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Monolithic integration of InP on Si by molten alloy driven selective area epitaxial growth

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We report a new approach for monolithic integration of III-V materials into silicon, based on selective area growth and driven by a molten alloy in metal-organic vapor epitaxy. Our method includes elements of both selective area and droplet-mediated growths and combines the advantages of the two techniques. Using this approach, we obtain organized arrays of high crystalline quality InP insertions into (100) oriented Si substrates. Our detailed structural, morphological and optical studies reveal the conditions leading to defect formation. These conditions are then eliminated to optimize the process for obtaining dislocation-free InP nanostructures grown directly on Si and buried below the top surface. The PL signal from these structures exhibits a narrow peak at the InP bandgap energy. The fundamental aspects of the growth are studied by modeling the InP nucleation process. The model is fitted by our x-ray diffraction measurements and correlates well with the results of our transmission electron microscopy and optical investigations. Our method constitutes a new approach for the monolithic integration of active III-V material into Si platform and opens up new opportunities in active Si photonics.

Introduction

In virtue of indisputable advantages, such as robustness, sustainable oxide and wide prevalence in nature, Si-based photonics and electronics empowered by complementary-metal-oxide-semiconductor (CMOS) technology are being continuously developed with the aim of lowering power consumption, high-speed operation and continual device miniaturization. Since the scaling on silicon devices approaches the physical limit, new materials need to be integrated on to the Si platform, to ensure further developments in electronics and to achieve new functionalities in photonic applications. III-V materials are, for example, required for active photonic and high-speed electronic applications, due to their direct band gap structures and high electron mobility. This makes monolithic integration of III-V materials onto the Si platform very attractive, as it will have a considerable impact on both research and the industrial production of novel integrated photonics, optical interconnects and high-speed electronic devices. Such an integration, however, is challenging, due to the large lattice mismatch between Si and III-V materials, their highly dissimilar thermal expansion coefficients and polar-on-nonpolar epitaxy problems, which typically lead to structures with defect density levels above what is acceptable for device fabrication.

In recent years tremendous effort has been devoted to implementing techniques that reduce the number of defects in the III-V nanostructures monolithically integrated into Si. They include wafer bonding, transfer printing and direct epitaxial methods of integration, such as growth on thick buffer layers, various selective-area growth (SAG) methods and different approaches utilizing liquid group III metallic droplet as a growth catalyst. The latter two families of techniques are more promising, due to their flexible integration into Si-based circuits and avoiding usage of III-V substrates, complicated wafer bonding techniques and difficulties in alignment of III-V elements to the Si-circuit. They have subsequently already been used to obtain high-quality III-V structures grown on Si for laser applications. Besides effective dislocation filtering, the SAG approaches have been used to produce highly ordered nanostructure arrays with precise positioning. The use of group III droplets enables the growth of III-V materials on Si with low defect density, due to a small nucleation area and close to thermodynamic equilibrium growth conditions.

Here, we present a new CMOS compatible epitaxial growth technique by metal-organic vapor phase epitaxy (MOVPE), named molten alloy driven selective-area growth (MADSAG). It is based on using a droplet of the group III element of the final crystal combined with the advantages of SAG. Methods of direct selective area epitaxy on silicon usually suffer from crystal...
defects, which cannot be trapped in at least one of the directions\textsuperscript{13}, as well as from anti-phase boundaries induced by multinuclear growth\textsuperscript{38}. In our method, the molten alloy is used to promote crystallization with a single nucleus for every nanoinsertion.

Implementing this method, we obtain high crystalline quality InP nanoinsertions into (100) oriented Si without using a buffer layer. While our MADSAG method can be used to grow different III-V compounds on Si, the choice of materials here is motivated by further applications in telecom-wavelength compatible photonics\textsuperscript{39}, and the otherwise scarce reports of the epitaxy of InP on Si substrates in the literature.

