Understanding reflectance anisotropy: Surface-state signatures and bulk-related features

W. G. Schmidt and F. Bechstedt
Institut für Festkörpertheorie und Theoretische Optik, Friedrich-Schiller-Universität, Max-Wien-Platz 1, 07743 Jena, Germany

J. Bernholc
Department of Physics, North Carolina State University, Raleigh, North Carolina 27695-8202

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Using a real-space multigrid method and ab initio pseudopotentials we calculated the reflectance anisotropy spectra for (001) growth structures of GaP and InP as well as for stepped Si(111):H and hydrogenated Si(110) surfaces. Our results, obtained within DFT-LDA, show good qualitative agreement with recent experiments. Further improvement is achieved by using a numerically efficient GW approach with approximations for local-field effects and dynamical screening. We find contributions to the anisotropy signal related either directly to surface states or to transitions between surface-perturbed bulk wave functions. Our results demonstrate the high sensitivity of RAS to the surface structure and chemistry and show that the optical transitions causing the anisotropy signal take place mainly in a few uppermost atomic layers of the substrate. © 2000 American Vacuum Society. [S0734-211X(00)04104-4]

I. INTRODUCTION

Optical spectroscopy has become an important tool of surface analysis in the last years due to its high sensitivity and in situ applicability. In particular, reflectance anisotropy spectroscopy (RAS), also called reflectance difference spectroscopy (RDS), is increasingly used to characterize static surfaces and to monitor dynamic surface processes in various environments. To fully exploit the technological potential of RAS, a theoretical understanding of the underlying physical mechanisms and reliable schemes to calculate RAS spectra are needed. The progress in the computational modeling of RAS has been rather slow, however. The agreement between the theoretical spectra, usually calculated by using slabs consisting of a few atomic layers, and the measured data is often unsatisfactory. This gave rise to suggestions that long-range effects, hardly accessible to slab calculations, are at least partially responsible for the surface optical anisotropy. A dynamic photon-induced localization of the initial and final states over the range of the penetration depth of light of several hundred Å was suggested to account for anisotropy signals close to the bulk critical point (CP) energies. In Ref. 8 another long-range effect, the quenching of bulk-state wave functions near the surface, was made responsible for the appearance of peaks at bulk CP energies in the surface spectra. These hypotheses are currently subject to both experimental and theoretical investigations. The present article aims at the understanding and interpretation of the surface optical anisotropy.

Realistic surfaces, and growth structures in particular, usually contain defects such as surface steps. The latter have been shown experimentally to give rise to distinct optical anisotropies, the origin of which is presently under debate. The theoretical description of these step-related optical anisotropies is important for a correct interpretation of RAS spectra measured during growth and contributes also to a more complete understanding of the origin of RAS in general.

In the present work we combine converged calculations based on density functional theory (DFT) in the local density approximation (LDA) with a numerically efficient approximate GW approach or a scissors operator approach to calculate the reflectance anisotropy. As model systems we study the (001)(2 × 4) surfaces of GaP and InP, steps on Si(111):H surfaces as well as the hydrogenated Si(110) surface. We find good agreement with experiment. Both bulk-related (“intrinsic”) and directly surface-state related (“extrinsic”) contributions in the RAS are reproduced by well-converged slab calculations. This indicates that the difficulty in theoretical modeling of RAS is not conceptual, but rather caused by: (i) convergence problems due to the large numerical expense required for calculations of surface optical properties, and (ii) the difficulty in accounting for the many-particle effects in the spectra in an efficient yet accurate manner.

II. METHODOLOGY

Our DFT-LDA calculations are performed using first-principles pseudopotentials together with a real-space finite-difference method. A multigrid technique is used for convergence acceleration. The real-space method enables efficient parallelization and is thus particularly suited for the large supercells and many electronic states required for the calculation of the surface dielectric function. The electronic wave functions were mapped on a grid with a spacing of about 0.24 Å. To model the III–V(001) surfaces we used periodic supercells containing 12 atomic layers, the bottom
layer saturated with pseudohydrogens, and a vacuum region equivalent to eight layers. The stepped Si(111):H and Si(110):H surfaces were described with symmetric slabs about 50 Å thick, separated by 14 Å of vacuum.

