

Formation of GaP/Si(100) heterointerfaces in presence of inherent reactor residuals

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Abstract

Adequate silicon preparation is a prerequisite for defect-free III-V growth on Si. We transfer the silicon processing from clean to GaP containing metalorganic vapor phase epitaxy reactors, where we monitor the entire process *in situ* with reflection anisotropy spectroscopy and analyze the chemical composition of the surface with X-ray photoelectron spectroscopy. Beyond a certain sub-monolayer threshold value of (Ga,P) residuals found on the Si(100) surface, GaP grows with an inverted majority sublattice. Analogously to III-V growth on two-domain substrates, the co-existence of Si–Ga and Si–P interfacial bonds at terraces of the same type causes anti-phase disorder in GaP epilayers.

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Keywords

Heterointerfaces, III-V on Silicon, optical in situ spectroscopy, photoelectron spectroscopy

The integration of III-V and silicon is highly desired in both microelectronics and photovoltaics. Adequate Si(100) surface preparation is decisive for the quality of subsequently grown GaP epilayers, which could serve as pseudomorphic III-V/Si(100) quasisubstrates. Single-domain Si(100) surfaces, for example, are a prerequisite for anti-phase domain free III-V heteroepitaxy.¹ Si(100) surface preparation has been studied in great detail: When prepared in H₂ ambient, monohydride-terminated Si dimers form at the Si(100) surface.² Their orientation corresponds to the step structure at the surface: The dimer orientation with respect to the step edges is changing from parallel (B-type, (2×1)) to perpendicular (A-type, (1×2)) at adjacent terraces separated by steps of odd numbered atomic heights.³ A preference for energetically unfavorable⁴ A-type Si(100) with 2° misorientation towards [011], however, could be explained by kinetics: The anisotropic diffusion of Si vacancies preferably along dimers rows and their annihilation at the step edges creates A-type terraces.⁵ Temperatures of about 750 °C and high H₂ pressures are essential here.⁵ Identical conditions applied to almost exactly oriented Si(100) with larger terraces, in contrast, causes layer-by-layer Si removal,⁶ which needs to be avoided in order to prepare smooth, single-domain A-type surfaces. We recently studied GaP nucleation on Si(100) and suggested a kinetically limited formation of abrupt heterointerfaces with either Si–P or Si–Ga bonds depending on the (Ga,P) chemical potential during nucleation.⁷ For GaP grown on Si(100) in P-rich conditions, we could directly evidence the existence of Si–P bonds with X-ray photoelectron spectroscopy (XPS).⁸ Though III-V residuals are inherently present in a realistic processing ambient, Si(100) studies were mainly performed in clean metalorganic vapor phase epitaxy (MOVPE) reactor conditions. The impact of reactor residuals on the formation of the heterointerface is still unclear.

Here, we will discuss the crucial influence of surface misorientation and sub-monolayer coverages of (Ga,P) residuals on the Si(100) preparation and subsequent GaP nucleation in MOVPE ambient (Aixtron AIX-200). We apply reflection anisotropy spectroscopy (RAS, LayTec EpiRAS-200) throughout processing and benchmark these optical *in situ* signals to

in situ mass spectrometry (MS, Hiden HAL301) as well as low energy electron diffraction (LEED, Specs ErLEED 100-A) and XPS (Specs Focus 500 and Phoibos 100) accessible after contamination-free sample transfer.⁹ RAS measures here the difference in complex reflection along $[011]$ and $[0\bar{1}1]$ and is particularly sensitive to (100) surfaces of cubic crystals.¹⁰ Throughout the discussion, we will focus on Si(100) with $0.1^\circ \pm 0.05^\circ$ misorientation towards $[011]$. Important differences compared to higher misorientations will be discussed. We varied reactor pre-conditioning and Si preparation to control the amount of (Ga,P) residuals at the Si(100) surface: (1) Samples prepared in a GaP reactor, which was baked at least 30 min at 1010°C in H_2 and eventually coated with Si, are marked with the index *clean* and (2) samples prepared in presence of intentionally higher amounts of residual (Ga,P) species (e.g. by reducing the bakeout time and applying quicker pressure ramps after Si homoepitaxy) are indexed with *cont.* Identical process parameters were applied for thermal deoxidation and Si homoepitaxy with silane (SiH_4).² In case of GaP/Si samples, subsequent tertbutylphosphine (TBP) and triethylgallium (TEGa) pulses were offered at 420°C prior to pseudomorphic growth of about 40 nm GaP.⁷

