Topical Review

Growth of long III-As NWs by hydride vapor phase epitaxy

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Abstract
In this review paper, we focus on the contribution of hydride vapor phase epitaxy (HVPE) to the growth of III-As nanowires (NWs). HVPE is the third epitaxial technique involving gaseous precursors together with molecular beam epitaxy (MBE) and metal-organic VPE (MOVPE) to grow III–V semiconductor compounds. Although a pioneer in the growth of III–V epilayers, HVPE arrived on the scene of NW growth the very last. Yet, HVPE brought different and interesting insights to the topic since HVPE is a very reactive growth system, exhibiting fast growth property, while growth is governed by the temperature-dependent kinetics of surface mechanisms. After a brief review of the specific attributes of HVPE growth, we first feature the innovative polytypism-free crystalline quality of cubic GaAs NWs grown by Au-assisted vapor–liquid–solid (VLS) epitaxy, on exceptional length and for radii down to 6 nm. We then move to the integration of III–V NWs with silicon. Special emphasis is placed on the nucleation issue experienced by both Au-assisted VLS MOVPE and HVPE, and a model demonstrates that the presence of Si atoms in the liquid droplets suppresses nucleation of NWs unless a high Ga concentration is reached in the catalyst droplet. The second known issue is the amphoteric behavior of Si when it is used as doping element for GaAs. On the basis of compared MBE and HVPE experimental data, a model puts forward the role of the As concentration in the liquid Au–Ga–As–Si droplets to yield p-type (low As content) or n-type (high As content) GaAs:Si NWs. We finally describe how self-catalysed VLS growth and condensation growth are implemented by HVPE for the growth of GaAs and InAs NWs on Si.

Keywords: HVPE, III–V, GaAs nanowires, InAs nanowires, epitaxy, NWs

(Some figures may appear in colour only in the online journal)

Abbreviation

nanowire NW;
nanowires NWs;
vapor liquid solid VLS;
vapor solid VS.

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1. General introduction
Semiconductor crystals have been synthesized for more than 60 years by means of epitaxial growth techniques, driven by the societal need for information and communication technologies. The whole story started with the controlled growth of bulk 3D millimeter scale crystals [1, 2] and planar 2D thick buffer layers [3], followed by the race to reduce thicknesses of the 2D epilayers and control their stacking in low
dimensionality heterostructures, including quantum wells and quantum dots, at the heart of microelectronic and optoelectronic devices. Together with these planar geometries, morphology-controlled 3D semiconductor structures were grown at the micrometer and submicrometer scales such as grading, rods, tips, but also whiskers, to end with filamentary 1D nanowires. Whatever the geometry, shape, dimension of the epilayers or structures, the issues at stake are always the same: defect-free growth, control of the chemical composition of the layers, control of the crystal phase, control of the doping, control of the abruptness of interfaces in heterostructures, control of the facet growth anisotropy when it comes to the synthesis of 3D structures.

Three epitaxial processes are being implemented to grow the diversity of these objects: liquid phase epitaxy [4–6], vapor phase epitaxy (VPE) and molecular beam epitaxy (MBE). MBE and metal-organic vapor phase epitaxy (MOVPE) are the most widespread growth techniques for III–V compounds since the 1980s. The prevalence of MOVPE and MBE is still at play when it comes to nanowire (NW) growth. NWs are synthesized by a wide range of methods: top-down photolithography [7, 8], but above all bottom-up catalyst-free selective area growth (SAG) and catalyst-assisted VLS and vapor–solid–solid (VSS) growth. Whatever the route, either VS condensation (in SAG) or liquid–solid solidification (in VLS), the main growth techniques involved in the synthesis of III (Ga, In)–V (P, As) NWs are still MBE and MOVPE. MOVPE is often considered as the most promising for the industrial mass manufacture of semiconductor NW arrays. First micro- and nano-sized wires were synthesized for Si and III–V materials by VLS growth [9, 10]. Continuous efforts have been put forth into catalyst-assisted VLS growth of III–V NWs and most commonly used GaAs NWs for more than fifteen years. VLS growth is particularly well mastered nowadays for both MBE and MOVPE epitaxial tools and enables growth of complex NW structures, including ordered arrays of NWs by catalyzed-SAG on patterned substrates [11–15]. We invite the reader to consult some reviews and books coping with growth, properties and applications of semiconductor NWs [16–37].