For the relaxation of the electronic and atomic degrees of freedom we sampled the surface Brillouin zone (SBZ) with four special \( \mathbf{k} \) points in the irreducible part. The surface dielectric function was calculated with uniformly distributed \( \mathbf{k} \) points corresponding to densities of 256 (GaP), 1024 (InP), and about 500 points (Si) in the full \( (1 \times 1) \) SBZ. Further details of the DFT-LDA calculations can be found in Refs. 15–18.

Calculations within DFT-LDA neglect the quasiparticle character of electrons and typically underestimate excitation energies, an effect known as the band-gap problem. For example, we calculate an indirect band gap of 0.58 eV for Si, while the room temperature experiment yields 1.11 eV. A similar underestimation of about 0.5 eV occurs for the \( E_1 \) and \( E_2 \) CPs of the bulk band structure, for which we obtain 3.0 and 3.8 eV, respectively, and for the surface state energies of Si. In order to compare the calculated RAS spectra for Si surfaces with experiment, we therefore apply an upward rigid shift of 0.5 eV to the conduction bands and renormalize the momentum matrix elements accordingly. The same simple recipe has been followed in other recent \textit{ab initio} studies of the optical properties of Si surfaces.

Such a rigid shift (or scissors operator approach) fails, however, if the underestimation of the excitation energies in DFT-LDA is less uniform than in Si. For bulk GaP, for example, it varies between 0.6 and 0.9 eV for states close to the fundamental gap. Furthermore, as will be shown below, different shifts are obtained for bulk and surface state energies in the case of GaP and InP. Rather than applying a rigid shift as for Si we therefore use GW calculations to self-energy effects into account for the III–V (001) surface. In order to obtain accurate excitation energies, the one-electron states should be obtained by solving the quasiparticle Schrödinger equation, i.e., the local exchange and correlation potential \( V_{XC}(r) \) used in DFT-LDA should be replaced by the nonlocal and energy-dependent self-energy operator \( \Sigma(r, r'; E) \) to describe the electron–electron interaction correctly. However, even in its lowest so-called GW approximation, where \( \Sigma \) is expressed as a convolution of the single-particle propagator \( G \) and the dynamically screened Coulomb interaction \( W \), its calculation remains a formidable task. In particular the large number of states entering the surface dielectric function makes calculations very expensive. Recently a linear parametrization of the quasiparticle shifts with respect to the surface localization of the corresponding electronic states was therefore suggested.

We used a different approach and introduced further approximations, following the schemes developed by Hybertsen and Louie and Bechstedt et al. In first order perturbation theory the GW quasiparticle energies are obtained from the DFT-LDA eigenvalues by

\[
E_{n,k}^{\text{GW}} = E_{n,k}^{\text{LDA}} + \frac{1}{1 + \beta_{n,k}} \left( \Sigma_{n,k}^{\text{st}} + \Sigma_{n,k}^{\text{dyn}}(E_{n,k}^{\text{LDA}}) - V_{n,k}^{XC} \right),
\]

where the self-energy operator \( \Sigma \) has been divided into static and dynamic contributions. \( \beta_{n,k} \) is the linear term of the expansion of \( \Sigma^{\text{dyn}} \) in energy around the DFT-LDA eigenvalue \( E_{n,k}^{\text{LDA}} \). The static part can be further divided into two parts:

\[
\Sigma^{\text{st}}(r,r') = \frac{1}{2} \sum_{n,k} \psi_{n,k}(r) \psi_{n,k}^*(r') \left\{ W(r,r';0) - v(r-r') \right\} - \sum_{n,k} \psi_{n,k}(r) \psi_{n,k}^*(r') W(r,r';0),
\]

representing the Coulomb hole \( \Sigma_{\text{COH}} \) and the screened exchange \( \Sigma_{\text{SEX}} \). The \( \psi_{n,k} \) are the DFT-LDA wave functions and \( W(r,r';0) \) and \( v(r-r') \) denote the statically screened and bare Coulomb potentials, respectively (see, e.g., Ref. 19 for a comprehensive review). For the practical calculation of \( W \) we model the dielectric function as

\[
\varepsilon(q) = 1 + \left( (\varepsilon_{\infty} - 1)^{-1} + (q/q_{\text{TF}})^2 \right)^{-1} + 3 q^4/(4 k_F^2 q_{\text{TF}}^2) \varepsilon_{\infty}^{-1},
\]

