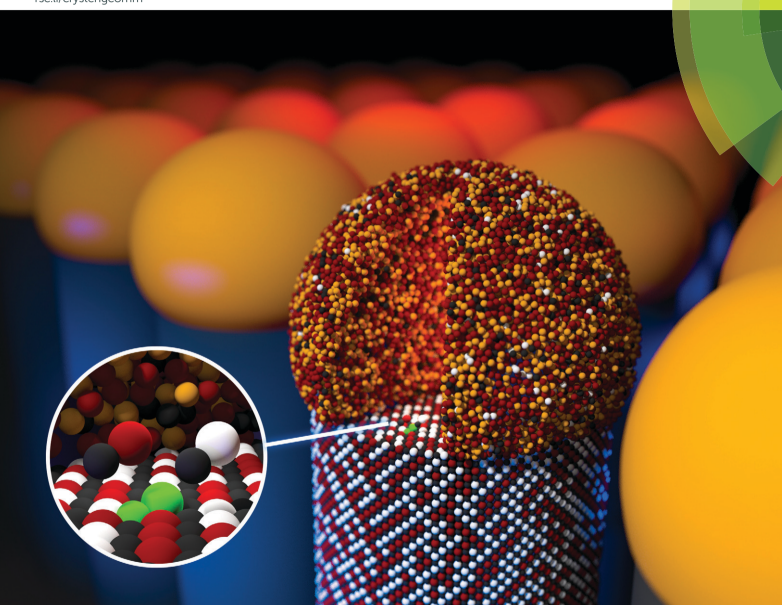
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Nucleation-limited composition of ternary III-V nanowires forming from quaternary gold based liquid alloys†

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We derive an analytic expression for the composition of a ternary solid material nucleating from a quaternary liquid melt. The calculations are based on the two-component nucleation theory with realistic descriptions of the liquid and solid phases. We apply this theory to gold-catalyzed, nucleation limited vaporliquid-solid growth of ternary III-V nanowires. We consider ternary gallium, indium, and aluminum arsenides and antimonides and discuss growth conditions for optimum composition control in these materials. Furthermore, we compare our calculations with the results of an equilibrium thermodynamic model.

Introduction

Semiconductor nanowires (NWs) present a promising class of nanoscale objects whose properties are dramatically different from the bulk counterparts. Inspired by the enhanced functionality, III-V NWs have received tremendous research interest1-3 that led to development of new applications. Particularly, NWs can be used as basic functional elements for a wide range of electronic and optical devices including high brightness light emitting diodes, transistors, low cost solar cells, flexible panel displays,7 and logic gates.8 However, applying NW devices to industrial scale requires advanced growth technology with a precise control of the NW composition, heterojunction properties, morphology, and crystal structure. The vapor-liquid-solid (VLS) mechanism9 is very promising since it enables growth of size-uniform NW arrays with a high degree of crystal quality. This method involves feeding semiconductor materials into a liquid catalyst droplet from the ambient vapor, which leads to vertical crystal growth as a result of supersaturation in the droplet. Self-catalyzed NW growth is a widely used VLS approach¹⁰⁻¹⁴ allowing one to avoid contamination by foreign metals. However, gold remains one of the most attractive catalysts due to its versatility and simplicity of the control over the NW diameter and position.

The critical step in NW-based device design is the ability to control the composition of ternary III-V NWs, which involves the bandgap tunability by variation of the compound concentrations in the ternary alloy. In recent years, enormous progress has been made toward improving the growth technologies by broadening the spectrum of the NW materials. Particularly, growth of AlGaAs and InGaAs NWs using molecular beam epitaxy (MBE)15 and metal-organic vapor phase epitaxy (MOVPE)16 have been studied. Sb-based NWs such as GaAsSb, ¹⁷ InAsSb (ref. 18) and InGaSb (ref. 19) are nowadays of particular interest in terms of mid-infrared band gap engineering. In these investigations, growth of ternary NWs over a broad range of compositions has been demonstrated. However, the liquid droplet composition has not been systematically investigated which impedes a direct comparison between experimental and theoretical results. NW synthesis resulting in compositions within the miscibility gap can, for systems where this is relevant, be explained in terms of growth kinetics, 20 whereas the present work is devoted to thermodynamically stable compositions.