Indeed, VPE processes involving chloride (e.g. not metal-organic) gaseous growth precursors were the very first VPE techniques developed in the 1960s and 1970s for the growth of III–V semiconductor layers [2, 38–40]. Yet, chloride-based growth techniques entered the nanowire growth scene the last by far. In hydride VPE (HVPE), the anion elements are transported by gaseous hydrides such as AsH3 for III-As compounds, but the specific attribute of HVPE growth is the use of chloride molecules for supplying the cation elements Ga or In. These volatile chloride GaCl or InCl precursors have a direct impact on the reactor design and the thermodynamic conditions of growth. HVPE is a hot wall technique to ensure the stability of the chloride gaseous species. The growth reactions involving chloride precursors are exothermic while they are endothermic for MOVPE. The growth temperature is about 100 °C higher for HVPE than it is for MOVPE or MBE for the same grown material. HVPE makes use of high partial pressures of precursors, which results in fast growth with some 30–50 μm h⁻¹ for standard growth rate, up to more than 100 μm h⁻¹ with a record at 300 μm h⁻¹ for GaAs layers [41].

These features, high material input and fast growth, certainly justified the reluctance to use HVPE for the growth of small size objects. Moreover, by the time of the first VLS processes implemented in MBE and MOVPE at 400 °C–600 °C for the growth of GaAs NWs involving gold as catalyst, the high temperature needed for HVPE growth of GaAs (700 °C) was considered incompatible with a GaAs solidification from the triple VLS phase. Was HVPE worth the audience for current semiconductor issues, especially growth of NWs, arrays of NWs, nay integration of NW arrays with silicon? If we were to retain only two salient properties of HVPE, of course in addition to the high growth rate property, it would be: (i) perfect selective growth capability, (ii) the ability to reach large crystal growth anisotropy yielding unusual and high aspect ratio structures. The point was to apply these attributes to grow and shape matter at the nanometer scale. As a matter of fact, the road was drawn with the work of Lieber’s group with the first demonstration of millimeter-long silicon nanowires grown at some 30 μm min⁻¹ [42].

In this review paper, we address HVPE growth of III–V NWs. A brief analysis of HVPE thermodynamics is undertaken in section 2, together with a discussion of the growth mechanisms and significant kinetic features of the growth of III–V by HVPE. Section 3 describes the ability of HVPE to grow high crystal quality GaAs NWs with high aspect ratio on GaAs substrates, by VLS-HVPE assisted by gold. Section 4 moves to the integration of GaAs NWs on silicon substrates by Au-catalysed VLS growth. Special emphasis is placed on the nucleation issue and on the doping of GaAs NWs with the silicon element, solving the enigma of n-type or p-type doping encountered with Si depending on the growth technique. Integration of GaAs and InAs NWs on Si by self-catalysed VLS growth is discussed in section 5. We finally summarize the input of HVPE to the synthesis of III–V NWs.

2. Thermodynamics and kinetics of HVPE growth

The growth of (Ga, In)–As layers by HVPE involves gaseous GaCl and InCl molecules synthesized inside the reactor in an upstream zone, by the use of HCl flows reacting with separate Ga and In liquid sources at a temperature greater than 700 °C. As a matter of fact, HVPE is a hot wall reactor technique, needed to produce the III-Cl precursors and insure their stability downstream to the substrate. Element V is transported as arsine (AsH3) which is completely decomposed into (As₂, As₂₂) gaseous molecules by entering the hot wall reactor [43]. HVPE is usually implemented at atmospheric pressure, with H₂ as carrier gas (at 31 min⁻¹). One of the specific attributes of HVPE is that the substrate surface interacts with a complex vapor composed of III-Cl, HCl, H₂ and As₄/As₂, with the last
two species being considered at equilibrium. In HVPE, we discuss results as a function of the III/V ratio calculated as the ratio between the partial pressure of gaseous III-Cl over the partial pressure of gaseous As₄ interacting with the substrate surface in the growth zone. Note that for the temperature range set for the growth of GaAs or InAs, the partial pressure of As₄ is a hundred times greater than the As₂ partial pressure.

III-As deposition can be summarized by the global reaction:

$$\frac{1}{4} \text{As}_4(g) + \text{III} - \text{Cl}_2(g) + \frac{1}{2} \text{H}_2(g) \rightleftharpoons \text{III} - \text{As}_4(c) + \text{HCl}_g,$$

where index g is assigned to gaseous molecules and index c is for crystalline species.