where \( k_F \) and \( q_{\text{TF}} \) represent the Fermi and Thomas–Fermi wave vectors, respectively. This expression interpolates between the correct behaviors at high and low \( q \) vectors and by construction correctly obtains the static dielectric constant for \( q = 0 \). This model reproduces very well the random phase approximation results for semiconductors. Using the LDA for \( W \), \( \Sigma_{\text{COH}} \) can now be calculated analytically, resulting in a potential that depends on the local charge density. For the calculation of \( \Sigma_{\text{SEX}} \) we retain only diagonal elements in the Fourier transform of \( W \). The local fields are included, however, by using state-dependent charge densities

\[
\rho_{n,k} = \int d^3r \rho(r) \left| \psi_{n,k}(r) \right|^2,
\]

in the calculation of \( k_F \) and \( q_{\text{TF}} \). The dynamic terms \( \beta_{n,k} \) and \( \Sigma^{\text{dyn}} \) in Eq. (1) can be approximated by simple integrals of the dielectric function. For the practical calculation we use Eq. (3) together with a single plasmon pole approximation to describe the frequency dependence. Local-field effects are again included using the mean-density approximation, Eq. (4).

After applying this simplified GW approach to the DFT-LDA bulk band structure of GaP we obtain interband transition energies of 3.2.7 and 4.8 eV for \( E_0 \) and \( E'_0 \), respectively. \( E( X_{1c} ) \) and \( E( X_{3c} ) \) are shifted to 2.2 and 2.4 eV. In the case of bulk InP, the \( E_0 \) gap at \( \bar{\Gamma} \) changes from 0.9 eV in DFT-LDA to 1.4 eV. The transition energies \( E_1 \) and \( E'_0 \) are shifted from 2.5 to 3.2 eV and from 4.2 to 5.0 eV, respectively. These values agree with the experiment within 0.2 eV. The application of this GW scheme to the surface seems problematic, since the model requires the input of the dielectric constant \( \varepsilon_{\infty} \). Northrup and Rohlfing et al. have shown, however, that the inaccuracies caused by the use of bulk dielectric constants for surface GW calculations are very small.
Finally, we use the electronic structure to determine the reflectance anisotropy from the slab polarizability according to the formalism developed by Del Sole \(^{30}\) and Manghi \emph{et al.} \(^{31}\)

**III. RESULTS**

**A. GaP(001)(2×4)**

Total-energy calculations \(^{17,32,33}\) for GaP(001) predict a sequence of (2×4) single-dimer reconstructions for cation-rich surfaces. \(β2(2×4)\) and \(c(4×4)\) reconstructions, known from GaAs(001), form for more P-rich surfaces. A top view of the GaP surface models is given in Fig. 1. These theoretical findings agree with the recent experiments, as discussed in detail in Refs. 17 and 33.

In Fig. 2 we show the RAS spectra calculated within DFT-LDA for the four (2×4) surface structures shown in Fig. 1. The top Ga dimer and mixed-dimer structures show a pronounced negative anisotropy in the low energy region, with minima (denoted \(S\) in Fig. 2) between 2.0 and 2.3 eV. The strength of this anisotropy is directly correlated with the number of Ga–Ga bonds along the \([1\bar{1}0]\) direction. Its magnitude is the highest for the top Ga dimer model with eight bonds, is slightly reduced, and shifted to lower energies for the mixed-dimer geometry with six cation–cation bonds, and is flattened for the \(δ\) structure with only two such bonds. The calculated spectra also show a strong dependence on the structural details at higher energies. For the \(β2\) geometry with three P–P dimers oriented along \([1\bar{1}0]\), we find a relatively broad positive anisotropy between about 2.4 and 4.4 eV. Maxima of the anisotropy appear around 3.2 and 4.1 eV and a shoulder exists at 2.8 eV. The shape of this anisotropy is roughly preserved for the \(δ\) structure, which features one P–P dimer. Its magnitude is, however, somewhat reduced and the spectrum is shifted downwards. An even further reduction in positive anisotropy occurs for the mixed-dimer and top Ga dimer structures, featuring single Ga–P or Ga–Ga dimers, respectively, on top of a Ga-terminated substrate. The evolution of the spectra in the high energy region shows thus a correlation between the positive anisotropy and the formation of P–P dimers.