Analytical models which link the solid-liquid and solidvapor compositions have been reported by Dubrovskii et al.21,22 and Glas.23 However, a full description of ternary NWs that form from quaternary alloys (three NW constituencies and gold) is still lacking because of unknown thermodynamic and kinetic constants such as chemical potentials and crystallization rates of different III-V pairs.

The current investigation is aimed at improving the fundamental understanding of the ternary NW formation from quaternary alloys and is based on realistic thermodynamic descriptions of all the considered phases. In our approach, binary and ternary interactions are taken into account. We calculate and analyze compositional diagrams for highly relevant III-V materials including the In-Ga-As-Au, Al-Ga-As-Au, In-Ga-Sb-Au and In-Sb-As-Au systems. Furthermore, a

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comparison between self-catalyzed and Au-catalyzed NW growth in these materials systems is presented. The obtained results provide a basis for understanding the ternary NW formation and can be useful for choosing the growth conditions to tune the NW composition to the desired values.

Calculations

Paper

The formation energy of a binary, or two-component, nucleus can be written as 20,24

$$F(x,s) = -\Delta\mu(x,y)s + a\sqrt{s},$$
 (1)

where s is the size and x is the composition of the nucleus, so that s = i + j and x = i/(i + j), where i and j are the number of pairs of the two binaries that the nucleus consists of. The parameter y is the composition of the mother phase, $\Delta \mu(x,y)$ is the chemical potential difference between the mother and daughter phases, and a is a surface energy term. The composition and the size of the critical nucleus are thus given by the simultaneous solutions to the equations

$$\frac{\partial F}{\partial x} = -\frac{\partial \Delta \mu}{\partial x} s + \frac{\mathrm{d}a}{\mathrm{d}x} \sqrt{s} = 0, \quad \frac{\partial F}{\partial s} = -\Delta \mu + \frac{a}{2\sqrt{s}} = 0 \tag{2}$$

It has been argued, however, that the surface energy of the critical nucleus is at a minimum due to surface segregation effects.²⁵ This property corresponds to the condition da/dx = 0, which considerably simplifies the calculations and leads to the expression

$$\frac{\partial \Delta \mu}{\partial x} = 0 \tag{3}$$

for the composition of the critical nucleus.

We will consider the formation of a ternary alloy, $A_xB_{1-x}D$, from a quaternary solution mother phase containing A, B, D, and U, where U can be thought of as a solvent, often gold. For this situation, the chemical potential difference of the ternary alloy in the liquid and solid can be expressed as²⁶

$$\Delta \mu = x(\mu_{\rm A}^{\rm L} + \mu_{\rm D}^{\rm L} - \mu_{\rm AD}^{\rm S}) + (1 - x)(\mu_{\rm B}^{\rm L} + \mu_{\rm D}^{\rm L} - \mu_{\rm BD}^{\rm S}), \tag{4}$$

where the superscripts L and S refer to the liquid and solid phases, respectively. In order to express chemical potentials, we need thermodynamic models for the Gibbs free energy of the respective phases. The molar Gibbs free energy of the liquid phase is generally given by

$$G^{L} = \sum_{I} c_{I} \mu_{I}^{0} + RT \sum_{I} c_{I} \ln c_{I} + \sum_{I} \sum_{J=I+1} c_{I} c_{J} \left[\omega_{IJ} + \omega_{IJ}' \left(c_{I} - c_{J} \right) \right] + \sum_{I} \sum_{J=I+1} \sum_{K=I-1} c_{I} c_{J} c_{K} \omega_{IJK},$$
(5)

where c_I is the molar fraction of element I in the liquid phase, μ_I^0 is chemical potential of pure component I in the

liquid state, R is the gas constant, and T is temperature. The ω are interaction parameters accounting for deviations from ideality. For binary interactions, we assume a sub-regular solution model with composition dependent interactions, whereas the ternary interaction parameters are assumed to be independent of composition. All ω parameters are allowed to be temperature dependent. For the composition dependent binary interactions, one needs to be careful with the order of the indices. In CALPHAD, one uses an alphabetical order convention and writes the composition dependent part of the AB interaction term as $x_A x_B \omega'_{AB} (x_A - x_B)$. If, for instance, A is interpreted as In and B as Ga, the alphabetical order is broken and we need to construct $\omega'_{InGa} = -\omega'_{GaIn}$, where ω'_{GaIn} is the standard tabulated parameter (because Ga comes before In according to alphabetical order). For the composition independent interaction terms (binary and ternary), the order does not matter.