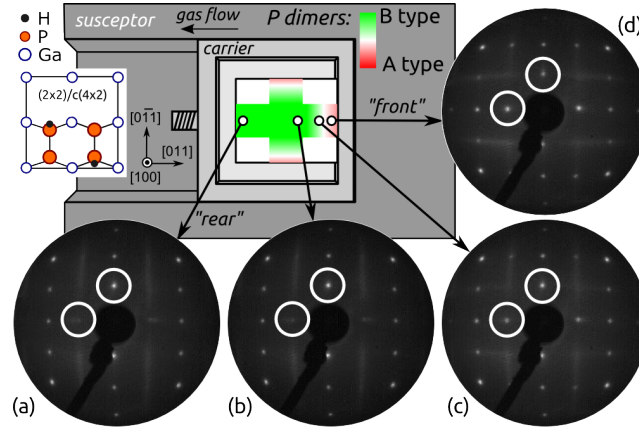


Figure 1: LEED patterns ($E=102\text{ eV}$) of $(2 \times 2)/c(4 \times 2)$ reconstructed GaP/Si_{clean} and qualitative sketch of the sample (length $\approx 2\text{ cm}$) with color-coded domain imbalance along a scan parallel (a,b,c,d) and perpendicular to the flow direction.

The “P-rich” $(2 \times 2)/c(4 \times 2)$ surface reconstruction of GaP(100) is formed by buckled P dimers, which are stabilized by one H atom each.^{11,12} Its LEED patterns consist of spots

at half-order (from the (2×1) -like P dimers) and streaks, both in parallel to the P dimer axis. Due to the zincblende crystal structure, the antiphase domain content at GaP/Si(100) surfaces is reflected in domains of mutually perpendicular P dimers.¹³ Fig. 1 shows LEED patterns of $(2 \times 2)/c(4 \times 2)$ reconstructed GaP/Si_{clean}. At about the center of the sample, the GaP/Si(100) surface is almost single-domain B-type as indicated by spots at half order along $[0\bar{1}1]$ in Fig 1(b). These LEED patterns do not change significantly over large areas of the sample. Towards the front edge, however, the spots at half order along $[011]$ increase in intensity (cf. Fig 1(c)) and at the very edge, the A-type domain even prevails (cf. Fig 1(d)). Similar behaviour is observed at the top and bottom edges. At the rear, in contrast, the B-type majority domains persist (cf. Fig 1(a)). These findings clearly point to an effect at edges close to the susceptor. Two possible explanations are that (i) already the corresponding domain ratio at the Si(100) surface is affected analogously, or (ii) that diffusion of residual atoms influences the chemical ambient during GaP nucleation.

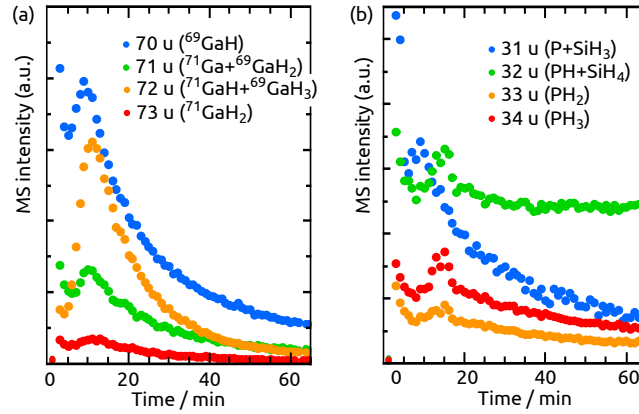


Figure 2: Mass spectrometry during heating (no sample, 950 mbar H₂, 1010 °C reached at 16 min) after a GaP/Si(100) process. (a) Ga-related species. (b) P-related species.

