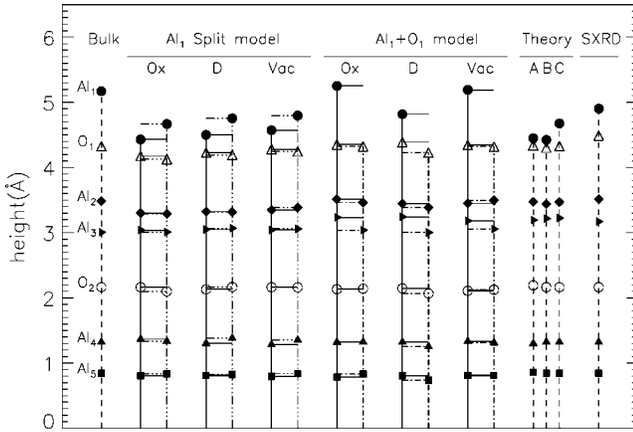


FIG. 5. Graphical representation of the atomic positions perpendicular to the surface in both terminations for the Al1-split and Al1+O1 models, compared to bulk values and to results from theory [A (Ref. 14), B (Ref. 11), and C (Ref. 12)] and x-ray diffraction (Ref. 6). Each vertical line represents one termination, and gives the optimized height of each atomic layer (labeled by individual layer-specific symbols) above the first fixed O layer (height 0 Å). Pairs of connected lines correspond to pairs of terminations that were optimized together, showing resulting height differences.

cable to all the various R factors commonly used in LEED, and the ratio should exceed 3 to indicate real improvements.

V. RESULTS

For each of the three different sample preparations, we have performed the most exhaustive structural examination of α -Al₂O₃(0001) to date by examining 22 different surface models within six distinct model classes (Table I). The results of the optimized fitting of the various structural models are summarized in Fig. 4 and Table II in terms of the Pendry R factor (R_P) and the Hamilton ratio (H_r). We are looking for structures with a low R_P , preferably lower by 20% than other structures, and with a relatively large Hamilton ratio, preferably larger than 3. Additionally, we need to exclude physically unrealistic structures, namely those that have unacceptable bond lengths; these structures are indicated by a pound sign in Fig. 4.³⁸ None of the ideal terminations (models 1–5) or the “hydroxyl” surface (model 6) adequately describes the data. The single-species models that included diffractationally inequivalent domains (models 12–15) also gave unacceptable R_P values (higher than 0.4) or unphysical

TABLE II. R_P factors, number of fitting parameters and Hamilton ratio for the mixed-species terminations, for the split model and for the disorder model. The R_{small} and the P_{small} values correspond to the best single-specie terminated model. In all cases the number of diffracted peaks n was 85.

Model	$R_P = R_{large}$	P_{large}	R_{small}	P_{small}	H_r
16					
‘Ox’	0.33	21	0.48	10	6.5
‘Vac’	0.33	21	0.46	10	5.5
‘D’	0.32	21	0.47	11	7.4
17					
‘Ox’	0.35	21	0.48	10	5.2
‘Vac’	0.34	21	0.46	10	4.9
‘D’	0.33	21	0.47	11	6.6
18					
‘Ox’	0.35	21	0.48	10	5.2
‘Vac’	0.34	21	0.46	10	4.9
‘D’	0.33	21	0.47	11	6.6
19					
‘Ox’	0.41	21	0.48	10	2.2
‘Vac’	0.44	21	0.46	10	0.5
‘D’	0.40	21	0.47	11	2.4
20					
‘Ox’	0.44	21	0.48	10	1.1
‘Vac’	0.44	21	0.46	10	0.5
‘D’	0.40	21	0.47	11	2.4
21					
‘Ox’	0.30	22	0.48	10	8.2
‘Vac’	0.29	22	0.46	10	8.0
‘D’	0.29	22	0.47	11	9.4
22					
‘Ox’	0.39	22	0.48	10	2.7
‘Vac’	0.39	22	0.46	10	2.1
‘D’	0.38	22	0.47	11	3.1