The chemical potentials of the species in the liquid are expressed by differentiation of G^{L} in eqn (5) according to

$$\mu_J^{L} = G^{L} + \frac{\partial G^{L}}{\partial c_J} - \sum_I c_I \frac{\partial G^{L}}{\partial c_I}.$$
 (6)

The zincblende (ZB) solid phase, $A_xB_{1-x}D$, is modeled as a binary regular solid solution with chemical potentials given by

$$\mu_{AD}^{S} = \mu_{AD}^{0} + RT \ln x + (1 - x)^{2} \left[\omega_{S} + (4x - 1)\omega_{S}' \right]$$
 (7a)

$$\mu_{BD}^{S} = \mu_{BD}^{0} + RT \ln(1-x) + x^{2} \left[\omega_{S} + (4x-3)\omega_{S}'\right],$$
 (7b)

where μ_{AD}^0 and μ_{BD}^0 are the chemical potentials of pure AD and BD binaries, respectively, and ω_S , ω_S' are the temperature dependent interaction parameters of zero and first order.

By explicitly expressing the chemical potentials of the species in the liquid, we construct the chemical potential difference according to eqn (4). Differentiating this expression according to eqn (3) yields an analytical expression for the composition of the critical nucleus, which serves as a first approximation of the composition of the solid phase,

$$RT \ln\left(\frac{x}{1-x}\right) - 2\omega_{\rm S}\left(x - \frac{1}{2}\right) - \omega_{\rm S}'\left(6x^2 - 6x + 1\right) = \varphi(y). \tag{8}$$

Here, we introduce the $\varphi(y)$ function given by

$$\varphi(y) = \alpha + RT \ln\left(\frac{y}{1-y}\right) - 6c_{\text{tot}}^2 \omega_{AB}^2 y^2$$

$$-2c_{\text{tot}} \left[\omega_{AB} - 3c_{\text{tot}} \omega_{AB}^\prime - c_D(\omega_{AB}^\prime + \omega_{BD}^\prime) - c_U(\omega_{AU}^\prime + \omega_{BU}^\prime) + c_D \omega_{ABD} + c_U \omega_{ABU}^\prime\right] y,$$
(9)

with

$$\alpha = \Delta \mu_{AD}^{0} - \Delta \mu_{BD}^{0} + (\omega_{AB} - c_{tot}\omega_{AB}' - 2c_{D}\omega_{BD}' - 2c_{U}\omega_{BU}' + c_{D}\omega_{ABD} + c_{U}\omega_{ABU})c_{tot} + (\omega_{AD} - \omega_{BD})c_{D} + (\omega_{AU} - \omega_{BU})c_{U} - (\omega_{AD}' - \omega_{BD}')c_{D}^{2} - (\omega_{AU}' - \omega_{BU}')c_{U}^{2} + (\omega_{ADU} - \omega_{BDU})c_{D}c_{U}.$$
(10)

In the above equations, we have made the substitutions $c_{\text{tot}} = c_{\text{A}} + c_{\text{B}}$ and $y = c_{\text{A}}/c_{\text{tot}}$. In some relevant cases (see the discussion in ref. 20 for self-catalyzed growth), there is a closed form approximation for eqn (8) given by

$$y = \frac{x}{x + (1 - x)e^{\frac{2\omega_3(x - 1/2) + \omega'_3(6x^2 - 6x + 1) + b}{RT}}}.$$
 (11)

Here,

$$b = \alpha - 6c_{\text{tot}}^2 \omega_{AB}'$$

$$-2c_{\text{tot}} [\omega_{AB} - 3c_{\text{tot}} \omega_{AB}' - c_D (\omega_{AD}' + \omega_{BD}') - c_U (\omega_{AU}' + \omega_{BU}')$$

$$+ c_D \omega_{ABD} + c_U \omega_{ABU}]$$

$$(12)$$

for systems requiring values of y close to one to get any appreciable amount of AB in the solid and

$$b = \alpha \tag{13}$$

for systems with the reversed behavior.