Fig. 2 demonstrates that out-gassing of both Ga and P species from reactor parts needs to be considered: After a typical GaP/Si growth run, we heated the reactor to 1010 °C (950 mbar H₂) and monitored the ratio of atomic weights : degree of ionization, which relate to P and Ga, with a MS connected to the reactor outlet. While we cannot directly translate the measured ionization currents to partial pressures in the reactor, both Ga, GaH_x, P and

PH_x species are clearly present in the gas phase.

In order to study the effect of background residuals quantitatively, we performed XPS measurements. Coverages were quantitatively estimated applying a model described in Ref.,¹⁴ including the Si 2p plasmon loss peak. One ML would correspond to the thickness of a quarter of a lattice constant with every atomic substrate site replaced by the overlayer species. Here, we compare Si_{clean} to Si_{cont} , where we increased the reactor pressure gradient after Si buffer growth. Both samples were transferred to XPS⁹ after Si(100) processing and prior to pulsed GaP nucleation. Fig. 3 shows selected PE lines and Tab.1 gives binding energies (E_B) and coverages. For both samples, the Ga 2p_{3/2} photoemission (PE) line (Fig. 3(a)) clearly consists of two components: a bigger component (Ga_2) and a smaller one at lower binding energies (Ga_1). Compared to recently published data for a thicker GaP/Si(100) reference,⁸ the entire spectrum is shifted towards higher binding energies as the Fermi level at the surface here is located closer to the conduction band. Considering that shift, Ga_2 matches the position of GaP. Due to the small cross-section and lower surface-sensitivity, the P 2p PE line (Fig. 3(b)) of Si_{cont} is not intense enough to reliably distinguish between the existence of two or just one component.⁽ⁱ⁾ Si_{cont} , however, is covered by about 9 times more Ga and about 2 times more P compared to Si_{clean} . When Si_{cont} is measured at 30° exit angle to raise surface sensitivity, the ratio $\text{Ga}_1:\text{Ga}_2$ increases, whereas $\text{Ga}_1:\text{P}$ remains constant (see Tab.1).⁽ⁱⁱ⁾ This implies that the Ga_2 signal is more attenuated than that of Ga_1 and P. We interpret this as Ga_2 being covered by another species, while Ga_1 and P are not. One possible explanation is that three different adsorbate species coexist at the surface: (i) about 9 % of a monolayer (ML) GaP, where Ga (the Ga_2 component) is situated below P, (ii) about 8 % ML of P-species not bound to Ga, and (iii) about 1 % ML Ga-species (the Ga_1 component) not bound to P. The Si 2p PE line (Fig. 3(c)) cannot sufficiently be

⁽ⁱ⁾ The different ratio of $\text{Ga}_2:\text{P}$ between the samples could explain the shift of the P 2p line position, as the two chemically shifted components, one would expect, cannot be reliably resolved.

⁽ⁱⁱ⁾ Intensity ratios were calculated using SPECS ASF factors assuming a homogeneous element distribution. For thin layers, this underrates PE lines at higher kinetic energies, which explains the different ratio when compared to the ML coverage.

fitted by neither one single component, nor by two components, due to the low coverages. Consequently, the Si 2p PE line does not allow a clear conclusion on whether the Ga and the P species are chemically bound to Si. Position-dependent measurements at 90° exit angle of Si_{cont} suggest a decreasing Ga:P ratio in flow direction (rear: $\text{Ga}_1:\text{P}=0.14$, $\text{Ga}_2:\text{P}=0.36$; front: $\text{Ga}_1:\text{P}=0.17$, $\text{Ga}_2:\text{P}=0.74$) with higher amounts of Ga at the front and more P at the rear of the sample.