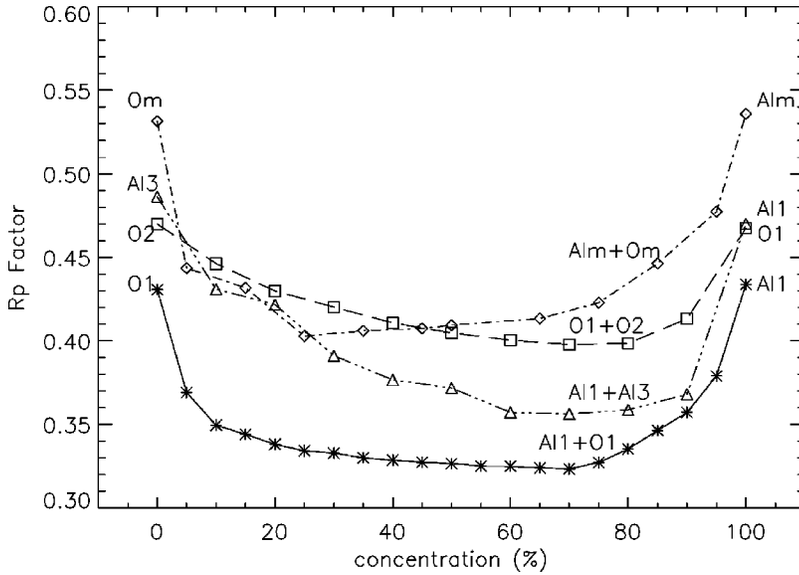


FIG. 6. The goodness of the model fits (Pendry R factor, R_p) for representative surface models having a mixture of domains with different terminations. On the left-hand axis (0%) and the right-hand axis (100%), the models have a single domain of the labeled type. In between, the surface is a mixture of the two terminations. The extreme insensitivity to the domain concentration suggests that the mixed-domain classes of models are inappropriate. While the results shown are from the D experimental data set, the other surface preparations gave similar results.

bond lengths. This fact establishes that the inadequacy of the aluminum-terminated surface model (Al1) is not simply due to the omission of the diffractationally inequivalent terraces. Clearly, additional effects beyond diffractationally inequivalent domains must be included to adequately model the LEED data. In models Al1 + Al3 and O1 + O2 (models 12

and 13), there are additional degrees of freedom available because the relative atomic positions on the two domains are not constrained to be the same. The fitting artificially used these degrees of freedom to produce low R_p values by making unphysical bond lengths. That these fits are artificial is reflected in the low Hamilton ratios for these models.

TABLE III. Pendry R factors R_p , Hamilton ratios H_r , and the change in the first two interlayer spacings (with representative uncertainties) for the best fits to models Al1 + O1 and Al1-split, for the three sample preparations. For Al1-split, and the Al domains of the Al1 + O1 models, Δd_{12} is the Al1-O1 spacing (averaged for Al1-split) and Δd_{23} is the O1-Al2 spacing (averaged for Al1-split). For the O domain of the Al1 + O1 model, Δd_{12} is the O1-Al2 spacing, and Δd_{23} is the Al2-Al3 spacing. Also shown are the changes in the first two interlayer distances provided by x-ray diffraction and theory. The rotation of the oxygen atoms in the first oxygen layer (O1) obtained from LEED (this work), from x-ray-diffraction experiments, and from first-principle calculations is also shown.

	R_p	H_r	Δd_{12} (%)	Δd_{23} (%)	O1 rot. ($^\circ$)	O1 exp. (%)
Al1 + O1 (Ox)	0.33 ± 0.05	6.5				
Al domain			$+5.0 \pm 8.0$	$+0.2 \pm 7.0$		
O domain			+2.4	-11.8		
Al1 + O1 (Vac)	0.33	5.5				
Al domain			0.0	-1.7		
O domain			+5.0	-14.8		
Al1 + O1 (D)	0.32	7.4				
Al Domain			-38.4	0.0		
O domain			+7.5	-20.2		
Al1-split (Ox)	0.30	8.2	-52.8 ± 5.0	$+1.5 \pm 5.0$	2.7	6
Al1-split (Vac)	0.29	8.6	-50.0	+6.3	3.2	4
Al1-split (D)	0.29	9.4	-50.6	+5.8	2.8	6
X ray (Ref. 6)			-50.8	+16.0	6.7	4.2
Theory (Ref. 14)			-87.4	+3.1	3.05	3.20