From the thermodynamic point of view, the selected states $(c_{\text{tot}}, y, c_{\text{D}})$ and T of the homogeneous liquid particle are not equilibrium states but rather represent a supersaturation with respect to the ZB solid phase. This does not guarantee that there is no thermodynamic driving force for other solid phases to form. According to the Gibbs phase rule, as many as six phases can coexist in a quaternary system. However, we assume that our system is at a non-equilibrium steady state where only the ZB phase forms from the supersaturated liquid and the formation of other solid phases (if they can exist at the given conditions) is kinetically inhibited. As far as possible, we will choose conditions based on the reported experimental results for the respective systems.

Results and discussion

In this section, we calculate and discuss the compositional diagrams for the four materials systems which are technologically highly relevant. The phase diagrams have been constructed using the Thermo-Calc software.²⁷ The list of the interaction parameters and the Gibbs energy functions is presented in the ESI.†

We start our investigation with the analysis of goldcatalyzed In_xGa_{1-x}As NWs. The InAs-GaAs pseudobinary interaction parameter is large and gives rise to a significant miscibility gap where the formation of the homogeneous solid solution is thermodynamically forbidden at the relevant growth temperatures. From the GaAs-InAs vertical section of the phase diagram of the In-Ga-As-Au system presented in Fig. 1 ($c_{\text{tot}} = 0.5$ and $c_{\text{As}} = 0.02$), it is seen that the In_xGa_{1-x}As phase exists over the entire range of relevant temperatures. Fig. 2 shows the liquid droplet composition versus the $In_xGa_{1-x}As$ solid composition at T = 477 °C and $c_{As} = 0.02$ in a wide range of the total concentrations of group III elements from $c_{\text{tot}} = 0.98$ (corresponding to self-catalyzed growth without gold) to $c_{\text{tot}} = 0.3$, where gold dominates in the catalyst droplet. It should be noted that, whereas the NW composition can be measured accurately using high-resolution electron microscopy techniques, 17 the composition of the droplet during growth is generally unknown. However, an experimental value of group III elements concentration in the Au-III droplet has been estimated to $c_{tot} = 0.5$ by Harmand et al.28 The amount of group V elements is very small due to the low solubility of arsenic in Au-III alloys. In spite of its negligible amount, the concentration of group V elements has drastic influence on the chemical potentials and, consequently, on the growth rates and crystal structures of ternary NWs.²⁹ As seen from Fig. 2, reasonable agreement between our analytical model and a purely thermodynamic phase segregation model (represented by the open circles) is observed. This model is based on segregation of a hypothetical, homogeneous supersaturated liquid with composition y into a solid phase with composition x, and a remaining liquid in equilibrium with the solid phase. The phase segregation data were evaluated using the Thermo-Calc software.

The slope of the y(x) curve is practically vertical over almost the entire ν range and changes only for $\nu > 0.97$. Therefore, very high y values are needed to tune the composition of In_xGa_{1-x}As NWs. As noticed above, a direct comparison bethe experimental and theoretical liquid-solid

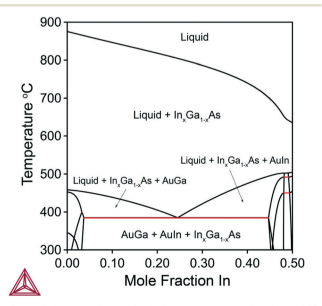


Fig. 1 Vertical section of the In–Ga–As–Au system at c_{tot} = 0.5 and c_{As} = 0.02.

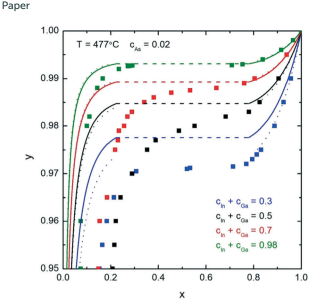


Fig. 2 Analytical calculations (solid curves) for the liquid-solid composition dependence for different $c_{\rm tot}$ at T=477 °C and $c_{\rm As}=0.02$. The dashed parts of the curves correspond to the miscibility gap. The small open circles correspond to phase segregation, evaluated using the Thermo-Calc software. The squares represent results from ref. 24.

composition dependences is seriously hampered because of in most cases unknown compositions of the catalyst particles. However, it is possible to describe the main observed trends. So, for example, to synthesize self-catalyzed $\text{In}_x\text{Ga}_{1-x}\text{As}$ NWs with x=0.03-0.05 as obtained in ref. 30, $y\approx 0.96-0.98$ is required. Moreover, the use of Au catalyst droplets allows one to increase the In fraction up to x=0.22. This is consistent with the obtained theoretical results: the In fraction in the nanowire decreases with increasing c_{tot} at constant y and T (Fig. 2).

