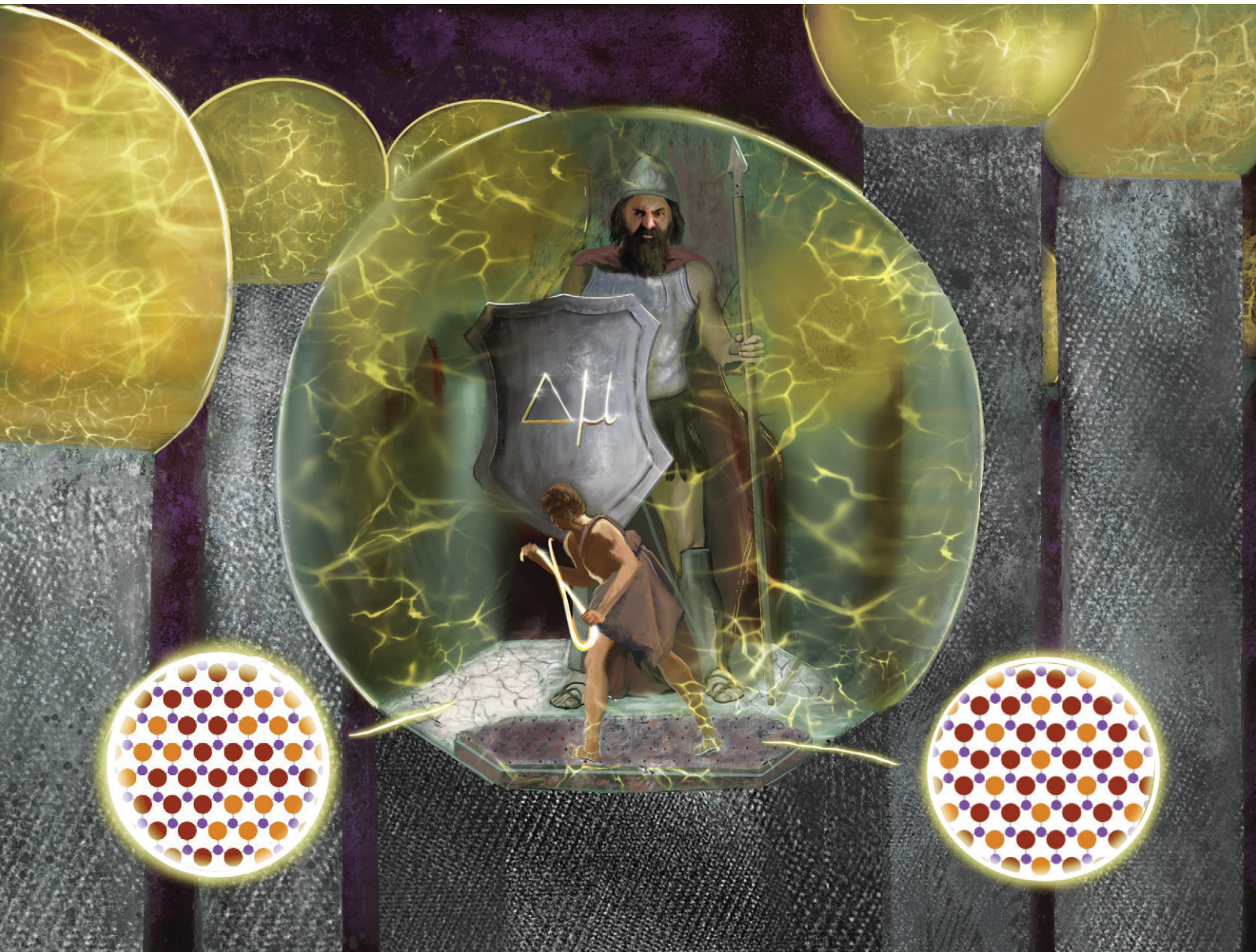


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## Surface energy driven miscibility gap suppression during nucleation of III–V ternary alloys

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The existence of a miscibility gap limits the range of solid compositions which are possible to achieve under near-equilibrium growth conditions. Circumventing the miscibility gap is of paramount importance for device fabrication. We propose that one of the suppression mechanisms is the nucleus surface energy and explain how the surface energy influences the miscibility gap during nucleation from a liquid melt. By doing this, we start with the formation of the critical nucleus of a ternary alloy considering its surface energy as a combination of the binary ones. For certain nucleation scenarios in the InGaAs materials system we show that the compositional independence of the surface energy term is a good approximation due to similar values for InAs and GaAs pairs. However, a large difference in the surface energies of the two materials may lead to a significant modification of the liquid-solid composition dependence and a complete suppression of the miscibility gap.

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### Introduction

Growth of ternary alloys formed by mixing two binary compounds is of paramount importance in materials science and engineering.<sup>1</sup> This is due to the possibility to tune one or more properties (band gap and lasing wavelength, lattice constant, magnetic properties) of the structure by varying its composition. Among structures of ternary alloys of different dimensions and morphology, ternary nanowires<sup>2–4</sup> have attracted great attention due to a broad spectrum of its possible application in optoelectronics, biotechnology and energy harvesting.<sup>5</sup> In contrast to elemental and binary compound semiconductors, the emission wavelength of optoelectronic devices based on ternary alloys can be adjusted by choosing from a continuous range determined by the two binaries.<sup>6,7</sup> Mastering the optoelectronic device design is impossible without an accurate composition control. Nevertheless, having the ideal growth technique, in many cases not all the solid compositions can be achieved for all materials combinations because of the miscibility gap, resulting in a wide range of thermodynamically forbidden compositions.<sup>8</sup> The reason for the miscibility gap is a net repulsion between dissimilar atoms and is, in the regular solution model,<sup>9</sup> described by the pseudobinary interaction parameter in the solid,  $\omega_s$ . It can readily be shown that the miscibility gap appears in ternary alloys with  $\omega_s > 2RT$ , where  $R$  is the gas constant and  $T$  is the temperature in Kelvin.<sup>10</sup>