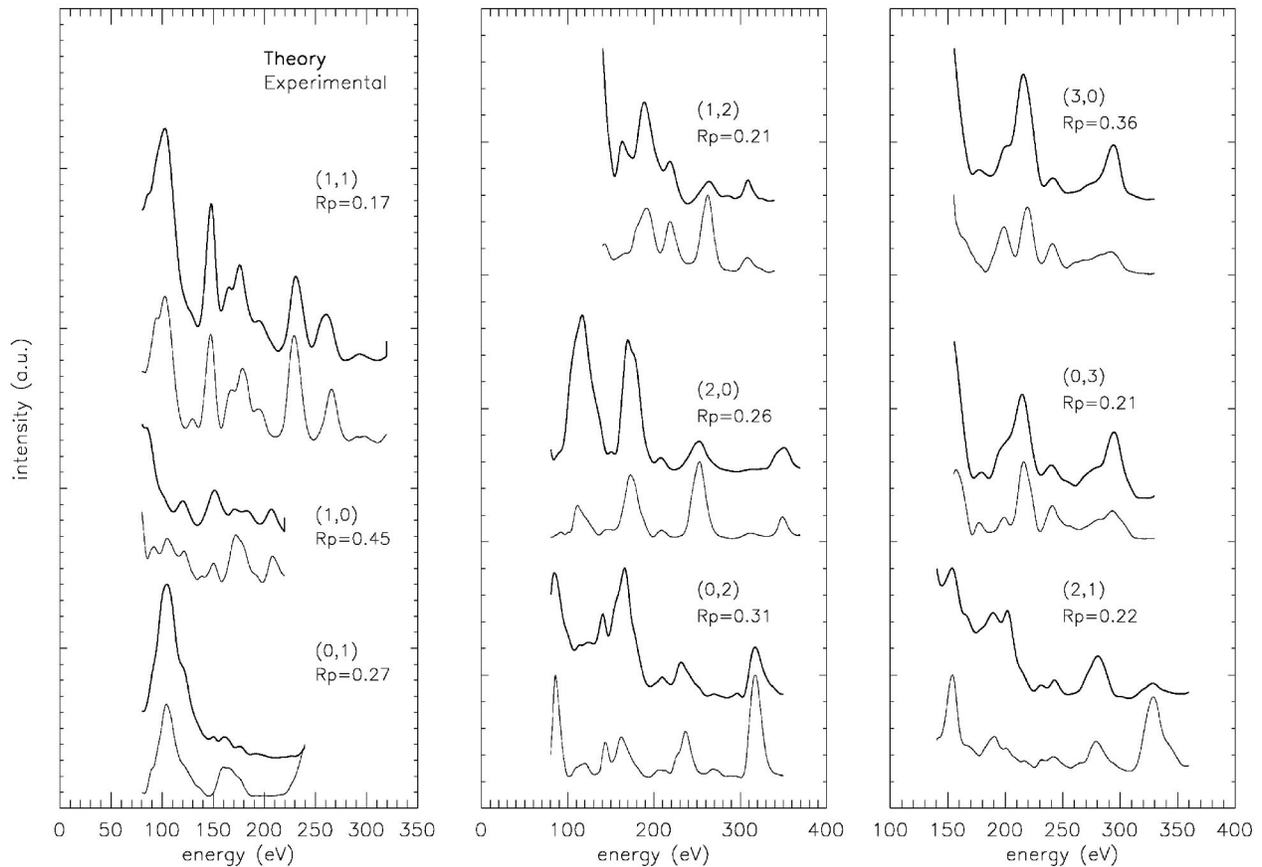


FIG. 7. Experimental (thin lines) and theoretical (thick lines) LEED $I(V)$ curves for the split-position model for the Ox experimental data set. The (i,j) notation gives the index of the diffraction spots.

The models with split positions (model 21) and Al1 + O1 mixed terminations (model 16) are clearly favored over the other models, irrespective of the surface preparation, based upon their low Pendry R factors and high Hamilton ratios. For all three preparation methods, the Al1-split model has the lowest R_p 's and the highest H_r 's. However, since the Al1 + O1 model has R_p values that are only $\sim 10\%$ larger and has acceptable H_r values, this model cannot be immediately discarded. However, additional considerations that are discussed below allow us to clearly favor the Al1-split model.