The squares are the numerical results obtained using the previous two-component nucleation model including surface energies and a specific VLS term offsetting the chemical potential difference.²⁴ The main difference between that nucleation model and the one presented here is that the previous model that includes surface energies admits NW compositions within the miscibility gap. This is because the composition dependent surface energies influence the saddle point coordinates in such a way that it turns the two minima that would correspond to the miscibility gap compositions into one shallow minimum between these two minima.

The relationship between the solid and liquid compositions for different growth temperatures at the fixed $c_{\rm tot}$ = 0.5 is presented in Fig. 3. Clearly, the width of the miscibility gap shrinks with increased temperature and disappears completely at T=543 °C. This would enable the thermodynamically stable growth of ternary ${\rm In}_x{\rm Ga}_{1-x}{\rm As}$ NWs with any x value. However, such high temperatures can be unfeasible for MBE and MOVPE growth due to increased desorption and potential decomposition of III–V materials. Thus, as an example, thermodynamically stable growth of ${\rm In}_x{\rm Ga}_{1-x}{\rm As}$ NWs is not possible in the range 0.22 < x < 0.78 at T=477 °C (see Fig. 2).

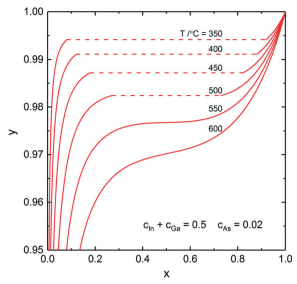
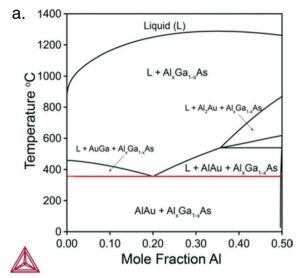


Fig. 3 Analytical calculations (solid curves) for the liquid-solid composition dependence for different temperatures at $c_{\rm in}+c_{\rm Ga}=0.5$ and $c_{\rm As}=0.02$. The dashed parts of the curves correspond to the miscibility gap.

To sum up the results of this section, the required *y* values for obtaining reasonable InAs fractions in ternary InGaAs NWs increase with the total concentration of group III elements, whereas the III/Au ratio does not change the miscibility gap region which is entirely determined by the temperature-dependent pseudobinary interaction parameter.

Next, we consider nucleation of gold-catalyzed Al_xGa_{1-x}As NWs. Fig. 4a shows a vertical section of the quaternary Al-Ga-As-Au system with $c_{\text{tot}} = 0.5$ and $c_{\text{As}} = 0.02$. While many different phases can be present in the case of Au-assisted NW growth, the Al_xGa_{1-x}As is a dominating solid phase at almost any Au-III ratios and in a wide range of NW growth temperatures. Fig. 4b shows a vertical section of the ternary Al-Ga-As system which is relevant for self-catalyzed growth of Al_xGa_{1-x}As NWs. Fig. 4a reveals that the minimum temperature for Au-catalyzed growth of Al_xGa_{1-x}As NWs should be higher than the eutectic point of the system (T = 353 °C). The minimum VLS growth temperature decreases as the Au concentration decreases. For the Al_xGa_{1-x}As system, the pseudobinary interaction parameter has a low enough value leading to the small dome height in comparison with the $In_xGa_{1-x}As$ system: the critical temperature at which the miscibility gap disappears equals -142°. This enables the fabrication of $Al_xGa_{1-x}As$ NWs with any solid composition at relevant MBE and MOVPE growth temperatures. The relationship between the solid and liquid compositions for different c_{tot} values at T= 610 °C is presented in Fig. 5. For obtaining self-catalyzed $Al_xGa_{1-x}As$ NWs with non-zero x, very low y values are necessary, while an almost horizontal slope cumbers precise compositional control. Indeed, a small addition of Al to the liquid results in a tremendous increase of the Al concentration in the solid. However, with decreasing c_{tot} , the slope of y(x)changes drastically and the required y values increase by several orders at high enough Au concentrations ($c_{tot} = 0.3$). The