The composition tuning range is limited in (In,Ga)As (ref. 11) and (In,Ga)N (ref. 12) alloys due to prominent miscibility gaps. In some cases such as In(As,Sb), the miscibility gap is shifted and non-symmetrical because of the composition-dependent pseudobinary interaction parameter.<sup>13</sup> The miscibility gap shrinks with temperature. However, high growth temperatures are often unfeasible because of the increased decomposition rate of the solid. For example, the miscibility gap in InGaAs materials system disappears at 543 °C,<sup>14</sup> while typical growth temperature of InGaAs nanowires by the molecular beam epitaxy and vapor phase epitaxy methods is in the range of 400–500 °C.<sup>4</sup> Thus, a large composition range of certain ternary alloys is thermodynamically forbidden. Understanding how to tune the composition of solid solutions over the entire range is needed to improve the quality of modern devices and to design promising structures such as vertical quantum wires.<sup>15</sup>

Considering the immiscibility in bulk alloys,<sup>16</sup> growth of nanostructures with compositions inside the miscibility gap is often possible *via* formation of metastable states. For example, the miscibility gap in thin films can be suppressed by elastic stress.<sup>17,18</sup> In the case of growth of axial nanowire heterostructures at temperatures lower than the critical one,<sup>14</sup> the wide range of thermodynamically forbidden compositions should lead to the formation of an atomically abrupt interface. However, a compositionally graded interface over the entire compositional range is observed in heterostructured nanowires of materials systems with a high pseudobinary interaction parameter which proves either the metastable state formation inside the miscibility gap, or the suppression of the miscibility gap.<sup>19,20</sup> Despite a number of

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experimental investigations on the growth of nanowires with composition inside the miscibility gap,<sup>21,22</sup> the suppression mechanism in these nanostructures remain unclear. Some kinetic theories<sup>23,24</sup> assume that the miscibility gap might shrink under relatively high supersaturations in the liquid. Indeed, it has been shown that high concentrations of group V elements in the droplet can “straighten” the initially S-shaped liquid–solid composition dependence curve. Another possible explanation is the size-dependent effect of spinodal decomposition,<sup>25</sup> or the chemical-potential-dependent surface reactions resulting in a miscibility gap shrinkage with reducing particle size.<sup>26</sup>

The composition-independence of the surface energy term of the critical nucleus is a standard assumption substantiated by the investigation by Wilemski,<sup>27</sup> where it was assumed that the component with the lower surface energy segregates to the surface so readily so that the nucleus always obtains the surface energy of that component. This is used in the majority of kinetic models based on the incorporation rate,<sup>23,24,28</sup> mass balance models,<sup>29</sup> equilibrium models<sup>30</sup> and nucleation-limited models.<sup>14,31</sup> The omitted surface energy term is a good approximation for the relatively large size of the stable growing clusters, making the surface energies insignificant. Using the Butler equations<sup>32</sup> for calculation of the composition-dependent interface energies, the robustness of the existence of the saddle point is presented in.<sup>33</sup> Finally, the effect of a composition-dependent edge energy on the interface profiles of the axial heterostructures in nanowires has been studied by Glas<sup>34</sup> for the (Al,Ga)As materials system, in which the growth temperatures are far above the miscibility gap.

In the current paper, we aim to analytically explore the miscibility gap suppression driven by the surface energy of the ternary nucleus. The nucleus surface energy is introduced in a general way giving an opportunity to consider the Vegard's law case, the Wilemski approach, and their intermediate cases as well as a convex (second degree polynomial) model which resembles the Butler equations case. We construct the miscibility gap and analyse it with respect to the composition of group V elements and the effective surface energy ratio.

There are a number of simplifications and assumptions utilized in this paper. (i) The investigation is based on classical nucleation theory and implies the capillarity approximation<sup>35–37</sup> within which molecular clusters are treated as macroscopic objects. This is a basic model and might lead to some invalid results such as overestimation of the nucleation rate of crystalline monolayers.<sup>38</sup> No renormalization because of the finite size of the considered system is utilized. Such self-consistency renormalization of the Zeldovich nucleation rate for calculation of nanowire elongation rates and description of crystal structure can be found in.<sup>39</sup> The critical nucleus size depending on the supersaturation could be relatively large (>200 atoms (ref. 33)). However, more accurate calculations involve quantum corrections and consideration of the nanoscale nature.<sup>40,41</sup>

(ii) The Gibbs–Thomson effect of elevated chemical potential in the droplet due to a curved surface<sup>42</sup> is not considered. Its effect can often be ignored for liquid particles with sizes larger than about 20 nm. (iii) The obtained results describe the composition of the critical nucleus. Thus, it applies to nucleation-limited growth. At high supersaturation the final composition might differ from the one of the critical nucleus.<sup>24</sup> (iv) Finally, there is an uncertainty in the estimated values of the surface energies and the concentration of group V elements in the liquid. Despite these shortcomings, classical nucleation theory has proven to be valuable for modelling and understanding several aspects of nanowire growth.<sup>14,31,43,44</sup>

Due to the theoretical nature of our work, we are able to vary the parameters whose values are very difficult to study experimentally, namely the surface energy ratio and concentration of group V elements. This allows us to discuss some general trends, which can be of high value to experimentalists. Moreover, we have estimated the effect of the uncertainty of surface energy values on the liquid–solid composition dependence and miscibility gap.