HVPE is known as a near-equilibrium condensation process: one can understand it as reacting almost immediately upon an increase of the supersaturation of the vapor phase that causes the return to equilibrium, which produces the solid [44]. On decreasing the supersaturation, for instance by the use of additional HCl in the vapor phase, reversible processes occur at the VS interface due to the volatility of the chlorides. First consequence is that HVPE features very fast growth (condensation growth rate ranging from 3 to 100 μm h⁻¹) upon setting high material input of growth precursors. As a matter of fact, III-As growth are implemented with partial pressures of As₄ and III-Cl of some $7 \times 10^{-4}$ atm (70 Pa) and $3 \times 10^{-3}$ atm (300 Pa) above the substrate respectively. Please note that HVPE uses III/V ratio greater than one to compensate very substantial desorption of the volatile III-Cl precursors starting around 600 °C. The HVPE reactor design and the experimental parameters can be chosen in such a way that gas-phase mass transport is not a limiting factor for III-As layers growth. Growth is actually governed by the temperature-dependent kinetics of the surface mechanisms: adsorption and decomposition of growth precursors, desorption of by-products, and diffusion of ad-species, as demonstrated by the strong temperature dependence of the GaAs growth rate under standard growth conditions (figure 1) [45, 46]. Among the aforesaid surface kinetics involved in the HVPE process, the III-Cl growth precursor decomposition kinetics appears to be the slowest, so that it is considered as the actual limiting rate for HVPE growth [45]. Note that the decomposition kinetics of III-Cl precursors is still faster than that of other chemical precursors such as metal-organic ones. Record HVPE growth rate (such as 300 μm h⁻¹ for GaAs [41]) is often associated with the use of high supersaturation conditions in the mind of growth physicists. As a matter of fact, HVPE is considered as a high supersaturation process, which is not true, because one may confuse high supersaturation and high material input. Indeed, the Gibbs free energy difference that maintains the stationary state between solid GaAs and a vapor composed of As₄/As₂ and volatile GaCl molecules is some 450 cal mol⁻¹ at 715 °C, which is about eight times lower than the thermodynamic driving force for MOVPE and MBE [47]. It turns out that it is easy to supersaturate the vapor phase to yield high growth rates in HVPE. This highly reactive system needs very few material amount (10 sccm for the gaseous AsH₃ flow, 10 sccm for the gaseous HCl flow over the Ga source or the In source).

Here comes the first question about using fast growth for growing NWs, even by VLS-mediated growth. It is generally assumed that a high growth rate yields lower crystalline quality. Early HVPE experiments demonstrated that high quality epilayers had been grown at growth rates higher than 100 μm h⁻¹ [41]. Use of rapid growth was addressed for the IV compounds with the synthesis of silicon and germanium NWs with lengths up to 30 μm in a pioneer work [42]. Highly uniform millimeter-long Si NWs were successfully grown at a mean rate of 31 μm min⁻¹ and used as building blocks to fabricate field effect transistor arrays consisting of some 100 devices per nanowire [48]. NWs, a few tens of micrometers long, provide larger surfaces to be functionalized, which is of great interest for sensor and detection applications [48, 49]. Due to the rapid decomposition of the III-Cl growth precursors with respect to other chemical precursors, one could expect HVPE to be an efficient alternative tool for the synthesis of III–V alloy NWs with exceptional length at an acceptable growth rate, provided the control of the morphology and crystalline homogeneity of the nano-objects. Very interestingly, Joyce et al used rapid growth rate in VLS-MOVPE to synthesize less tapered, planar crystallographic defect-free III–V NWs [50]. The authors actually observed that increasing the precursor flows and consequently the growth rate, decreased the NW twin density [50]. One should promote untapered morphologies for laser applications where uniform diameters are expected to enhance the NW performance as a resonant cavity. All these results suggest that low defect density semiconductor NWs can be grown very fast.

When it comes to VLS growth, the second issue is related to temperature. HVPE growth of III-As compounds is implemented at some 715 °C for GaAs and 650 °C for InAs, the temperatures at which the thermal activated dechlorination of the III-Cl precursors works well. Investigating VLS in the
HVPE environment addresses the question of how catalyst-assisted VLS could be carried out at such high temperatures. These usual process temperatures in HVPE result in high condensation planar growth rates (standard 30–40 μm h⁻¹ for growth on (001)). Another interest is to assess the competition between VLS solidification of NWs expected to take place in the ⟨111⟩B direction and condensation planar growth on the [001] direction. The first HVPE growth experiments of GaAs NWs described in the next section, answered these two questions.