Our calculations are in accord with the available experimental data \(^{17,34,35}\). The RAS spectra measured for the Ga-rich phase of GaP(001)(2×4) feature a strong negative peak around 2.4 eV. Both the top Ga dimer model and the mixed-dimer model are thus plausible candidates to explain the Ga-rich surface phase. The measured spectrum for the Ga-rich phase shows a maximum between the energies of the \(E_0\) and \(E_1\) CPs. This peak, which should be observed between 2.1 and 2.9 eV in the calculated spectrum, is absent in case of the top Ga dimer structure. It appears, however, as a weak shoulder (denoted \(A\) in Fig. 2) for the mixed-dimer model. As will be discussed below, the agreement between the calculated spectrum for the mixed-dimer structure and the measured RAS improves substantially upon inclusion of self-energy effects, in particular with respect to the shoulder \(A\). The RAS spectra thus indicate that the Ga-rich phase of GaP(001)(2×4) corresponds to the mixed-dimer geometry. This assignment is corroborated by the recently measured surface core-level shifts. \(^{36}\) The spectroscopic results indicate that the extreme Ga-rich limit, which according to the total-energy calculations \(^{17,33}\) is characterized by the top Ga dimer model, may actually not be reached experimentally.

The measured spectrum for the less Ga-rich (2×4) surface phase with an approximately balanced surface stoichiometry is dominated by a “camelback” spectrum with maxima between the energies of the \(E_0\) and \(E_1\) CPs and at \(E'_0\). \(^{17}\) No negative anisotropy appears. The only computed spectrum with no (or very little) negative anisotropy belongs to the \(β2(2×4)\) structure. Maxima appear at 3.2 and 4.1 eV, ...
close to the calculated energies of the \( E_1 \) and \( E_0' \) CPs. Our results thus indicate that the less Ga-rich phase of GaP(001) \((2 \times 4)\) corresponds to the \( \beta_2(2 \times 4) \) structure.

In the following, we focus on the Ga-rich surface phase, which corresponds to the mixed-dimer structure as discussed above. This structure is reproducibly obtained in experiments.\(^{17,35,36}\) To determine the origin of the optical anisotropy we spatially separated the contributions to the RAS from transitions between subsurface/surface electronic states. Solid lines represent the calculations for the complete slab.

The optical anisotropy calculated for the complete slab (solid lines in Fig. 3) corresponds roughly to a superposition of the bulk-related features and surface peaks discussed above. This is a theoretical confirmation of the early work by Aspnes and Studna,\(^{37}\) who discriminated between two components of the reflectance anisotropy: “intrinsic” contributions from surface effects on bulk wave functions and “extrinsic” contributions related to the surface structure.

In Fig. 3 two sets of theoretical data are presented, obtained in DFT-LDA and GWA. As expected from the shifts calculated for the GaP bulk band structure, we observe a distinct blueshift of the spectrum upon inclusion of self-energy effects. However, the shift is nonuniform: it amounts to about 0.6 eV for the surface-state related features (dotted...
lines) and 0.8–1.0 eV for the bulk-related ones (dashed lines). The nonuniformity of the calculated quasiparticle shifts agrees with experiment: As indicated by the arrows in the lower part of Fig. 3, the energy shift between the measured data and the spectrum calculated in GWA is nearly constant and amounts to about 0.3 eV. The inclusion of self-energy effects in the calculations not only leads to energy shifts, but also to pronounced changes in the line shape. The weak shoulder $A$ at 2.8 eV in the DFT-LDA spectrum develops into a maximum at 3.5 eV, in good agreement with experiment.

While the agreement between the measured and the calculated spectra improves much upon inclusion of GW corrections, there remain discrepancies: the excitation energies are somewhat overestimated and the line shapes, in particular for higher energies, do not agree very well. This is not too surprising: one limitation of our study is the neglect of electron–hole interactions. Furthermore, the LDA-like treatment of local-field effects in our GWA model only approximates the screening across the highly inhomogeneous surface region. Our calculations refer to zero temperature and are performed with the theoretical lattice constant and amounts to about 0.3 eV. The inclusion of self-energy effects in the calculations not only leads to energy shifts, but also to pronounced changes in the line shape. The weak shoulder $A$ at 2.8 eV in the DFT-LDA spectrum develops into a maximum at 3.5 eV, in good agreement with experiment.

The major experimental features are qualitatively reproduced in the DFT-LDA calculations, although at much lower photon energies. Inclusion of self-energy effects shifts the peak positions nonuniformly to higher energies, aligning them with experiment within a few tenths of an eV. The shifts for the bulk related features are between 0.7 and 1.0 eV, somewhat larger than the shifts of 0.4–0.5 eV for the surface peaks. The changes of the line shape due to self-energy effects distinctly improve the agreement with experiment.