The best-fit Al1 + O1 model has several questionable properties. First, fitting the two "cleanest" preparation methods (Ox and Vac) with the model gave surfaces that are essentially bulklike, while the preparation method that involved exposure to deuterium (D data) at low temperatures produced a significant contraction of the first interlayer spacing (see Fig. 5 and Table III). This is counter to the usual expectation that clean surfaces are contracted, and that the adsorption of hydrogen results in a return of the first-interlayer spacing to one that resembles the bulk value.^{39,40} Second, the Al1 + O1 models were extremely insensitive to the relative amounts of Al and O domains. In fact, the uncertainty in the mix ratio is on the order of $\pm 40\%$ for all three preparation methods, as can be seen from Fig. 6 for the case of the D data. This insensitivity is physically very un-

satisfying. Finally, the Al1 + O1 model is entirely inconsistent with theoretical predictions for the clean surface, which show that the Al1 model has lowest energy over sapphire's full range of stability,^{12,13,15} precluding a phase-separated surface such as the Al + O models. Figure 6 also shows that the same insensitivity of R_p to composition is obtained for other mixed-domain models considered in this work. Again, a similar R_p behavior was also observed for the other two sample preparation procedures.

In contrast, the simpler Al1-split model gives consistent and physically reasonable results. As seen in Fig. 5, the only significant difference between the two domains is the separation of the top Al layer. This supports the validity of the model since the additional degrees of freedom in the layers below the surface could have been changed to give a good fit. That is, if the model was unsuitable, the domains would differ significantly beyond the first layer, using these additional degrees of freedom to best fit the data. This observation is also consistent with the split-position model having a Hamilton ratio larger than all the other models. In the split-position method, the difference in the position of the Al atoms in the two domains is related to the vibrational amplitude of the outermost Al atoms. The large difference we observe, ~ 0.24 Å, is indicative of an anharmonic enhancement of the perpendicular vibrational mode of the outermost

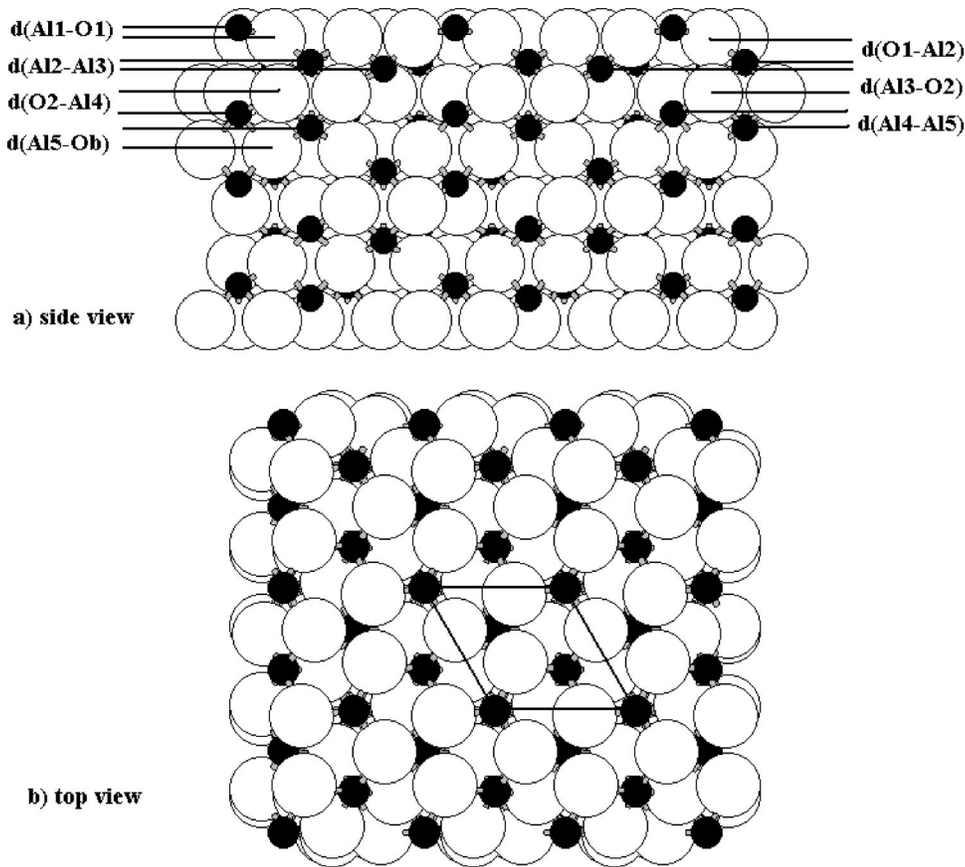


FIG. 8. Plan views of the best-fit structure (the split-position model, No. 21) of α - $\text{Al}_2\text{O}_3(0001)$. Above: side view. Below: top view. The aluminum atoms are the small solid circles, while the oxygen atoms are the large open circles.