Last point addresses morphology-controlled 3D micro- and nanostructures. Such morphology-controlled growth is usually carried out by SAG on patterned substrate. HVPE performs very well on that matter as volatile chloride precursors do not adsorb on masked areas whatever the geometry of the pattern, even on dark field patterns [51]. HVPE also gives access to the large growth anisotropy of current III–V crystals. The competition for precursors between masked and open areas being non-existent, one can expect the 3D morphologies to be ruled by surface kinetics, that is by the intrinsic growth anisotropy of crystals. For convex growth, the shaping rule is: a face is all the more developed that it grows slowly. In a growth regime governed by surface kinetics, the growth rate of the faces can be set by varying the experimental parameters of the temperature and composition of the vapor phase, both III/V ratio and supersaturation with the use of additional HCl. HVPE offers the possibility to set a large number of growth rate ratios as a wide range of growth rates are available whatever the orientation [44]. This huge anisotropy allows to shape structures with various morphologies and aspect ratios, in one-step short growth time, capitalizing upon the main advantage of the bottom-up approach: structures are defined by their crystalline properties with no scattering defects as typically induced by techniques involving etching. This is one of the key points of HVPE: as the growth is governed by surface kinetics, one can tailor experimental parameters to increase growth anisotropy, to generate high-aspect-ratio defect-free structures at the micrometer scale such as stripes [52], tips [53], and at the nanometer scale such as gratings of high aspect ratio stripes [51].

3. GaAs NWs grown by Au-catalysed VLS-HVPE

3.1. Brief recall of the issues of early growth experiments of GaAs NWs

As already quoted, the first bottom-up approach for the synthesis of NWs was the catalyst-assisted VLS growth as introduced by Wagner and Ellis in 1964 to explain the epitaxial growth of micron-sized Si whiskers [9]. The second fundamental work was provided by Hiruma et al who demonstrated the feasibility of the VLS growth of III–V nano-whiskers [10]. In VLS growth, a suitable metal catalyst alloys with the metal atoms of the III–V surface introducing a local liquid–solid interface. The constituent materials are supplied from the vapor phase and incorporated into the metal catalyst which acts as a seed for the NW growth. For these pioneer studies, GaAs NWs were mostly grown from Au catalytic particles.

The period between 2000 and 2010 was a very dynamic production time for semiconductor NWs VLS growth. Let us cite major works implementing MBE [54–57], Chemical Beam Epitaxy (CBE) [58], Electron Beam Epitaxy [59], and MOVPE [60–65]. The metallic clusters determine the shape and position of the NWs. Size-selected catalytic Au aerosol particles were manipulated by means of atomic force microscopy, in order to nucleate nano-whiskers in specific positions [66]. Patterning of the catalyst on the substrate surface was also carried out by electron beam lithography [67], or nano-imprint lithography [68], followed by catalyst evaporation and lift-off.

Whatever the structure device is, high material crystal quality is required. The control of the crystal phase of III–V NWs has been a challenging task for a while [69–72]. Indeed, GaAs NWs featured spontaneous zinc blende (ZB)-wurtzite (WZ) polytypism and stacking faults could form between alternating WZ and ZB layers along the ⟨111⟩ axis of the NWs [73–77]. The issue of the crystal phase of NWs is not trivial. It is related to a complex combination of surface energy values, catalyst droplet contact angle, wetting of the NW side facets and applied growth conditions. In pioneering studies of Au-catalysed MOVPE, high V/III ratio and low temperature (T) were demonstrated to favor predominantly ZB NWs, while low V/III and high T were suitable for pure WZ [23, 78]. Crystal phase engineering involving controlled axial stacking of WZ and ZB segments was addressed soon [79, 80]. Lehmann et al refined the conditions and showed that transitions between ZB and WZ were actually related to growth regimes being As-limited (ZB to WZ) or Ga-limited (WZ to ZB) [81]. The role of the droplet contact angle and wetting of NW side facets was clarified later, introducing distinct nucleation locations to yield WZ or ZB segments (see section 3.3).