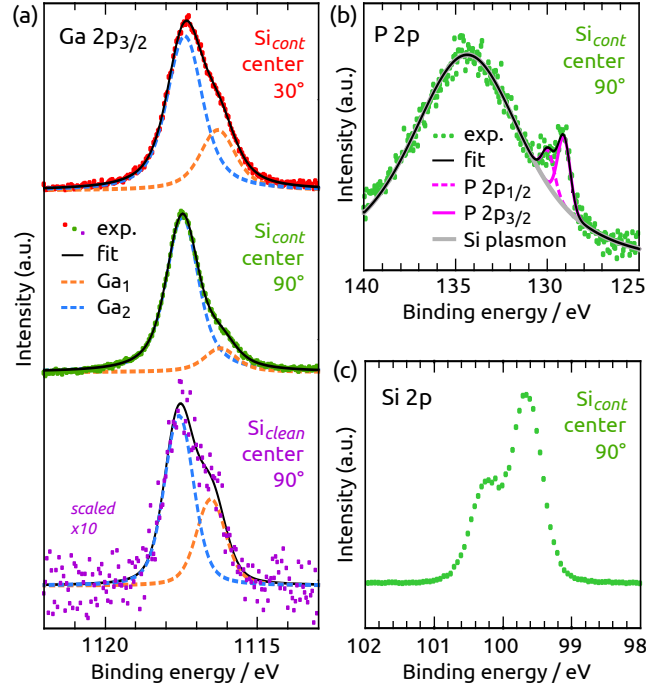


Figure 3: XPS measurements (monochromated Al K_{α}). (a) Ga 2p_{3/2} PE line and fit components of Si_{cont} measured at normal exit angle (green) and 30° exit angle (red) as well as of Si_{clean} (violet). (b) P 2p PE line of Si_{cont} measured at normal exit angle and fit components including the Si 2p plasmon loss peak at 134.37 eV (fwhm=7.4 eV). (c) Si 2p PE line of Si_{cont} measured at normal exit angle.

Fig. 4 shows the RA spectra of one single Si_{cont} sample directly prior to nucleation as well as of $(2 \times 2)/c(4 \times 2)$ reconstructed GaP/ Si_{cont} after heteroepitaxy. The Si(100) signal (green, solid line) is similar to that of H-terminated Si(100), though there are slight differences: Beyond 3.9 eV, RAS of Si_{cont} does not clearly show the peak-like contribution known from clean H-terminated Si(100),^{5,15} but an additional broad peak seems present around 2.2 eV. The clear peak at the E_1 interband transition of Si, however, corresponds to a preferential A-

Table 1: Coverage \mathcal{C} in ML, 14 E_B (wrt. E_F), intensity ratios (corrected with atomic senistivity factors (ASF)) of the $2p_{3/2}$ PE components shown in Fig. 3

| $PE\ line$ | Si_{clean} | | | | Si_{cont} | | 90° | 30° |
|-----------------|--------------|------------------|----------|------------------|----------------------------------|------|------------|------------|
| | E_B/eV | \mathcal{C}/ML | E_B/eV | \mathcal{C}/ML | $ratio$ | | | |
| Ga ₁ | 1116.53 | 0.4 % | 1116.28 | 1.3 % | Ga ₁ :Ga ₂ | 0.17 | 0.39 | |
| Ga ₂ | 1117.58 | 0.7 % | 1117.48 | 8.6 % | Ga ₁ :P | 0.10 | 0.11 | |
| P | 129.38 | 8 % | 129.09 | 17 % | Ga ₂ :P | 0.68 | 0.28 | |
| Si | 99.76 | - | 99.65 | - | | | | |

type Si dimer orientation, which is indicated in the inset (b). RA spectra of $(2 \times 2)/c(4 \times 2)$ reconstructed GaP/Si(100) are related to the buckled P dimers and the sign of the RAS signal (at the surface-modified bulk transition below E_1 of GaP and at the transition at about 2.5 eV) corresponds to the P dimer orientation.^{11–13,16} The GaP/Si_{cont} RA spectrum measured here (orange, dash-dotted line) corresponds to majority A-type P dimers. The intensity, however, indicates that antiphase disorder did not completely annihilate.¹³ Hence, in this case, RAS proves growth of majority A-type GaP on preferential A-type Si. Within an abrupt interface model,^{7,17} the case A \rightarrow A implies Si–Ga bonds at the heterointerface as indicated in the inset (c) of Fig. 4.