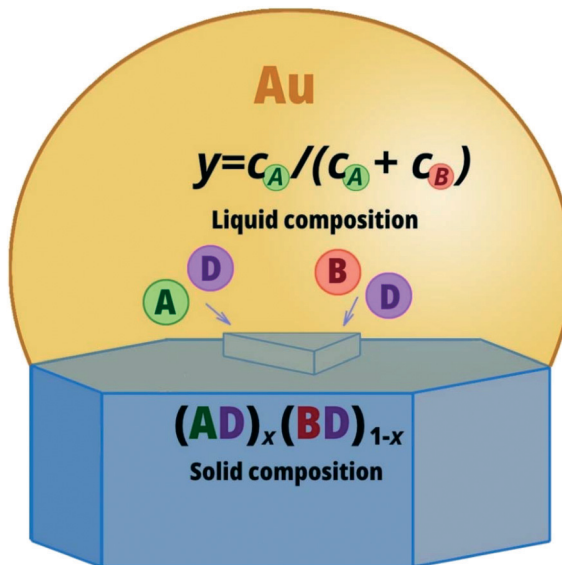
## Calculations

Conceptually, there are two main approaches to fabricate micro- and nanostructures.<sup>45</sup> The first one is the top-down approach which implies the surface patterning by electron beam lithography,<sup>46</sup> nanoimprint lithography<sup>47</sup> and other methods followed by etching. The second one is the bottom-up approach in which structures are synthesized by stacking atoms of provided semiconductor materials onto each other. Nucleation occurs from the liquid phase within vapor–liquid–solid growth or from the vapor within vapor–solid–solid growth.

To fabricate ternary alloy structures, bottom-up approach with the involved liquid phase is the most suitable because the solid composition can be accurately controlled by composition of the liquid while the layer-by-layer growth process provides excellent crystal quality. Therefore, in this paper we consider the formation of a critical nucleus of a ternary  $A_xB_{1-x}D$  alloy from a quaternary liquid droplet which contain A, B, D, and U, where U is gold or another catalyst. The foreign catalyst changes the chemical potential in the liquid and simplifies nanostructure growth. In the context of nanowires, liquid catalyst droplets act as collection sites for semiconductor material resulting in selective growth which can be ordered if the patterned mask is used or randomly distributed otherwise. It is also possible to synthesize nanowires without a foreign metal. This is the so called self-catalyzed growth<sup>48</sup> where the liquid droplets contain only the group III nanowire constituent and small amount of group V constituent.

Nucleation is schematically presented in Fig. 1. Nucleation occurs at the liquid–solid interface while the ratio of incorporated A and B atoms determines the solid composition  $x$ . The presented model allows us to discuss growth of both bulk structures and nanowires from the liquid





**Fig. 1** Schematic illustration of the formation of an  $A_xB_{1-x}D$  nucleus from the quaternary liquid phase at nucleation in the center of the liquid–solid interface.

phase. Assuming the droplet and substrate to be “infinitely” large, the theory can be applied for the description of bulk structures grown by the liquid phase epitaxy.<sup>49</sup> For the nanoscale droplet particles, the model is applicable for nanowires grown *via* the vapor–liquid–solid (VLS) mechanism.<sup>50</sup> This is an attractive bottom-up approach of nanostructure fabrication that involves nucleation and growth from supersaturated liquid droplets. The difference between nucleation at the triple phase line and in the center of the liquid–solid interface is determined by the vertical edge of the 2D nucleus. It is a weighted sum of the vapor–solid, liquid–solid and liquid–vapor surface energies at the triple phase line nucleation, while it is a function of the liquid–solid surface energy in the case of nucleation in the center of the liquid–solid interface (see Appendix A).

The change of Gibbs energy during nucleation is composed of three parts. The first one decreases linearly with the nucleus size  $s$  with the chemical potential as proportionality factor. The second term increases with the nucleus perimeter due to the formation of new surfaces. The last term is VLS specific and refers to the transformation of the liquid drop surface due to the nucleus formation and is often ignored because of its small value. Thus, the nucleus formation energy is given by

$$F = -\Delta\mu s + \alpha\sqrt{s}. \quad (1)$$

Here  $\alpha$  is the effective surface energy of the nucleus of a ternary alloy and  $\Delta\mu$  is the chemical potential difference between the quaternary liquid melt and the solid (see Appendix A). The form of the chemical potentials which contain the ternary and composition-dependent binary interaction parameters can be found in.<sup>31</sup>

According to classical, two-component nucleation theory, the size and composition of a critical nucleus can be found from the simultaneous solution of the following system of differential equations:

$$\frac{\partial F}{\partial x} = -\frac{\partial\Delta\mu}{\partial x}s + \frac{da}{dx}\sqrt{s} = 0, \quad (2)$$

$$\frac{\partial F}{\partial s} = -\Delta\mu + \frac{a}{2\sqrt{s}} = 0. \quad (3)$$

The critical size is a result of maximization of the formation energy in the size and can be expressed from eqn (3), yielding  $\sqrt{s} = a/(2\Delta\mu)$ . Then, the nucleation barrier required for the formation of the nucleus of the critical size is  $F_* = a^2/4\Delta\mu$ . Now, one should minimize the nucleation barrier  $F_*$  in the solid composition. This procedure is equivalent to elimination of the nucleus size in eqn (2), and gives the relationship between the terms related to the phase transition and the nucleus surface energy in the form

$$\frac{\partial\Delta\mu}{\partial x} = \Delta\mu \frac{2}{a} \frac{da}{dx}. \quad (4)$$

As a consequence of the chemical potential being a partial molar property of a binary system, the derivative of the chemical potential difference with respect to solid composition equals the chemical potential difference of the AD ( $\Delta\mu_{AD}$ ) and BD ( $\Delta\mu_{BD}$ ) pairs,  $\partial\Delta\mu/\partial x = \Delta\mu_{AD} - \Delta\mu_{BD}$ . Thus, eqn (4) might be reduced to

$$\frac{\Delta\mu_{AD}}{\Delta\mu_{BD}} = 1 + \frac{\frac{2}{a} \frac{da}{dx}}{1 - \frac{2}{a} \frac{da}{dx} x}. \quad (5)$$

In this form, it is clearly seen that in the general case, there is no equality between the chemical potential difference of AD and BD pairs. This equality (being equivalent to  $\partial\Delta\mu/\partial x = 0$  considered in ref. 14 and 31) is only the case if  $da/dx = 0$ .