In order to trace the origin of the surface-related anisotropies in detail, we extended earlier calculations of the InP(001) surface electronic structure to resonant surface states. We find four partially bound surface states, $V_1$–$V_4$, close to the VBM. $V_1$ is formed by the bonds between first- and second-layer cations, $V_2$ corresponds to the P dangling bond at the mixed dimer and $V_3$ arises from the second-layer In–In bonds at the anion site of the mixed dimer (cf. left panel of Fig. 6). $V_4$ is complementary to $V_3$ and comparatively weakly localized at the second-layer cations at the cation site of the mixed dimer. The latter two states are degenerate with bulk bands for most parts of the SBZ (see also Fig. 5). For low photon energies we obtain two pronounced negative peaks (denoted $S1/2$ and $S3$). In the high-energy region features appear close to the $E_1$ and $E_0$ energies. They form a characteristic “three-buckle” shape. Separating spatially the contributions to the optical anisotropy from different slab regions, we find that $S1/2$ and $S3$ originate entirely from the uppermost four atomic layers. This is consistent with experiment, as the measured negative anisotropy for low energies is extremely stoichiometry dependent and therefore surface related. The features at the CP energies arise from transitions between bulk-like electronic states that are perturbed by the surface. In earlier calculations, we have shown that these high-energy features are rather insensitive to the atomic surface structure. This agrees with experiment, where changes in the surface conditions had very little effect on the three-buckle shape.

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In addition, there are surface resonances. The strongly localized mixed-dimer bond \( V_5 \) forms a very flat-band at about 2.0 to 2.1 eV. The bonds between the uppermost P and the second-layer In atoms, \( V_6 \), disperse in energy between 2.1 and 2.2 eV. The lowest unoccupied surface states, \( C_{1-3} \), correspond to empty dangling bonds localized at the surface cations. \( C_1 \) is also localized at the mixed dimer (see also Fig. 5 in Ref. 16). Its energy agrees with the available inverse photoemission data.\(^{48}\) In order to gain an intuitive understanding of the optical anisotropy, we investigated the existence of surface state “signatures” by calculating the RAS due to transitions within pairs of the above identified surface states. We find that some of these transitions give rise to pronounced anisotropy features, the superposition of which accounts nearly for the low-energy part of the spectrum. Figure 6 shows the calculated contribution to the RAS from transitions involving \( V_1 - V_5 \) and \( C_1 - C_3 \). This analysis provides a complete understanding of the surface features of the optical anisotropy: \( S_1 \) is caused by the bonds between first- and second-layer cations. \( S_2 \) arises from transitions mainly involving the second-layer In–In bonds. The symmetry break induced by the mixed dimer on the In–In bonds of the second atomic layer together with the dimer bond itself are responsible for \( S_3 \).

The calculated RAS in GWA reproduces the overall experimental data rather well. The distinct contributions from \( V_1 \) and \( V_3 \) at 2.0 and 2.2 eV are not resolved however. Instead, we obtain one relatively broad minimum at 2.1 eV, due to the finite energy broadening necessary to account for the still limited number of wave vectors used to sample the SBZ. Apart from that, however, all experimentally found features can be identified and quantitatively described by slab calculations that take the electronic transitions within the uppermost atomic layers of the substrate into account.

**C. Hydrogenated Si surfaces**

As an example of step-induced optical anisotropies we study bilayer steps at the hydrogenated Si\((111)\) surface. Since all dangling bonds are saturated, only intrinsic contributions to the RAS are expected. The two principal directions of miscut for the \( \{111\} \) surface are \((1\bar{1}2)\) and \((\bar{1}1\bar{2})\). Assuming straight steps and barring any reconstruction, the first miscut (A in Fig. 7) gives step atoms with one dangling bond each, and the second with two dangling bonds each. In the latter case two different atomic structures are possible: a dihydride structure forms that either lies in a plane parallel (B1 in Fig. 7) or perpendicular to the step edge (B2 in Fig. 7). The optical response of these three configurations is investigated in the following. More precisely, adapting the accepted nomenclature for vicinal samples, the Si\((3\{111\}–(1\bar{1}2))\) and Si\([2\{111\}–(\bar{1}1\bar{2})]\) surfaces are studied, which consist of regularly spaced terraces that are three/two Si atoms wide along the \([1\bar{1}2][\bar{1}1\bar{2}]\) direction, separated by single bilayer steps. In the following we refer to these structures as \((1\bar{1}2)\) and \((\bar{1}1\bar{2})\) steps.