The first quote of wire growth by HVPE was actually found in 1993 [82]. InP wires were selectively grown on bare (001) GaAs substrates which were patterned with focused ion beam to create steps and kinks for further localised InP nucleation [83–85]. This was made possible because HVPE growth is orientation dependent as was said in the previous section; a diffusion controlled process is inefficient in that case. Ten years have passed before new work on NWs growth by HVPE was published. As a matter of fact, not only very few HVPE research groups are still active for III-As and III-P compounds (Wisconsin-Madison University in the USA; KTH (Sweden), Thales Research and Technology/Thales-Alcatel III–V Laboratory and Institut Pascal (France) in Europe), but also the HVPE high process temperature and fast growth property were considered as major issues for implementing VLS epitaxy of III–V NWs.

3.2. Long untapered rodlike cubic GaAs NWs

The first synthesis of GaAs NWs by Au-assisted VLS growth in the HVPE environment was reported in 2010 [86]. For this preliminary study, non optimum conditions were chosen like
The arrow shows the (111)B growth direction. Wedge angles were determined around 60°. Amorphous native oxide covers the NW. (c) Electron diffraction patterns of a GaAs NW: left [-110], right [-211] zone axis in the cubic ZB structure [86]. (d) HRTEM image of a GaAs NW with a diameter of 22 nm. All HRTEM images were taken along the [1-21] zone axis [87].

First insight of the growth mechanisms involved in Au-assisted VLS-HVPE explains such aspect ratios. VLS-HVPE is supposed to occur preferentially via the adsorption of GaCl molecules on fully liquid Au–Ga–As droplets [87] at growth temperature higher than 700 °C, followed by fast decomposition and desorption of the Cl atoms in the H2 ambient atmosphere. The liquid phase seems plausible (VSS should be excluded) with respect to the process temperature and the observed NW high growth rate: a rich Ga content Au–Ga liquid phase takes place upon the pre-growth annealing of the Au–GaAs substrate, and then the wetting of the catalyst particle with Ga and As insures the NW extension. VLS-HVPE GaAs NWs have uniform diameter from the base to the top, over length greater than 10 μm. A simple argument excludes any diffusion of growth precursors from the substrate surface along the nanowire sidewalls: the length of the wires is much longer than the surface diffusion length of whatever ad-species. If precursors diffused from the substrate surface, then one would have observed a larger base on the NWs. Indeed, the rapid condensation rate on the planar substrate is high enough to incorporate the incoming species. Rapid Ga incorporation into the droplet is favored by the high dechlorination frequency: GaCl molecules are quickly decomposed, probably as soon as they adsorb on the surface of the Au–Ga–As droplet. Such a rapid growth involves condensation when axial growth is based on solidification. The growth rate anisotropy between axial and radial growth is high enough to promote preferential axial development of the NWs and limited radial growth. The post-growth energy dispersive x-ray spectroscopy (EDS) analysis showed a Ga content below 1% in the tip of grown NWs [87]. Such an after-growth low Ga content can be explained by a quick diffusion of the Ga element to the liquid–solid interface, as a consequence of the concentration gradient established between the droplet surface and the heterogeneous liquid–solid interface where a rapid solidification kinetics yields an almost complete and instantaneous consumption of the Ga species. Rapid Ga incorporation into the droplet is favored by the high dechlorination frequency: GaCl molecules are quickly decomposed, probably as soon as they adsorb on the surface of the Au–Ga–As droplet. Such a reactant decomposition kinetics-limited growth regime was also demonstrated for the VLS synthesis of millimeter-long Si NWs [48].

Let us now return to the link between the growth regime and the crystal homogeneity of the NWs. Changes in the
shape of wires were observed when the growth mechanism switched from surface diffusion to direct impingement [57, 88]. Plante et al have shown that modifications of the crystal structure of NWs took place where growth by surface diffusion was dominant [54]. As already discussed, one expect VLS-HVPE to rely on a preferential axial growth mode through the catalyst droplet. The absence of competition between radial and axial growth, the latter one being very limited, almost non existent, favors the crystalline homogeneity of the NWs and probably the absence of twins. Nevertheless, it is not possible to conclude about VLS-HVPE and its capability to control crystal homogeneity unless growth of thin NWs of diameter less than 20 nm has been addressed, knowing that structure variations, from cubic to wurtzite, were observed for very thin GaAs NWs produced by MOVPE and MBE.