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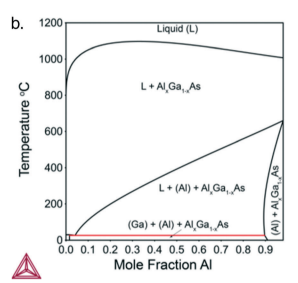


Fig. 4 a. Vertical section of the Al-Ga-As-Au system at $c_{tot} = 0.5$ and c_{As} = 0.02. b. Vertical section of the Al-Ga-As system at c_{tot} = 0.98 and $c_{As} = 0.02.$

change of temperature also influences y(x) but not so significant as c_{tot} does. As seen in Fig. 5, the results are well fitted by the one-parametric (ε) Langmuir expression given by x = $\varepsilon y/(1+y(\varepsilon-1))^{15}$, whereas tremendous discrepancy in comparison to the phase segregation approach (dotted curves) is observed.

Further, we investigate the nucleation of In_xGa_{1-x}Sb NWs from quaternary In-Ga-Sb-Au alloy. As seen from the vertical section of the In-Ga-As-Au system at $c_{\text{tot}} = 0.64$ and $c_{\text{Sb}} =$ 0.06 presented in Fig. 6, the compositional tuning of $In_xGa_{1-x}Sb$ throughout the entire compositional range is possible at T < 430 °C, which is the lowest temperature of the liquid single phase region. In comparison to As-based system, a relatively high Sb concentration $c_{Sb} = 0.06$ is needed to ensure positive difference of chemical potentials and, consequently, a positive supersaturation. Fig. 7 shows the liquid

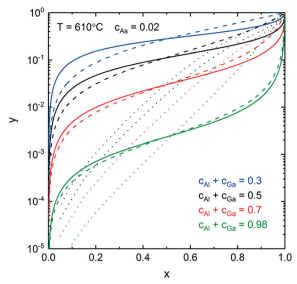


Fig. 5 Analytical calculations for the liquid-solid composition dependence of $Al_xGa_{1-x}As$ NWs for different c_{tot} at T = 610 °C and c_{As} = 0.02. The dotted curves are obtained using the phase segregation model. The dashed curves correspond to the model based on the oneparametric Langmuir expression with ε = 425 for c_{tot} = 0.98, ε = 32 for $c_{\rm tot}$ = 0.7, ε = 7.8 for $c_{\rm tot}$ = 0.5, and ε = 1.43 for $c_{\rm tot}$ = 0.3.

composition at $c_{\text{tot}} = 0.64$ and $c_{\text{Sb}} = 0.06$ plotted against the solid composition for different growth temperatures. The blue dashed curve at T = 450 °C indicates the absence of the solid phase and corresponds to the In molar fraction range within the 0.31-0.54 range on the GaSb-InSb vertical section of the In-Ga-Sb-Au phase diagram. When y decreases below 0.47, the solid phase occurs again (not shown in Fig. 7). The black dotted curve corresponds to the phase segregation model and is obtained using the Thermo-Calc software at T =430 °C, c_{tot} = 0.64, and c_{Sb} = 0.06. It is seen that the analytical

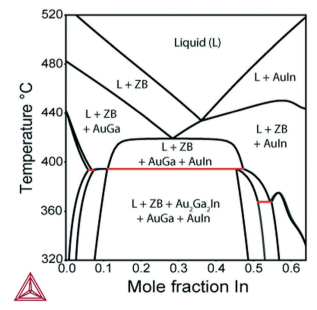


Fig. 6 Vertical section of the In-Ga-Sb-Au system at $c_{\text{tot}} = 0.64$ and $c_{Sb} = 0.06$

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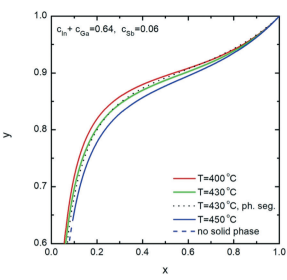


Fig. 7 Analytical calculations (solid curves) for the liquid-solid composition dependence of $In_xGa_{1-x}Sb$ NWs for different temperatures at $c_{In}+c_{Ga}=0.64$ and $c_{Sb}=0.06$. The dashed blue curve indicates the absence of solid $In_xGa_{1-x}Sb$ at these y-values. The dotted curve is obtained using the phase segregation model (T=430 °C).

model given by eqn (8) agrees well with the results of the phase segregation model. Fig. 8 shows the comparison of exact [given by eqn (8)] and approximate [eqn (11) and (12)] solutions for different total concentrations of group III elements and demonstrates a good agreement between them, especially for high x values. However, a significant divergence between exact and approximate solutions is observed for self-catalyzed growth.