![Fig. 7. Side view of the relaxed step configurations studied in this work. Empty (filled) circles represent Si (H) atoms.](image-url)
The side views of the relaxed step geometries are shown in Fig. 7. All surface bonds are saturated by hydrogen. The Si atoms thus remain in bulk-like configurations and the relaxations are rather small and local. We find that the formation of a vertical dihydride structure in a plane perpendicular to the step edge (B2 in Fig. 7) is energetically preferred by 0.19 eV per step atom compared to the B1 configuration. The formation of vertical rather than in-plane dihydride structures on \( \langle \bar{1}12 \rangle \) steps is in agreement with infrared spectroscopy and Raman spectroscopy. The calculated RAS spectra for the three investigated step geometries are presented in Fig. 8. All steps give rise to strong anisotropy features close to the bulk CP energies. The main differences between the spectrum calculated for \( \langle 112 \rangle \) and \( \langle \bar{1}12 \rangle \) steps are a blueshift of about 0.1 eV and the occurrence of negative anisotropies at around 3 eV for the former and at around 4 eV for the latter two spectra. Based on the striking similarity of the measured step-induced RAS to that of Si(110):H it was suggested in Ref. 11 that the optical response of \( \langle 112 \rangle \) steps arises from transitions in deeper layers. Furthermore, it was stated that the surface local-field effect has to be included to obtain reasonable agreement with experiment.11 In Ref. 53 it was concluded that the existence of Si and H vacancies at the Si(110):H surface is a necessary condition for calculating line shapes in rough agreement with experimental data. Furthermore, it was noted that the anisotropy signal for Si(111):H vicinally cut toward \([11\bar{2}]\) depends nearly linearly on the step density. A similar dependence was recently obtained for Si(001).13 Furthermore, our results, as well as earlier work, show that the atomic relaxations of the step edges are rather small and local. Therefore, it is expected that our calculations can be extrapolated to allow for a meaningful comparison with the measured data.

In Fig. 10 we show the experimental data for surfaces vicinally cut \( \pm 5^\circ \). The upper curve in Fig. 10 shows the RAS of the surface cut toward \([1\bar{1}2]\). The main characteristics of the spectrum are the pronounced anisotropy maxima near the \( E_1 \) and \( E_2 \) CPs of the bulk band structure. This is in agreement with experiment.11 Our results, however, show that even an ideal Si(110):H surface gives rise to a dielectric function-like RAS spectrum and do not indicate a particular importance of the surface local-field effect for that surface.

A direct comparison of the data calculated for stepped surfaces with experiment is not possible, as the existing measurements were performed on surfaces with considerably smaller step densities. Unfortunately, the lateral periodicities needed to accommodate wider terraces are still out of reach for \textit{ab initio} calculations of the optical response. In Ref. 11 it was noted, however, that the anisotropy signal for Si(111):H vicinally cut toward \([11\bar{2}]\) depends nearly linearly on the step density. A similar dependence was recently obtained for Si(001).13 Furthermore, our results, as well as earlier work, show that the atomic relaxations of the step edges are rather small and local. Therefore, it is expected that our calculations can be extrapolated to allow for a meaningful comparison with the measured data.

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qualitative agreement with the corresponding calculated spectrum for \(\langle 11\bar{2}\rangle\) steps (A in Fig. 7). The measured anisotropies are about one order of magnitude smaller than the calculated values. This can be explained partially by the lower step density: a miscut angle of \(\theta = 5^\circ\) corresponds to an average terrace width \(n\) of 11 rows \([\tan \theta = 2\sqrt{3}/(3n - 1)]\), as opposed to the width of three rows in our calculation. Temperature effects and sample imperfections neglected in our calculation also reduce the measured anisotropy signal. The calculated line shape, in particular around the \(E_1\) energy, is only an approximate description of the measured signal. This is not surprising, however. The neglect of excitonic effects in the calculations leads to a bulk dielectric function where the \(E_1\) peak is underestimated by about 50\% and reduced to a weak shoulder of the \(E_2\) peak.