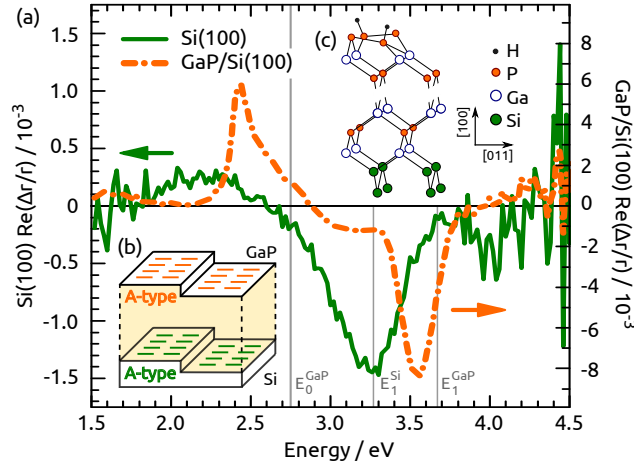


Figure 4: (a) RA spectra of Si_{cont} directly prior to GaP nucleation (420 °C) and of subsequently grown, $(2 \times 2)/c(4 \times 2)$ reconstructed GaP (50 °C). (b) Corresponding orientations of P and Si dimers (prior to nucleation). (c) Sketch of the heterointerface in the abrupt model. Vertical grey lines indicate the critical point energies of Si¹⁸ and GaP¹⁹ respectively.

Both for GaP growth on Si_{clean} (see Fig. 1(a)) and Si(100) with 2° offcut,⁷ in contrast, the GaP sublattice orientation is inverted, which implies Si–P bonds at the heterointerface.⁷ In principle, the inversion of the GaP sublattice could also be explained by Si–P bonds combined with either (i) B-type Si(100), (ii) substitutional P adsorption or (iii) an additional Ga/P interlayer forming one ML below the uppermost Si atoms. However, these processes would then only occur during pulsed nucleation on Si_{cont} and not on Si_{clean} .⁽ⁱⁱⁱ⁾ We observed Si–P bonds and a GaP sublattice orientation in disagreement with substitutional P adsorption of a whole Si ML in Ref.⁸ during GaP nucleation in comparably clean systems. In case of Si_{cont} , our XPS analysis indicates that P is adsorbed on top of Si. Consequently, it seems unlikely that P will substitute Si atoms during subsequent GaP nucleation. Ga was found to promote B-type Si(100) terraces at Si(100), which was Joule-heated to 600°C in UHV.²² To explain our findings, rearrangement towards Ga-covered B-type Si terraces²² would have to take place during pulsed nucleation simultaneously with replacement of the uppermost Ga atoms by P, which we find unlikely. Ga droplet growth on Si(100) terraces with precursor supply may lead to pyramidal etching.^{23,24} The droplet formation was found to be significantly above 1 ML of Ga and can be reduced by higher V:III ratios.²⁵ Here, Ga coverages prior to nucleation are clearly below 1 ML and XPS results even imply that the larger Ga_2 component, which is not located at the very surface, is bound to P whereas the Ga_1 component is on top. Regarding Ga coverages below 0.5 ML, UHV studies suggest that Ga–Ga dimers form on top of Si dimers which remain unbroken.^{26–28} Though experimental conditions are quite different here (monohydride terminated Si, presence of P), intact A-type Si dimers below the adsorbed Ga could explain the Si dimer related RAS minimum at E_1^{Si} (Fig. 4) and possibly also the slight differences in lineshape above and below E_1^{Si} compared to Si(100) prepared in absence of III-V residuals.⁵ The increased amount of Ga available during GaP nucleation also shifts the (P,Ga) chemical potential towards more Ga-rich conditions. *Ab*