Experimental

The MADSAG experimental procedure is schematically shown in Fig. 1. The samples investigated here were prepared on Si (100) wafers covered by a Si$_3$N$_4$ mask containing arrays of circular openings 200 nm in diameter and arranged with a pitch of 800 nm. After etching the samples in KOH, inverted pyramidal holes confined by the (111) family of planes are formed in each mask opening. The details of the mask fabrication and Si surface preparation are provided in section S1 of the Supplementary Information (SI). The epitaxial growth on the patterned wafers, in turn, consisted of depositing P-rich In melts with subsequent annealing in a PH$_3$ ambient. Details of the epitaxial growth are described in section S2 of the SI. A series of samples were synthesized at different temperatures and PH$_3$ fluxes, in order to understand the InP nucleation by the MADSAG mechanism, through monitoring the evolution of the morphology, chemical composition and crystal structure and quality of the grown nanoinsertions. The structures were studied in detail using scanning electron microscopy (SEM), high resolution scanning transmission electron microscopy (HR STEM), energy-dispersive X-ray spectroscopy (EDX) and X-ray diffractometry (XRD). Their optical properties were investigated using photoluminescence (PL) measurements.

Results and discussion

We implement our approach to obtain organized arrays of high crystalline quality InP insertions into (100) oriented Si substrates. We present and discuss the results of detailed structural, morphological and optical studies of the obtained structures to identify the growth conditions leading to no defect formation. We identify the impact of the PH$_3$ flux and substrate temperature during the annealing to optimize the process for obtaining dislocation-free InP nanostructures grown directly on Si and buried below the top surface.

Optimization of the group V flux

We start by comparing three samples grown identically but annealed under different PH$_3$ fluxes, in order to investigate the impact of the PH$_3$ flux during annealing on the nucleation of InP inside the openings in Si(100). While changing the PH$_3$ flux, the substrate temperature was kept at 600 °C. Moreover, the total number of PH$_3$ molecules per unit surface was kept constant during annealing, by adjusting the annealing time. Images of three samples with the deposited P-rich In melts not annealed, and annealed at 1.2 x 10$^{-2}$ mol/min (15 min) and 2.2 x 10$^{-2}$ mol/min (8.1 min) PH$_3$ flux rates are presented in Fig. 2, respectively. Fig. 2a-c show plan-view SEM images of the structures, Fig. 2d – f show STEM images of the cross-sections of the samples, Fig. 2g – i are fast Fourier transforms (FFT) of the HR STEM images of the regions marked in Fig. 2d – f, and Fig. 2j – l are EDX composition maps of the samples. Details of the STEM and EDX measurements along with the HR STEM images corresponding to the FFTs in Fig. 2g – i can be found in section S4 of the SI. We find that InP does not nucleate in the sample without annealing, evident by the FFTs of HR STEM images from this sample matching the body-centered tetragonal (bct) crystal structure of In (Fig. 2g). Since the nucleation of the solid phase is known to be limited by the concentration of group V atoms\textsuperscript{40,41}, the P concentration in this case could not have been high enough to initiate an irreversible crystallization. The concentration of P of 12% (comparable with the method accuracy), measured by EDX in Fig. 2j, confirms this. Annealing under a PH$_3$ flow is expected to increase the P concentration in the melt, and subsequently promote the nucleation and further crystallization of InP. Images of the sample annealed at PH$_3$ flux 1.2x10$^{-2}$ mol/min suggest a partial nucleation of InP, as regions of both bct In and face centered cubic (fcc) InP can be observed in this sample (see Fig. 2h).

The EDX measurements support this by revealing a non-homogeneous distribution of P and its segregation in the regions with an fcc InP phase (Fig. 2k). The InP region has the same crystallographic orientation as the Si substrate (see section S4 in the SI) and therefore, we conclude that InP nucleates at the interface between the In melt and Si and it inherits the substrate orientation.

Increasing the PH$_3$ flux leads to increase in the volume of the crystallized InP, evident by the images in Fig. 2c, f, i and l, from the sample annealed at PH$_3$ flux 2.2x10$^{-2}$ mol/min. Expansion of the volume of material in the nanoinsertions due to complete conversion of In to InP can be clearly seen in the plan-view SEM image in Fig. 2c. HR STEM images and the corresponding FFTs from this sample only show the fcc InP crystal phase, and the P concentration in the EDX map in Fig. 2l has a uniform
distribution. We observe a sharp interface between the InP and Si lattices, with the crystal orientation of InP matched to that of the surrounding Si crystal (see section S4 of SI). Further increasing the PH$_3$ flux beyond this value did not considerably affect the outcome.