To combine all the considered cases, the surface energy of the ternary nucleus can be presented as

$$a = a_{BD} + x\Delta a + g(x), \quad (6)$$

with  $\Delta a = a_{AD} - a_{BD}$ . The composition-dependent function  $g(x)$  defines a particular model. So, a linear relationship can be obtained at

$$g_\delta(x) = \delta\Delta a(1 - x), \quad (7)$$

It is clearly seen that the Vegard's law case  $a = a_{AD}x + a_{BD}(1 - x)$  is implemented at  $\delta = 0$ , while the Wilemski case  $a = a_{AD}$  corresponds to  $\delta = 1$ . All the linear intermediate cases lying between the first and second curves can be described by  $0 < \delta < 1$ .

For the convex relationship, the function is given by

$$g_\alpha(x) = \alpha\Delta a x(1 - x), \quad (8)$$



where the bowing parameter  $\alpha$  defines the quadratic term and, thus, describes the deviation from linearity. Then the derivatives of the nucleus surface energy with respect to the solid composition are

$$\frac{da}{dx} = \Delta a(1 - \delta), \quad (9)$$

$$\frac{da}{dx} = \Delta a(1 + \alpha(1 - 2x)), \quad (10)$$

In the linear case of the Vegard's law ( $\delta = 0$ ), the derivative of the nucleus surface energy with respect to the solid composition is the surface energy difference of the binaries  $da/dx = \Delta a$ . In this case the solid composition of a ternary alloy can be found from the following equation

$$\begin{aligned} (\Delta\mu_{AD}^y a_{BD} - \Delta\mu_{BD}^y (a_{AD} + \Delta a)) - (\Delta\mu_{AD}^y - \Delta\mu_{BD}^y) x \Delta a \\ = (RT \ln x + \omega_s(1-x)^2)(a_{BD} - x \Delta a) \\ - (RT \ln(1-x) + \omega_s x^2)(a_{AD} + (1-x) \Delta a). \end{aligned} \quad (11)$$

Eqn (11) can be written as  $f(y) + u(y)x = g(x)$ . The superscript  $y$  in the chemical potential difference is explained in Appendix A. This mixing of solid and liquid compositions results in a complex dependence where the miscibility gap is a function of several parameters including As and Au concentrations in the droplet, growth temperature and the ratio of the binary surface energy terms. The In/III concentration ratio the liquid is denoted by  $y$ .

Considering the nucleus surface energy, the miscibility gap is shifted and not symmetrical around  $x = 0.5$  anymore. Its width at the fixed liquid composition can be found using the Maxwell construction or the Hessian determinant,

$$\det H = \begin{vmatrix} \frac{\partial^2 F}{\partial x^2} & \frac{\partial^2 F}{\partial x \partial s} \\ \frac{\partial^2 F}{\partial s \partial x} & \frac{\partial^2 F}{\partial s^2} \end{vmatrix}, \quad (12)$$

which in our case is given by

$$\det H = \frac{a}{4\sqrt{s}} \left( \frac{\partial^2 \Delta\mu}{\partial x^2} - \frac{1}{\sqrt{s}} \frac{d^2 a}{dx^2} \right) - \left( -\frac{\partial \Delta\mu}{\partial x} + \frac{1}{2\sqrt{s}} \frac{da}{dx} \right)^2. \quad (13)$$

In eqn (13),  $\partial^2 \Delta\mu / \partial x^2 = 2\omega_s - RT/(x(1-x))$ ,  $\sqrt{s} = a/(2\Delta\mu)$ , and  $d^2 a / dx^2 = 0$  for the linear model and  $d^2 a / dx^2 = -2\alpha\Delta a$  for the convex model. The spinodal is defined by the equation  $\det H = 0$ . The maximum of the Hessian determinant as a function of  $x$  corresponds to the largest instability of the system and gives the center of the miscibility gap.

The critical temperature  $T_c$  corresponds to the critical point where the binodal and spinodal curves coincide. In the case of  $x$ -independent nucleus surface energy,  $2RT_c = \omega_s$  ( $T_c$ ). Otherwise, to find the critical temperature as a function of the effective surface energy ratio at fixed concentrations of  $D$  and  $U$ , it is necessary to solve the equations  $d(\det H)/dx = 0$  and  $\det H = 0$  simultaneously with  $x$  being the composition of the critical nucleus.

## Results and discussion

We start the discussion with an overview of the different approaches to the surface energy of an arbitrary ternary nucleus. All the main cases calculated at  $a_{BD}/a_{AD} = 2$  are summarized in Fig. 2. The first curve (i) represents the Vegard's law where the surface energy term is a linear combination of the binary ones implemented at  $\delta = 0$ . Another extreme case (ii) corresponds to  $\delta = 1$  meaning the independence of the surface energy on the solid composition ( $da/dx = 0$ ). This approach was introduced by Wilemski<sup>27</sup> and is motivated by a phase segregation situation where the binary with lower surface energy concentrates at the nucleus periphery. In spite of the lack of justification for its application for solids<sup>51</sup> it remains very popular for modelling vapor-liquid-solid nucleation due to its simplicity.<sup>14,24,31</sup> The curve (iii) corresponds to the intermediate case calculated at  $\delta = 0.5$ . Finally, a non-linear case can be considered within the convex model (iv) calculated at  $\alpha = 1$ , while the curve coincides with the Vegard's law case at  $\alpha = 0$ . Due to the quadratic form in the convex model at non-zero  $\alpha$ , the results are expected to be similar to the Butler equation case.<sup>33</sup>

The liquid-solid composition dependence can be obtained from eqn (11) by finding the crossings of functions  $f(y) + u(y)x$  and  $g(x)$ . Under the condition of  $a_{GaAs}/a_{InAs} = 1$ , the function  $g(x)$  coincides with the previously considered case of  $g(x) = RT \ln(x/(1-x)) - 2\omega_s(x - 1/2)$ <sup>14</sup> while the decrease or increase of the ratio determines shifting the function up or down correspondingly. Fig. 3 shows the functions  $f(y) + u(y)x$  and  $g(x)$  calculated for different  $y$  at fixed  $a_{GaAs}/a_{InAs} = 1.3$ ,  $T = 450$  °C,  $c_{As} = 0.01$  and  $c_{Au} = 0.28$ . As seen from Fig. 3, there is only one solution in a wide  $y$  range except of  $y \approx 0.9506$  near which the curve  $f(y) + u(y)x$  crosses the curve  $g(x)$  three times. At such liquid compositions there is a miscibility gap.