⁽ⁱⁱⁱ⁾Note that our experiments differ from As-terminated Ge(100).^{20,21} Here, we vary the amount of sub-ML residuals on Si with A-type majority domains. Subsequently, pulsed nucleation with precursor supply is performed identically.

initio density functional theory calculations showed that the energetically favorable binding situation at the GaP/Si(100) heterointerface changes from Si–P to Si–Ga for more Ga-rich nucleation conditions,⁷ which can explain the observation of inverted GaP sublattice growth on Si_{clean} resp. Si_{cont}. It may also explain the Si–Ga bonds suggested recently.¹⁷

We believe that the Si surface preparation step after Si buffer growth is decisive here for the amount of residuals found on the surface. Annealing at 730 °C in 950 mbar H₂ after thermal deoxidation at 1000 °C is crucial for single-domain, A-type surface preparation of Si(100) 2°. ⁵ Quick temperature and pressure ramps, however, are necessary to avoid layer-by-layer removal on Si(100) 0.1° occurring in this temperature range.⁶ Particularly the pressure ramp is complicating Si(100) 0.1° surface preparation in reactors contaminated with III–V residuals: The rate of residuals desorbing from reactor parts (liner, susceptor, carrier) is increasing with decreasing pressure. This becomes particularly important at elevated temperatures, where desorption of residuals from reactor parts and diffusion of residuals on the sample are high while the H termination² is not stable yet. The consequence is that single-domain surface formation and avoidance of contamination conflict for nominal Si(100) substrates regarding process parameters. The residual atoms may also influence Si vacancy generation and diffusion, but *in situ* RAS reveals an A-type majority domains even for Si_{cont} so that we believe that the kinetic surface processes are comparable to Ref.⁶ Nucleation with a starting TEGa pulse (not shown here) was not sufficient to grow GaP with majority A-type P dimers. Nucleation with high residual amounts or high TEGa precursor supply—in order to reach high amounts of Ga for single-domain GaP—is not easy to control. The co-existence of areas with Si–P and Si–Ga bonds at single-domain Si(100) terraces, however, would cause antiphase disorder analogously to the existence of either Si–P or Si–Ga bonds at two-domain Si(100) terraces. We never observed A-type GaP/Si(100) surfaces in case of A-type Si(100) 2° misoriented substrates. Cooling to temperatures below 730 °C in 950 mbar H₂ (for annealing to prepare A-type terraces) prior to decreasing the reactor pressure for GaP nucleation seems to effectively hinder excessive Ga diffusion on the surface.

Both decreased desorption rates from reactor parts and the H termination of the Si(100) surface² are beneficial here. Nucleation on nominally oriented Si(100) may also benefit from even higher TBP partial pressures.

In conclusion, we showed that a compromise between the avoidance of layer-by-layer removal and diffusion of (Ga,P) residuals to the substrate must be made for Si(100) surface preparation. Sub-monolayer coverages of (Ga,P) residuals on the Si(100) surface—prior offering precursors for pulsed nucleation—strongly influence GaP nucleation. The majority GaP sublattice orientation changes for higher amounts of Ga present at the surface. We explain this by a dependency of abrupt heterointerface structures on the chemical potential during nucleation⁷ and the presence of Si–Ga nucleation seeds prior to offering the first precursor pulse. The coexistence of Si–P and Si–Ga bond domains can lead to antiphase disorder in the GaP epilayer. Consequently, the residual background pressure needs to be controlled precisely for well defined interface preparation. Time-resolved studies are planned in order to resolve the optical anisotropy of the presumably Si–Ga heterointerfaces.

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Graphical TOC Entry