Based on the results we conclude that the epitaxial growth in the MADSAG process consists of two stages: 1. deposition of a group III droplet and 2. nucleation and further crystallization of the III-V material during annealing under a group V flux. The nucleation and growth of the InP crystal are, however, driven by the concentration of P atoms in the In melt. This concentration, in turn, is determined by the difference between the absorption and desorption rates of P in the In melt. Assuming complete PH$_3$ pyrolysis, we expect that the absorption rate of P is primarily controlled by the PH$_3$ flux. Thus, when the PH$_3$ flux is kept constant, the P concentration in liquid In is determined by the substrate temperature, since the desorption of P is temperature dependent.

![Image](image_url)

Fig. 2. (a-c) SEM images of nanoinsertion grown in (100) Si and annealed under different conditions in plan-view geometry, (d-f) STEM images of the cross-sections of the nanoinsertions, (g-i) FFTs of HR STEM images of the regions in the nanoinsertions marked in (d-f), and (j-l) the corresponding concentration maps of In and P, illustrating the crystallization during the MADSAG process without annealing (a, d, g, j) and with annealing under PH$_3$ flux of 1.2x10$^{-2}$ (b, e, h, k) and 2.2x10$^{-2}$ mol/min (c, f, i, l). The crystallographic directions of the Si substrate in the plan-view and cross-sectional images are shown in (a) and (d). The EDX measurements have an accuracy of 10-15%.
Optimization of the substrate temperature

In order to better understand how the annealing temperature affects the MADSAG process, we have investigated the optical properties, morphology and crystalline quality of a set of nanoinsertions annealed at different temperatures under a PH₃ flux of 2.2x10⁻² mol/min. Fig. 3 compares the PL spectra acquired at 77K and 290K from the structures annealed at temperatures 550, 600 and 650°C. Details of the PL characterization are provided in section S5 of the SI. The PL signal from the three samples demonstrate similar shapes but have significantly different intensities. The highest PL intensity and the narrowest full width at half maximum (FWHM) are observed from the InP/Si structures annealed at 600°C, and the signals from the samples annealed at lower and higher temperatures both have decreased PL intensities. The signal from the sample annealed at 600 °C shows a narrow PL peak at the energy corresponding to InP in zincblende crystal phase and a contribution from type-II band alignment 43–45 (see section S5 of the SI for details). The observed PL peaks show FWHM below 25 meV at 77 K and 55 meV at 300 K, which is attributed to the high crystallinity of the grown InP structures.

Typical plan-view SEM images of arrays of the nanoinsertions after annealing at 550, 600 and 650°C are shown in Fig. 4a-c. We conceive that a square-base truncated pyramid shape of the structures in plan-view implies uniform growth of InP on all four etched {111} planes in Si and complete crystallization of In into InP. In the samples annealed at 600 °C (Fig. 4b) most of the structures have the truncated pyramidal shape. The structures annealed at slightly lower temperature (Fig. 4a) show similar morphology, while annealing at higher temperatures leads to irregularities in morphology (Fig. 4c). We relate the morphology variation to crystal defects and residual In after InP crystallization, which can exist under non-optimized growth conditions. In this case of non-optimal growth conditions, crystal defects tend to nucleate at the interface between the remaining In and InP. The crystallinity of these arrays of InP nanostructures were further examined by XRD.

Fig. 4d-f compares the XRD InP (111) pole figures acquired from the samples annealed at temperatures 550, 600 and 650°C, respectively. The peaks crystallographically aligned with the Si lattice are labeled with Miller indices, while the peaks arising from twinning in the InP crystal are marked with red arrows. The four-fold symmetry of the peaks from twinned structures is related to four possible symmetries inherited from nucleation on each of the four (111) Si planes formed in the mask openings. The samples annealed at different temperatures show very similar peak patterns.

However, the comparison of the integrated intensity of the signal corresponding to the defect-free InP (111) planes, summarized in Table 1, shows that the samples annealed at 550 and 600 °C have higher intensity of these reflections and thus better crystalline qualities. The comparison of the data with a reference InP wafer can be found in section S6 of the SI.