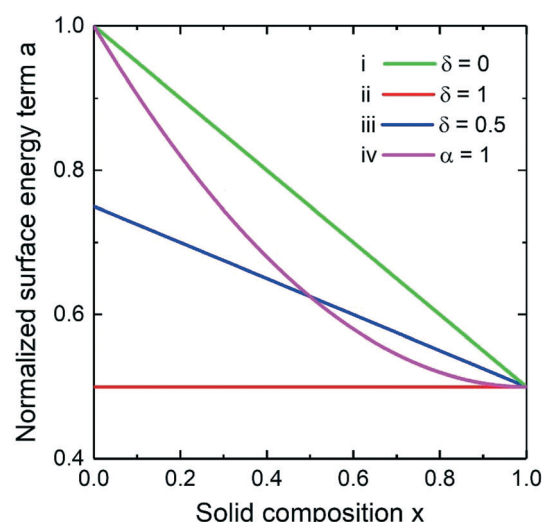
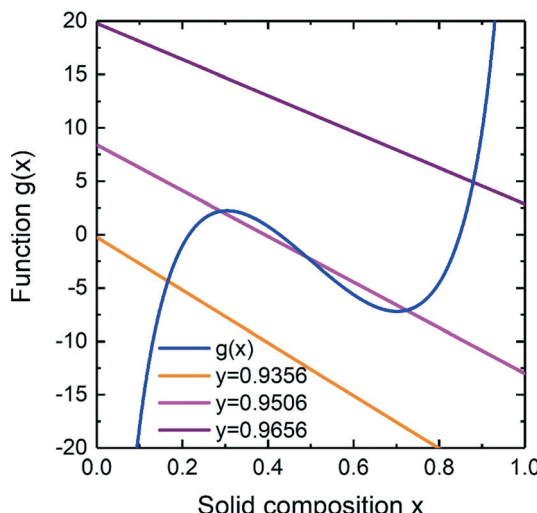


Fig. 2 Normalized surface energy term of a ternary nucleus for  $a_{BD}/a_{AD} = 2$  in the (i) Vegard's law case, (ii) Wilemski approach, (iii) their intermediate case and the convex model (iv).



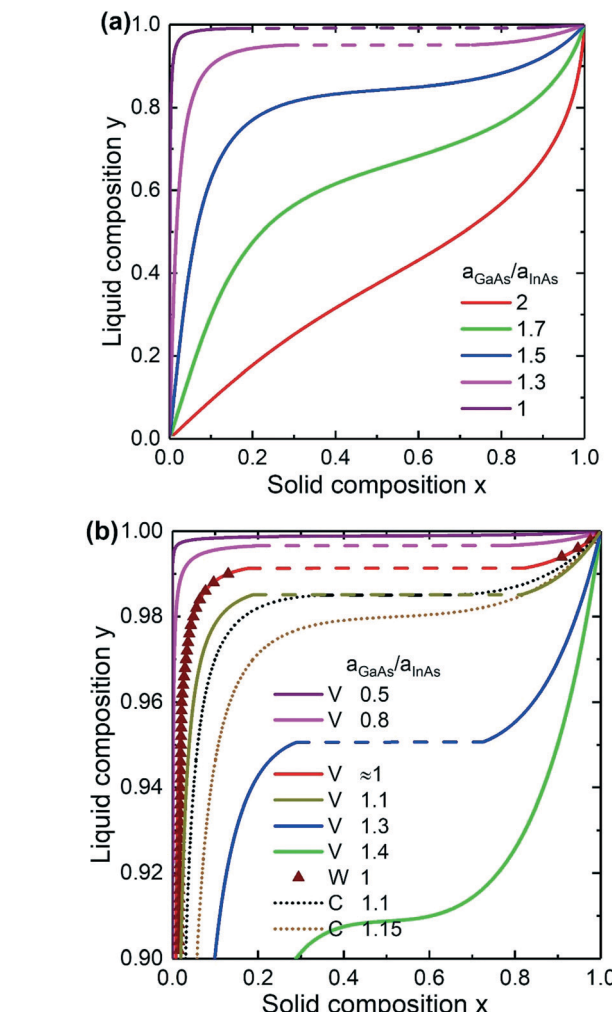


**Fig. 3** Illustration of the  $y(x)$  selection rule for InGaAs materials system. The blue solid curve is  $g(x)$  function; the tilted lines are  $f(y) + u(y)x$  calculated for different  $y$  and at fixed  $a_{\text{GaAs}}/a_{\text{InAs}} = 1.3$ .

Now let us consider the effect of the surface energy on the ternary nucleus formation in vapor–liquid–solid growth. Fig. 4a shows the liquid–solid composition dependence calculated within the Vegard's law for the critical  $\text{In}_x\text{Ga}_{1-x}\text{As}$  nucleus at different surface energies. The parameters are  $c_{\text{As}} = 0.01$ ,  $c_{\text{Au}} = 0.28$  and  $T = 450$  °C. It is seen that the liquid–solid composition dependence is significantly modified with increasing  $a_{\text{GaAs}}/a_{\text{InAs}}$ . It leads to a narrowing of the miscibility gap and to higher In content in the solid at fixed liquid composition. At the large enough ratio of  $a_{\text{GaAs}}/a_{\text{InAs}} = 1.5$ , the miscibility gap is completely suppressed, while at  $a_{\text{GaAs}}/a_{\text{InAs}} \approx 2$  the solid composition is almost equal to the liquid composition ( $x \approx y$ ).