3.3. The polytypism issue in GaAs NWs with diameter less than 10 nm

GaAs NWs grown by MOVPE or MBE with diameter lower than 30 nm were shown to exhibit a wurtzite structure with some ZB-WZ polytypism. The thinnest GaAs NWs published in 2009 were grown by VLS-MBE: NWs with diameter of 10 and 13 nm exhibiting a WZ structure over length of 5 μm with negligible intermixing of ZB stacking [89]. Thorough theoretical investigations of catalyst-assisted growth in the MBE and MOVPE processes have shown that suppression of nucleation at the triple phase line (TPL) (where the vapor, liquid and solid phases meet) helps to avoid ZB-WZ polytypism under optimized growth conditions [75]. This model was supported by experimental data provided by the VLS-MBE process with the Ga-catalysed growth of GaAs NWs [90]. As a matter of fact, several groups confirmed that self-catalysed III–V NWs adopted the ZB phase much more often than Au-catalysed one [90–92]. Zamani et al implemented polarity engineering to grow stacking disorder free (111)A-polar GaAs NWs under Ga-rich conditions in Ga-assisted MBE [93].

It appeared interesting to follow the evolution of the crystalline phase of NWs grown in the HVPE environment for the peculiar radius range (5–15 nm) where uncontrolled polytypism and conversion to WZ in (111)B NWs took place. GaAs VLS-HVPE growth was carried out on Au-coated (001) GaAs substrates [94]. Au droplets with diameters ranging from 50 to 10 nm were formed after annealing at 715 °C in the HVPE reactor of Au/GaAs substrates where a nominal amount of Au equivalent to 0.1 monolayer was deposited. GaAs growth was carried out at 715 °C for 30 min with a partial pressure of GaCl of 300 Pa, a partial pressure of As2 of 71 Pa (partial pressure of As2 was 0.5 Pa) and with a H2 vector flow of 3000 sccm including additional HCl flow (at 50 Pa) used to tune the supersaturation of the vapor phase [45, 46, 52]. These experimental parameters yielded a GaAs 2D planar growth rate of 38 μm h⁻¹ and 20 μm h⁻¹ on (001) and (111)B GaAs substrates, respectively.

Figure 3 shows the transmission electron microscopy (TEM) and HRTEM images of a GaAs NW 14 μm long with a constant radius of 15 nm grown by VLS-HVPE. Figure 3(c) displays the HRTEM image of a NW with a radius of 5–6 nm, indexed in the cubic ZB structure (see the fast Fourier transform (FFT) image in figure 3(c)). NWs were grown in the (111)B direction, exhibiting stacking-fault free pure ZB phase over the lengths which could be observed on individual NWs, that is up to 20 μm. Contrary to the NWs with radii of 25 nm or above, which were found hexagonal with (211) facets [86], thinner NWs have a more complex faceting. The prevalence of (211) ZB facets makes the observed crystal phase purity even more surprising, since it is known from different calculations [95–97] that these facets have a higher surface energy than (110) and therefore the formation of ZB structure is more energetically costly in this case. The defect-free area includes a very narrow zone just beneath the droplet where structural instabilities take place due to unsteady conditions in the droplet upon stopping precursors at the end of growth, resulting in progressive decreasing of the Ga liquid supersaturation after consumption upon crystallization.

The unexpected ZB phase for GaAs NWs of 5–15 nm radii calls for discussion. It is predicted from equilibrium formation energies that pure WZ phase should form for a radius of the order of 10 nm regardless of the growth conditions [97, 98]. In addition to these considerations related to intrinsic energetic morphological stability, crystal growth conditions play a major role through supersaturation state in the catalyst droplet which drives the nucleation path. The competition between the ZB and WZ structures is driven by the difference in the corresponding nucleation barriers rather than equilibrium energies of fully formed NWs. This increases the actual critical radius for the ZB-to-WZ transition to the value of the order of several tens of nanometers depending on the growth conditions [95]. VLS-HVPE yields NW axial growth rate of 100 μm h⁻¹ or higher, from liquid Au–Ga–As droplets with an original high percentage of Ga (higher than 30% if the droplets were pure binary Au–Ga alloys) [99].