XRD In (011) pole figures are displayed in Fig. 4g-i, revealing the impact of the annealing temperature on the presence of residual metallic In. The measurements from the three samples demonstrate similar peak positions corresponding to the four (110) crystallographic planes, while their intensities vary with...
annealing temperature (integrated reflection intensities are summarized in Table 1).

| Annealing temperature, °C | Integral intensity of In (011) reflection, a. u. | Integral intensity of InP (111) reflection, a. u. |
|---------------------------|-----------------------------------------------|-----------------------------------------------|
| 550                       | 621                                           | 16910                                         |
| 600                       | 250                                           | 17002                                         |
| 650                       | 1243                                          | 6842                                          |

This shows that the residual In in the structures is highly crystalline with a bct crystal structure (space group I4/mmm), which is consistent with what we have observed in HR STEM images. Comparing the integral intensity of In (011) reflections in the pole figures from the three samples in Table 1, we conclude that the sample annealed at 650 °C has the largest amount of monocry stalline In, whereas the sample annealed at 600 °C has the least amount of residual In.

The results so far suggest that InP crystals having a symmetrical truncated octahedral pyramid profile in plan-view contain no residual In and are free from defects. Fig. 5 shows images of two structures in cross-sections, one with a such a profile (Fig. 5a-d) and the other with an irregular and asymmetrical plan-view profile (Fig. 5e-h). The EDX maps in Fig. 5c and 5g clearly confirm that the structure with an irregular plan-view shape contains residual In, while the structure with a truncated pyramid shape does not.

Fig. 4. Top-view SEM images of the InP structures annealed at 550 (a), 600 (b) and 650 °C (c). InP (111) (d-f) and In (011) (g-i) pole figures acquired from MADSAG InP/Si annealed at 550, 600 and 650 °C respectively.
Moreover, evidence of dislocations in the nanoinsertions with irregular plan-view profile can be seen in bright field TEM images in Fig. 5 e and h. Dislocations, however, are typically absent in the structures with truncated pyramid plan-view profiles and the only evidence of crystal twinning can be found in these structures, Fig. 5a and d. Twins are known to have less destructive effect on device performance than other dislocation-type defects, since no dangling bonds form and all atomic bonds are saturated. However, they typically cause appearance of additional peaks in PL signal obtained from InP nanostructures. As we show in section S5 of the SI, those peaks in our measurements are significantly overlap with the signal from pure zincblende InP crystals. It results in asymmetric widening of cumulative PL signal and thus defines the abovementioned temperature-dependent FWHM in optimized samples with no remaining In.

We relate the incomplete consumption of In during InP formation with a lack of P atoms after full coverage of the top surface of the opening by InP crystal, which blocks further transport of P atoms to the In melt. This will be discussed further in the modeling describing the growth process. We assume that the In(011) intensity in the XRD data, $I_{\text{In}(011)}$, is directly related to the residual number of In atoms $N_{\text{res}}^{\text{In}}$. We therefore introduce a growth model to estimate the amount of residual In and understand the non-monotonic temperature dependence of the In(011) XRD signal.

**Growth modeling**

The residual number of In atoms can be calculated as the difference between the initial number of In atoms in melt $N_{\text{init}}$ and the number of In atoms in solid InP. The number of In atoms in InP equals to the number of P atoms, which consist of P dissolved in liquid In before nucleation $N_P^0$ and absorbed during the crystallization $N_P^{\text{cryst}}$, thus:

$$N_{\text{res}}^{\text{In}} = N_{\text{init}} - N_P^0 - N_P^{\text{cryst}}. \quad (1)$$

Nucleation and further crystallization of InP in the In melt are controlled by P concentration $C$ driven by the absorption and evaporation of P atoms. We assume that $C$ is rather uniform inside the In melt, due to the short characteristic time of the species mixing, which can be estimated as $A/D \sim 10^{-5}$ s, where $A$ is the area of the opening and $D$ is the diffusion coefficient of P in liquid In, which has the order of magnitude $\sim 10^{10} \text{m}^2/\text{s}$ at considered growth temperatures.