A comparison of the composition dependences calculated within different models is presented in Fig. 4b. The parameters are the same as in Fig. 4a. For center nucleation<sup>43</sup> of a InGaAs nucleus, the values of the surface energy terms (see Appendix A for definitions) for InAs (ref. 52) and GaAs (ref. 53) pairs are similar, namely  $a_{\text{GaAs}}/a_{\text{InAs}} = 1.0002$ . Thus the liquid–solid composition dependence is expected to be identical to the one in the Wilemski approach.<sup>31</sup> Modelling of nucleation from a quaternary liquid melt hampers the calculations because the surface energy of GaAs pairs at the solid–liquid interface increases by an order of magnitude during the transition from Au-catalyzed growth to self-catalyzed one.<sup>43,54</sup> However, in this paper the surface energy at the solid–liquid interface is treated as independent on the group III concentration since this phenomenon is poorly studied.

As seen from Fig. 4b the composition dependences corresponding to the Vegard's law at  $a_{\text{GaAs}}/a_{\text{InAs}} = 1.0002$  and the Wilemski approach are almost identical. This justifies the assumption of composition-independence of the surface energy for the InGaAs materials system. It is remarkable that within Vegard's law, the miscibility gap disappears at surface energy ratios quite far from 1 ( $a_{\text{GaAs}}/a_{\text{InAs}} \approx 0.5$  and  $a_{\text{GaAs}}/a_{\text{InAs}} \approx 1.4$  under the chosen conditions), while the surface energy ratio has larger impact within the convex model. Here the solid composition can be tuned throughout the entire range at  $a_{\text{GaAs}}/a_{\text{InAs}} = 1.15$ . The shape of the curve is similar to the result obtained in<sup>33</sup> using the Butler equations at  $c_3 \equiv c_{\text{In}} + c_{\text{Ga}} = 0.7$ ,  $T = 477$  °C,  $c_{\text{As}} = 0.02$  and the ratio of the pure InAs and GaAs surface energies of 1.15. It is important to note that in the case of nucleation at the triple phase line,  $a_{\text{GaAs}}/a_{\text{InAs}} \approx 1.15$ , which was the considered case in ref. 33. So, the miscibility gap is slightly modified in the Vegards law case and completely suppressed within the convex model. Thus, at the triple phase line nucleation the choice of model is crucial.



**Fig. 4** Liquid–solid composition dependences of InGaAs alloy for different surface energy ratio calculated within the Vegard model (a) and within Vegard (V), Wilemski (W) and convex (C) models (b). Triangles correspond to the Wilemski approach. The solid curves are the Vegard's law case and dots are the convex model with  $\alpha = 0.5$ . The dashed lines correspond to the miscibility gap.

$a_{\text{InAs}} \approx 1.4$  under the chosen conditions), while the surface energy ratio has larger impact within the convex model. Here the solid composition can be tuned throughout the entire range at  $a_{\text{GaAs}}/a_{\text{InAs}} = 1.15$ . The shape of the curve is similar to the result obtained in<sup>33</sup> using the Butler equations at  $c_3 \equiv c_{\text{In}} + c_{\text{Ga}} = 0.7$ ,  $T = 477$  °C,  $c_{\text{As}} = 0.02$  and the ratio of the pure InAs and GaAs surface energies of 1.15. It is important to note that in the case of nucleation at the triple phase line,  $a_{\text{GaAs}}/a_{\text{InAs}} \approx 1.15$ , which was the considered case in ref. 33. So, the miscibility gap is slightly modified in the Vegards law case and completely suppressed within the convex model. Thus, at the triple phase line nucleation the choice of model is crucial.

As it is widely known, for a composition independent  $\omega_S$  the temperature of the binodal can be written as  $T_b = \omega_S(1 - 2x)/(R \ln((1 - x)/x))$ . Introducing the surface energy, it is impossible to explicitly express the binodal temperature as a

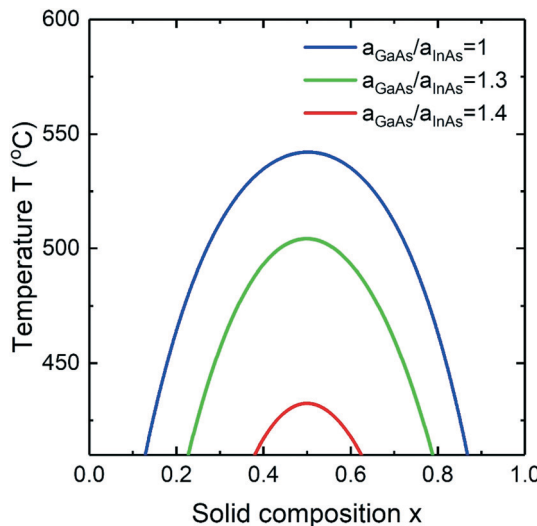


Fig. 5 The miscibility gap in  $\text{In}_x\text{Ga}_{1-x}\text{As}$  alloy at  $c_{\text{As}} = 0.01$  for different surface energy ratios.

function of solid composition in a similar closed form. Moreover, including the surface energies,  $T_b$  depends on the chemical potential in the liquid which is a strong function of group V element concentration. Fig. 5 shows the miscibility gap in an InGaAs alloy calculated for the Vegard's law case for different surface energy ratios at fixed  $c_{\text{As}} = 0.01$  and  $c_{\text{Au}} = 0.28$ . The largest binodal region corresponds to  $a_{\text{GaAs}}/a_{\text{InAs}} = 1$  and narrows the more this ratio deviates from 1. So, the critical temperature above which all the solid compositions are stable is  $T = 543$  °C at  $a_{\text{GaAs}}/a_{\text{InAs}} = 1$ ,  $T \approx 505$  °C at  $a_{\text{GaAs}}/a_{\text{InAs}} = 1.3$  and  $T = 432$  °C at  $a_{\text{GaAs}}/a_{\text{InAs}} = 1.4$ .