Al surface atoms.⁴¹ In addition, the Al1-split model has an average first-interlayer spacing that is in reasonable agreement with the previous x-ray⁶ and ion-scattering⁵ measurements. Importantly, the fact that our three sample-preparation methods result in essentially the same surface structure shows that the α - $\text{Al}_2\text{O}_3(0001)$ surface is very stable, and insensitive to processing conditions. The experimental and theoretical LEED $I(V)$ curves using the split-position model for the Ox data set are presented in Fig. 7. The same level of agreement between theory and experiment was obtained for the two other experimental data sets.⁴² Very recently, constant-stress, constant-temperature (10, 300, and 700 K) molecular-dynamics simulations were performed with shell-model potentials for an Al-terminated α - $\text{Al}_2\text{O}_3(0001)$ surface.⁴³ The interlayer relaxations obtained in that study for the first two interlayer distances ($\Delta d_{12} = -58\%$ and

$\Delta d_{23} = +4\%$) are in good agreement with our results for the Al1-split model (Table III). Their calculated vibrational amplitudes are also in good agreement with those from our LEED analysis.

Although the uncertainties in the displacements parallel to the surface are larger compared to the perpendicular ones, our results suggest a small rotation around a symmetry axis of the oxygen atoms in the first oxygen layer (O1). As shown in Table III, this work's values are in reasonable agreement with those obtained by and x-ray-diffraction experiments and first-principle calculations. A diagram of the best structure for the split-position model is presented in Fig. 8. The coordinates are the average of the atomic coordinates of the two Al domains presented in Table IV.

VI. DISCUSSION

While we find a large interlayer contraction at the surface, the contraction is significantly smaller than that predicted from state-of-the-art calculations.^{12,13,15} Since recent first-principles calculations^{15,16} showed that hydrogen adsorption on the aluminum-terminated surface reduces the contraction close to the value of this and other experimental studies,^{5,6} we next discuss hydrogen on the α - $\text{Al}_2\text{O}_3(0001)$ surface.

The α - $\text{Al}_2\text{O}_3(0001)$ surface is actually quite difficult to hydroxylate. While water undergoes dissociative chemisorption on the α - $\text{Al}_2\text{O}_3(0001)$ surface, extensive hydroxylation occurs only for vapor pressures above about 1 Torr.⁴⁴ For vapor pressures below 1 Torr, water adsorption produces

TABLE IV. Interlayer spacings for the “split-position” model for all three sample preparation procedures.

	Ox data	D data	Vac data
$d_{(Al1-O1)}$ (Å)	0.40	0.42	0.42
$d_{(O1-Al2)}$ (Å)	0.86	0.89	0.90
$d_{(Al2-Al3)}$ (Å)	0.27	0.26	0.32
$d_{(Al3-O2)}$ (Å)	0.89	0.91	0.89
$d_{(O2-Al4)}$ (Å)	0.78	0.85	0.84
$d_{(Al4-Al5)}$ (Å)	0.52	0.52	0.50
$d_{(Al5-Ob)}$ (Å)	0.82	0.82	0.82

only limited amounts of surface hydroxyl, presumably mainly at defect sites.⁴⁴ The fully hydrated surface has been shown experimentally to be oxygen terminated.^{45,46} First-principles calculations show that this surface is thermodynamically stable only for substantial pressures of H₂ or water.^{13,15,16} Furthermore, surface hydroxyl species are readily removed at very modest temperatures. Laser-induced thermal desorption and temperature-programmed desorption have shown that the hydroxyl coverage is negligible above 500 K.⁴⁷ Consistently, Coustet and Jupille found that their cleaning procedure of heating to 1000 K fully desorbed surface hydroxyl, as directly evidenced by vibrational (electron-energy-loss) spectroscopy.⁴⁸ Clearly, then, our “Vac” (heating in vacuum at 650 °C) and “Ox” (heating in O₂ at 650 °C) procedures should produce hydroxyl-free surfaces.