Before InP nucleation, P absorption at rate $F$ and evaporation at rate $C/\sigma$ are balanced at the considered growth temperatures, and thus $C$ is given by the corresponding steady-state value $C_0 = \sigma F \tau$ and does not depend on time. In our notations, $\sigma$ is the area of the vapor-
liquid interface per atom, and $\tau$ is the characteristic time of evaporation.

After nucleation, the amount of P adsorbed during crystallization can be determined by integrating the difference between the absorption and evaporation rates:

$$N^\text{cryst}_P = \int_0^t \left( F - \frac{C(t)}{\sigma t} \right) Ad t. \quad (2)$$

Analytical calculations for $C(t)$ and $t^\text{cryst}$ given in the Si allow the determination of $N^\text{cryst}_P$ and $N^\text{res}_P$ as functions of the annealing temperature $T$. Using the rough estimation of the integral intensity of In (011) reflections $I_{\text{in(011)}} \sim N^\text{res}_P$, our modeling result can be summarized as:

$$I_{\text{in(011)}} = I_0 + I_1 \exp \left[ -\frac{T_\text{evap}}{T_\text{D}} \right] + I_2 \exp \left[ \frac{T_\text{evap}+T_\text{D}}{T_\text{D}} \right] \quad (3)$$

With the temperature-independent term $I_0$, and increasing and decreasing exponential functions of the growth temperature with the corresponding prefactors $I_1$ and $I_2$. The temperature-dependent exponents are controlled by characteristic barriers for P evaporation $T_\text{evap}$ and diffusion $T_\text{D}$ in liquid In. Fig. 6 shows the integral intensity of the pole figure peaks corresponding to In(011) reflection fitted by Eq. (3) with fitting parameters $I_0 = 127150$, $I_1 = 228333$, $I_2 = 85.7$.

Eq. (3) brings together the effects of several processes, which contribute to the nonmonotonic temperature dependence of the residual amount of In, and thus the intensity of the In(011) XRD signal. The increasing exponent $I_1 \exp \left[ -\frac{T_\text{evap}}{T_\text{D}} \right]$ in Eq. (3) is proportional to $C_0$ and includes two contributions. First, according to our estimations (see section S7 in the SI), the P concentration before nucleation $C_0$ increases with temperature. Thus, less In remains at the end of the process, which corresponds to a decrease of the prefactor $I_1$ in Eq. (3) (see section S7 in the SI for details). Second, an increase in $C_0$ leads to faster motion of the growth front (shorter $t^\text{cryst}$). It may lead to the full coverage of the top surface of the opening by InP before the amount of P required for complete In conversion to InP gets absorbed. Thus, some residual In remains as we show in Fig. 5e, f. A similar situation can occur due to lowering the PH$_3$ flux during the annealing, as shown in Fig. 2c.

Meanwhile, during a faster crystallization stage less P evaporates from the melt and thus, less In remains, which is represented by the decreasing exponent $I_2 \exp \left[ \frac{(T_\text{evap}+T_\text{D})}{T_\text{D}} \right]$ in Eq. (3). Therefore, the competing effects of shorter crystallization time and increasing initial P concentration shape the observed nonmonotonic temperature dependence.

Conclusions

In conclusion, we have proposed and investigated a new method called MADSAG for the selective-area epitaxial growth of InP on (100) Si. The InP nanostructures grown under optimized conditions show a very low crystal defect density and sharp interfaces between the grown InP and hosting Si. They also demonstrate a high intensity optical response, verified by PL measurements. We have investigated the influence of the growth parameters on the nucleation and morphology of InP nanoinclusions by SEM, HR STEM, EDX and XRD. Under non-optimal conditions, residual In remains in the structures, which negatively impacts the crystalline quality and the PL signal of the InP nanoinclusions. The remaining In can be eliminated and complete crystallization of the In melt can be achieved by optimizing the growth temperature and PH$_3$ flux. We thus conclude that MADSAG is a very promising CMOS compatible growth method, which can facilitate the development of novel active hybrid nanodevices integrated into the Si platform.

Conflicts of interest

There are no conflicts to declare.

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