The dependence of the miscibility gap of InGaAs alloy on the As concentration at  $a_{\text{GaAs}}/a_{\text{InAs}} = 1.3$  and  $c_{\text{Au}} = 0.28$  is presented in Fig. 6. As seen with increasing As concentration the miscibility gap decreases. Changing the As concentration by an order of magnitude from  $c_{\text{As}} = 0.01$  to  $c_{\text{As}} = 0.1$  results

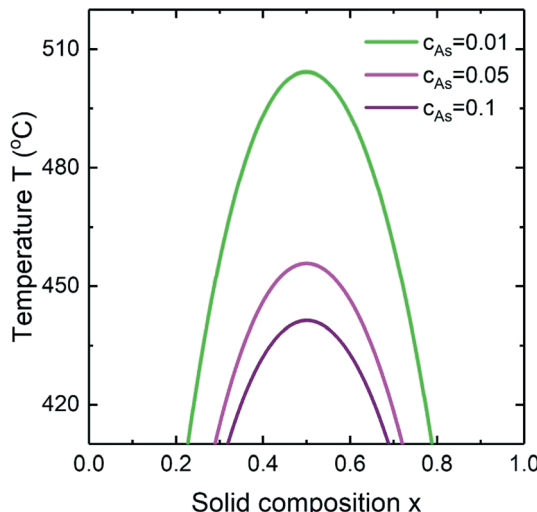


Fig. 6 The miscibility gap in InGaAs alloy at  $a_{\text{GaAs}}/a_{\text{InAs}} = 1.3$  for different As concentrations.

in shift of the critical temperature by 64 °C from  $T \approx 505$  °C to  $T \approx 441$  °C. Simplifying, the chemical potentials in the solid  $\mu_{\text{BD}}^s$  and  $\mu_{\text{AD}}^s$  are responsible for the binary separation and the appearance of the miscibility gap in a ternary alloy. Their difference is an essential part of the chemical potential term in the nucleus formation energy and has the largest impact on the thermodynamically forbidden compositional range. By including a composition dependence of the nucleus surface energy in the formation energy, the miscibility gap might be reduced in two possible ways. They are the increase of the second term in eqn (1) with raising the difference of the interface energies of the two binaries (Fig. 5) and the decrease of the first term in eqn (1) with raising the group V concentration in the liquid which leads to higher supersaturation (Fig. 6). Thus, even in ternary alloys with a high value of the pseudobinary interaction parameter, the miscibility gap might be suppressed completely at temperatures relevant for growth. It is interesting that the surface energy driven suppression mechanism acts at the nucleation stage in contrast to the kinetic growth picture previously discussed.<sup>23,24,28</sup>

The critical temperature  $T_c$  at which the miscibility gap vanishes is determined by both the composition of the liquid and the effective surface energy ratio. Fig. 7 shows the dependence of the critical temperature on  $a_{\text{GaAs}}/a_{\text{InAs}}$  for the InGaAs nucleus at fixed  $c_{\text{Au}} = 0.28$  calculated for different As concentrations within the Vegard's law model. It is clearly seen that both curves reach the maximum at equal surface energies ( $a_{\text{GaAs}}/a_{\text{InAs}} = 1$ ). Then the critical temperature decreases rapidly and almost symmetrically on both sides of  $a_{\text{GaAs}}/a_{\text{InAs}} = 1$ . So, if  $a_{\text{GaAs}}/a_{\text{InAs}} \approx 1.38$  and  $c_{\text{As}} = 0.01$ , the miscibility gap disappears at  $T_c = 450$  °C which is a relevant growth temperature for vapor phase deposition. The U-shaped curve of the critical temperature against effective

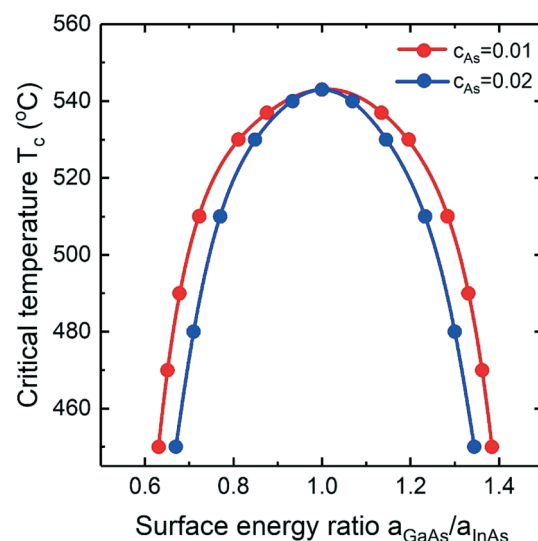


Fig. 7 Critical temperature versus the ratio of binary surface energy terms for InGaAs materials system at fixed As concentration of  $c_{\text{As}} = 0.01$  (red curve) and  $c_{\text{As}} = 0.02$  (blue curve). The dots represent numerical calculations and the solid curves are polynomial fits.



surface energy ratio explains the liquid–solid composition dependence presented in Fig. 4b: the solid phase exists over the entire compositional range at both high and low enough values of the effective surface energy ratio (the violet and green solid lines). Increasing the As concentration, the critical temperature decreases.

Finally, we discuss the impact of the uncertainty of the surface energies. The estimated error of the surface energy found in the literature is about 10%.<sup>55</sup> It influences both the critical nucleus size which is not calculated here and the liquid–solid composition dependence and the miscibility gap through the ratio of the surface energies. Assuming the error is 10%, the surface energy ratio is in a range of 0.9–1.1. As seen from Fig. 4b (compare the solid red and dark yellow curves), the liquid–solid composition dependence is slightly shifted to the bottom while the miscibility gap and the critical temperature (see Fig. 7) are almost the same. However, within the convex model (compare the solid red and dotted black curves), the miscibility gap is twice as small. Thus, in the context of the miscibility gap, the Vegard's law case is less sensitive to the surface energy error than the convex model.