These experimental results are in conflict with a recent ion-scattering study, which concluded that substantial amounts of hydrogen existed on the α -Al₂O₃(0001) surface even after heating to 1100 °C.⁵ The only way to resolve this contradiction with the desorption and vibrational-spectroscopy studies is if hydrogen exists in a nonhydroxyl form on the surface. While we cannot totally discount this possibility, it seems unlikely for several reasons. To begin with, there are only two thermodynamically stable surfaces—the clean (hydrogen-free) aluminum-terminated surface (at low hydrogen chemical potentials) and the fully hydrated surface (at high hydrogen chemical potentials).¹⁵ While hydrogen is calculated to bond directly to aluminum atoms of the Al-terminated surface (making a nonhydroxyl species) at 0 K, the bonding is weak.¹⁵ Indeed, simulations at room temperature using first-principles molecular dynamics revealed only OH species, not Al-H species.^{49,50} Given these observations, it is surprising that hydrogen can remain on the surface at 1100 °C,⁵ the approximate temperature at which substantial oxygen loss from the surface begins, leading to surface reconstructions.^{51,52} Why hydrogen would be more strongly bound than oxygen is unclear. Finally, the source of the surface hydrogen is also unclear—the bulk hydrogen concentration is extremely low in high-quality sapphire.⁵³ Clearly, more experimental work needs to be done on the hydrogen concentration of the α -Al₂O₃(0001) surface at elevated temperatures.

While the majority of experimental and theoretical results suggest that the (0001) surface of α -Al₂O₃ should be relatively free of hydrogen after heating in vacuum, it is not possible at this time to totally discount the presence of any hydrogen. However, it can be argued that hydrogen/hydroxyl is not responsible for the discrepancy between theory and experiment regarding the degree of surface contraction. Our current results find essentially the same surface structure (degree of contraction) despite three quite different processing conditions. Furthermore, our inward relaxation of about 51% agrees well with the values determined by the ion-scattering⁵ (63%) and x-ray-diffraction⁶ (51%) studies. Presumably, our different processing conditions and the other procedures used elsewhere would produce varying amounts of hydrogen/hydroxyl contamination, which would be manifested in differing surface contractions. Our analysis, however, does sug-

gest a possible physical origin for the discrepancy between theory and experiment.

In general, the vibrations of surface atoms are 30–40% larger than those in the bulk. However, using the Debye temperature for the Al atoms derived from the LEED $I(V)$ calculations (350 K), we calculate a bulk vibrational amplitude of 0.12 Å at room temperature. Our results suggest that the vibrational amplitude perpendicular to the surface is very large, about 0.24 Å. Thus, the vibrational amplitude at room temperature is approximately two times greater than the bulk value. Baudin and Hermansson using molecular-dynamics simulations, calculated the vibrational mean-square amplitudes ($\langle u^2 \rangle$) for the alumina surface atoms for different temperatures.⁴³ They concluded that, at room temperature, the $\langle u^2 \rangle_{\text{surface}} / \langle u^2 \rangle_{\text{bulk}}$ ratio for Al ions is ≈ 2.5 , in good agreement with the value suggested by our LEED analysis. While surprising, this result is not without precedent—large vibrations have been observed on other surfaces, e.g., Be(0001),⁵⁴ Ag(111),⁵⁵ Cu(111),⁵⁶ and H₂O(0001),^{57,58} and have also been predicted, but not yet detected, for oxides.^{17,59} Furthermore, as suggested by Harrison *et al.*, the discrepancy between theory and experiment over the amount of first-layer contraction may result from the failure of the zero-temperature calculations to account for large surface vibrations.¹⁷ [In the TiO₂(110) surface structure determined by x-ray diffraction,⁶⁰ the topmost oxygen row is actually contracted significantly more than predicted by first-principles calculations¹⁷]. Such large vibrations may have important implications for understanding the detailed surface properties of metal oxides. In the sapphire case, the presence of enhanced vibrations at the surface is easily visualized in terms of the reduced coordination—the Al-O bonds of the surface Al atoms are almost parallel to the surface, and thus the vibrations are primarily governed by bond-angle changes, which are generally softer than bond-length changes.

VII. SUMMARY

We have studied the α -Al₂O₃(0001) surface structure by examining an unprecedented number of model structures and emphasizing the sensitivity to the sample preparation method. We conclude that the surface termination of α -Al₂O₃(0001) is a single Al layer, that the first interlayer spacing is significantly contracted with respect to the bulk spacing, and that the surface structure is insensitive to our different processing methods, thus resolving contradictory experimental results in the literature. In addition, we suggest that the topmost Al layer has unusually large vibrational amplitudes at room temperature, although temperature-dependent experiments were not carried out to further support this assertion and exclude the possibility of static disorder. Such vibrations may account for the substantial difference between the interlayer contractions determined by zero-temperature calculations and finite-temperature experiments.

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