Finally, we analyze nucleation of $InSb_xAs_{1-x}$ NWs from the In-Sb-As-Au melt. According to the phase diagram for the In-Sb-As-Au system presented in Fig. 9, the VLS growth of

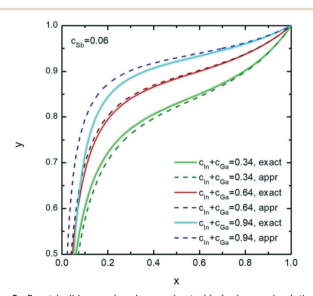


Fig. 8 Exact (solid curves) and approximate (dashed curves) solutions representing the liquid–solid composition dependences of solid $ln_xGa_{1-x}Sb$ for different c_{tot} values at T=430 °C and $c_{As}=0.06$.

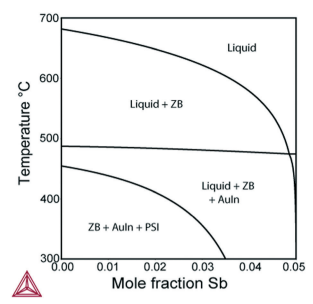


Fig. 9 Vertical section of the In–Sb–As–Au system at $c_{\rm In}$ = 0.45 and $c_{\rm tot}$ = 0.05.

InSb_xAs_{1-x} NWs is possible with any solid composition at T=450 °C. The main feature of this system is the compositional-dependent pseudobinary interaction parameter. This results in a composition-dependent, asymmetric miscibility gap. Fig. 10 shows the liquid composition *versus* the InSb_xAs_{1-x} solid composition for different temperatures in the cases of self-catalyzed ($c_{\text{tot}}=0.05$ and $c_{\text{In}}=0.95$) and gold-catalyzed growth at T=450 °C ($c_{\text{tot}}=0.05$ and $c_{\text{In}}=0.45$). Defect-free, self-catalyzed NWs with an Sb fraction greater than x=0.35 have been obtained at T=520 °C, as reported in ref. 32. Such a relatively high value corresponds to $y\approx0.975$. Clearly, it is

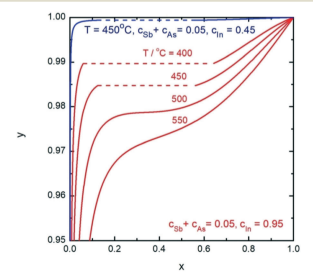


Fig. 10 Analytical calculations (solid curves) representing the liquid-solid composition dependences of solid $InSb_xAs_{1-x}$ NWs at $c_{1n}=0.95$ for different temperatures (red curves) and $c_{1n}=0.45$ at T=450 °C (blue curve). The dashed part of the curves correspond to the miscibility gap.

more difficult to control the NW composition in the case of Au-catalyzed growth because of the much steeper slope of the y(x) curve. Interestingly, in the case of $Al_xGa_{1-x}As$, the influence of the total concentration is reversed so that composition control would be easier in the Au-catalyzed case, as seen from comparing Fig. 5 and 10.

Conclusions

In summary, we have calculated the nucleation-limited compositions of ternary III-V NWs forming from quaternary alloys with gold. The calculations are based on realistic thermodynamic description and take into account ternary and composition-dependent binary interactions. We have analyzed the compositional diagrams for four highly relevant III-V materials systems. The results of this study show that the composition of ternary III-V NWs can be controlled over a wide range by tuning the composition of the catalyst droplet and temperature except for systems with high pseudobinary interaction parameters. For such systems (In_xGa_{1-x}Sb and In_xGa_{1-x}As), formation of ternary NWs is thermodynamically forbidden in a wide range of solid compositions at relevant temperatures due to the miscibility gaps. In addition, we demonstrated that in many cases our analytical calculations are in good agreement with a purely thermodynamic phase segregation model.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

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