## Conclusions

To summarize, the surface energy contribution in the formation energy of a two-component ternary alloy nucleus is considered. By doing this, we solve the system of equations which define the saddle point of the nucleus formation energy and eliminate the nucleus size. Within this model based on two-component nucleation theory, the composition of ternary alloys nucleating from a liquid melt is obtained as a function of the composition of the melt. The surface energy term of the ternary nucleus is a combination of the corresponding binary terms. In this regard, we analyse one convex model and three linear models, namely the Vegard's law case, the Wilemski approach and their intermediate case. Modelling the example of InGaAs VLS nanowire system shows that at the central nucleation the surface energy contribution is small and almost does not change the miscibility gap and the liquid–solid composition dependence because of the similar values of InAs and GaAs surface energy terms. This justifies the usage of the Wilemski approach which simplifies the calculations. For nucleation at the triple phase line, the convex and linear models give different results: in contrast to the Vegard's law case, the miscibility gap is completely suppressed within the convex model at  $a_{\text{GaAs}}/a_{\text{InAs}} = 1.15$ . In any case, a sufficient increase of the surface energy ratio results in a complete suppression of the miscibility gap. The model might be useful for the optimization of growth parameters for the fabrication of III–V semiconductor nanowires and bulk structures growing from the liquid phase with a specified ternary composition. Thus, the model could be applied to  $\text{In}_x\text{Ga}_{1-x}\text{N}$ ,  $\text{InAs}_x\text{Sb}_{1-x}$  or  $\text{In}_x\text{Ga}_{1-x}\text{Sb}$  alloys for instance. By doing this, one should find and substitute in the corresponding places of eqn (11)–(13)

the values of Gibbs free energies, interaction parameters and surface energies of the chosen material system. Moreover, the model might be used for modelling solid composition of material systems without miscibility gap at relevant growth temperatures because a large surface energy contribution may modify the liquid–solid composition dependence. With relevant chemical potentials and surface energies, our approach can also be generalized to other scenarios, such as direct vapor–solid nucleation and growth. However, care should be taken when applying the model and the obtained results because of the uncertainty in the surface energies.

## Appendix A

The difference of the chemical potentials between liquid and solid is given by

$$\Delta\mu = x\Delta\mu_{\text{AD}} + (1-x)\Delta\mu_{\text{BD}}, \quad (\text{A1})$$

where the chemical potential differences of AD and BD pairs in the liquid and in the solid state are given by

$$\Delta\mu_{\text{AD}} = \mu_{\text{A}}^{\text{L}} + \mu_{\text{D}}^{\text{L}} - \mu_{\text{AD}}^{\text{S}}, \quad (\text{A2})$$

$$\Delta\mu_{\text{BD}} = \mu_{\text{B}}^{\text{L}} + \mu_{\text{D}}^{\text{L}} - \mu_{\text{BD}}^{\text{S}}. \quad (\text{A3})$$

The chemical potentials in the liquid of the corresponding component are given by

$$\mu_{\text{A}}^{\text{L}} = \mu_{\text{A}}^0 + RT \ln c_{\text{A}} + \varphi_{\text{A}}, \quad (\text{A4})$$

$$\mu_{\text{B}}^{\text{L}} = \mu_{\text{B}}^0 + RT \ln c_{\text{B}} + \varphi_{\text{B}}, \quad (\text{A5})$$

$$\mu_{\text{D}}^{\text{L}} = \mu_{\text{D}}^0 + RT \ln c_{\text{D}} + \varphi_{\text{D}}. \quad (\text{A6})$$

Here  $\mu^0$  is the Gibbs free energy of the corresponding pure element,  $\varphi$  is a sum of products of corresponding interaction parameters and concentrations.<sup>23,31</sup>

The chemical potentials in the solid of AD and BD binary species are given by

$$\mu_{\text{AD}}^{\text{S}} = \mu_{\text{AD}}^0 + RT \ln x + \omega_{\text{S}}(1-x)^2, \quad (\text{A7})$$

$$\mu_{\text{BD}}^{\text{S}} = \mu_{\text{BD}}^0 + RT \ln(1-x) + \omega_{\text{S}}x^2, \quad (\text{A8})$$

with  $\omega_{\text{S}}$  being the pseudobinary interaction parameter and  $\mu^{\text{S}}$  is the Gibbs free energy of the corresponding binary compound. We use the following notation:

$$\Delta\mu_{\text{AD}}^{\text{S}} = \mu_{\text{A}}^{\text{L}} + \mu_{\text{D}}^{\text{L}} - \mu_{\text{AD}}^0, \quad (\text{A9})$$

$$\Delta\mu_{\text{BD}}^{\text{S}} = \mu_{\text{B}}^{\text{L}} + \mu_{\text{D}}^{\text{L}} - \mu_{\text{BD}}^0. \quad (\text{A10})$$

Next, the interface energy  $a_k$  with  $k = (\text{AD}, \text{BD})$  is a product of a geometrical factor and  $\Gamma_k$  being the effective surface energy of the vertical edge of the corresponding binary nucleus:

$$a_k = 2 \times 3^{3/4} \Gamma_k \sqrt{\Omega_S h}. \quad (\text{A11})$$

Here  $\Omega_S$  is the volume of a pair in the solid and  $h$  is the 2D nucleus thickness.  $\Gamma_k$  is a function of the surface energies of the solid–liquid ( $\gamma_{\text{SL}}$ ), solid–vapor ( $\gamma_{\text{SV}}$ ), and liquid–vapor ( $\gamma_{\text{LV}}$ ) interfaces and is given by

$$\Gamma_k = \chi \gamma_{\text{SL}} + (1 - \chi) \left( \gamma_{\text{SV}} - \gamma_{\text{LV}} \frac{\Omega_L}{\Omega_S} \sin \beta \right). \quad (\text{A12})$$

Here  $\Omega_L$  is the volume of a pair in the liquid and  $\beta$  is the wetting angle between the catalyst particle and the liquid–solid interface. Parameter  $\chi = 1$  for nucleation in the center and  $\chi = 2/3$  for triple line nucleation. It should be noted that in this paper we don't take into account the dependence of the surface energy of the solid–liquid interface  $\gamma_{\text{SL}}$  on composition of the liquid droplet<sup>39</sup> because it is poorly studied. The parameter values in the cases of nucleation at the triple phase line and in the center of the liquid–solid interface can be found in ref. 56.

## Conflicts of interest

The authors declare no conflicts of interest.

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