Since the step-induced reflectance anisotropy is to a large extent determined by transitions between bulk-like electronic states, the problematic description of the bulk dielectric function will cause differences in the line shape between the measured and calculated RAS spectra, in particular around the \(E_1\) energy, even if no further approximations had been made in describing the surface. Additionally, the surface local-field effect, neglected in our study, may influence the line shape. If these limitations are borne in mind, however, the agreement between experiment and theory for the \(\langle 11\bar{2}\rangle\) step-induced optical anisotropy is satisfactory.

This is seemingly not the case for \(\langle \bar{1}\bar{2}\rangle\) steps. The lower curve in Fig. 10 shows data measured for the Si(111):H surface miscut 5\(^\circ\) toward [\(\bar{1}\bar{2}\)]. Yasuda et al.\(^{11}\) assumed that the measured anisotropy is induced by bilayer \(\langle \bar{1}\bar{2}\rangle\) steps. This interpretation does not agree with the calculations. Theory predicts positive anisotropy features at the \(E_1\) and \(E_2\) energies, while the measured anisotropies are negative. Even considering all the limitations discussed above, it is implausible that the calculations should yield a qualitatively wrong result. From Fig. 10 it seems as if the \(\langle 11\bar{1}\rangle + 5^\circ\) spectrum was an inverted and somewhat reduced version of the \(\langle 11\bar{1}\rangle - 5^\circ\) spectrum. This fact has already been noted in Ref. 11. By means of infrared \(^{49,55}\) and Raman spectroscopy,\(^{52}\) as well as scanning tunneling microscope (STM) measurements,\(^{56,57}\) it has also been demonstrated that the atomic structures of Si(111):H surfaces miscut toward [\(\bar{1}\bar{1}\bar{2}\)] depend strongly on the preparation conditions. In particular, a transformation of straight dihydride-terminated \(\langle \bar{1}\bar{1}\bar{2}\rangle\) steps into a staircase-like arrangement of monohydride terminated \(\langle 11\bar{2}\rangle\) steps was observed. These newly formed steps form angles of 60\%/120\(^\circ\) with the original step edges. Assuming a complete transformation of straight \(\langle \bar{1}\bar{1}\bar{2}\rangle\) steps into zig-zag chains of ideal \(\langle 11\bar{2}\rangle\) steps, the optical anisotropy of the surface should from pure geometrical considerations correspond to the anisotropy measured for straight \(\langle 11\bar{2}\rangle\) steps, but multiplied by \(-\sqrt{3}\). In reality, the optical anisotropy of remaining, not-transformed parts of \(\langle \bar{1}\bar{1}\bar{2}\rangle\) steps will partially cancel the signal from the zig-zag chains. A further disturbance of the “ideal” RAS spectrum can be expected from contributions due to the edges of the staircase structures. Thus, it seems likely that the optical spectrum of a Si(111):H surface miscut toward [\(\bar{1}\bar{1}\bar{2}\)] is an inverted, reduced, and somewhat disturbed version of the spectrum measured for a surface miscut toward [\(\bar{1}\bar{1}\bar{1}\)]. Obviously, the actual spectrum will depend on the preparation conditions. This hypothesis, if correct, gives a natural explanation of the experimental findings of Ref. 11, which are seemingly at odds with our calculations. However, further experiments are needed to verify the suggested interpretation.

IV. SUMMARY

We have carried out extensive theoretical studies of the optical anisotropies of \((2 \times 4)\) reconstructed (001) surfaces of GaP and InP, stepped Si(111):H, and hydrogenated Si(110) surfaces. The RAS spectra calculated for the compound semiconductor surface structures show characteristic “signatures” from surface states. In particular, transitions between cation–cation bonding states and empty cation dangling bonds give rise to strong negative anisotropies at low photon energies. In addition, we find features caused by surface-modified bulk wave functions. Surface-perturbed bulk wave functions are also mainly responsible for the pronounced anisotropy features close to the bulk CP energies caused by the formation of steps on Si(111):H surfaces. Step orientation and geometry have little effect on the surface optical anisotropy. Inclusion of self-energy effects and a dense \(k\)-point sampling lead to a good agreement between slab calculations and experiments for the complete energy range of the spectrum. No long-range effects, such as photon-induced localization or quenching of bulk states, need to be invoked to explain the appearance of the surface optical anisotropy. The signal is mainly caused by optical transitions in the uppermost few atomic layers of the substrate.

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