In order to predict the crystal phase of NWs grown by Au-assisted VLS-HVPE, a model was developed in two steps [94, 100]. The first step expresses the liquid chemical
potential of GaAs in an Au–Ga–As droplet with respect to the stoichiometric ZB solid:
\[
\Delta \mu = \mu_3^1 + \mu_5^1 - \mu_{35}^S
\]
with \(\mu_3^1\) and \(\mu_5^1\) as the chemical potentials of Ga and As atoms in the liquid state respectively and \(\mu_{35}^S\) as the chemical potential of a GaAs pair in the solid state. \(\mu_3^1\) is written as a function of the Ga concentration in the liquid droplet \(c_3\) and is temperature-dependent. \(\mu_5^1\) and \(\mu_{35}^S\) do not contain dependence on the droplet composition. \(\mu_3^1\) is written as a function of the ratio between the As desorption flux and the As adsorption flux on the droplet surface. The prevalence of the WZ phase over ZB in VLS NWs requires that the \(\Delta \mu\) value is larger than two critical chemical potentials. The first one corresponds to the equality of the nucleation barriers for the critical WZ and ZB nuclei at the TPL and the second to the equality of the nucleation barrier for the WZ nucleus at the TPL and the ZB nucleus in the center (C) of the liquid–solid interface under the droplet. Consequently, the WZ structure forms when \(\Delta \mu > \max(\Delta \mu_{TPL}; \Delta \mu_C)\) with \(\Delta \mu_{TPL}\) and \(\Delta \mu_C\) written as a function of the bulk energy difference between the WZ and ZB structures, involving the surface energies of relevant ZB and WZ NW sidewall facets, the liquid–vapor surface energy of the droplet, the surface energy of the lateral solid–liquid interface of a two-dimensional island, and the fraction of the island perimeter at the TPL. The critical curve \(\max(\Delta \mu_{TPL}; \Delta \mu_C)\) is shown in figure 4. It is seen that the critical chemical potential is limited by the \(\Delta \mu_{TPL}\) for the Ga concentrations lower than 0.36–0.51, depending on the ZB facet type, while for higher \(c_3\), it is given by the \(\Delta \mu_C\), i.e., by the competition of the WZ TPL and the ZB C nucleation scenarios. Above \(c_3=0.65\), the WZ phase formation is completely suppressed on surface energetic grounds due to a low surface energy of a Ga-rich droplet, as in the case of pure Ga catalyst [90]. Hence, the WZ phase should be observed for high chemical potentials and low Ga concentrations in the top-left corner, while the ZB phase dominates below the critical curves. Very importantly, the ZB phase should form for all \(c_3 > 0.65\) actually regardless of the NW radius. This feature does not depend on the ZB facet type (i.e. (211) or (110)), because the nuclei are entirely surrounded by the liquid phase and do not feel the sidewalls. Therefore, the absolutely defect-free ZB phase of HVPE-grown GaAs NWs would be confirmed if we show that the Ga concentration in the droplets during growth is above 0.65 for all NWs.

Second step of the modelling writes the NW elongation rate \((dL/dt)\) as a function of the kinetics of As atoms arriving to and leaving from an Au–Ga–As droplet. The actual Ga concentration during growth can be estimated by equating the observed elongation rate (with the minimum of 100 \(\mu m h^{-1}\), or 28 nm s\(^{-1}\)) to the nucleation-mediated NW growth rate:

\[
\frac{dL}{dt} = \frac{3\sqrt{3}}{2} R^2 h \times A c_5 \frac{\Delta \mu}{k_B T} \exp \left( -\frac{\Delta \mu}{k_B T} \right)
\]

with \(R\) the radius of a two-dimensional nucleus of a monolayer height \(h\), \(A\) an unknown pre-factor in the Zeldovich nucleation rate, \(c_5\) the As concentration in the droplet. \(A G\) is the nucleation barrier for triangle two-dimensional nuclei of dimensions \(R\) and \(h\). All calculation details are given in [100].

The nucleation rates obtained at 715 °C and \(A = 7 nm^2 s^{-1}\) (as recommended by Glas et al at temperatures near 600 °C [101]) are shown by the red lines in figure 5 for different NW radii from 5 to 50 nm. It is seen that crossing points of these...
curves with the minimum experimental elongation rate of 28 nm s\(^{-1}\) correspond to the values of \(c_3\) which are systematically higher than 0.7. If we increase the value of \(A\) by a huge factor of \(10^4\) (which could partly be due to enhanced diffusivity of As at 715 °C compared to 600 °C), the Ga concentration would still remain higher than 0.65, as shown by the blue lines in figure 5. In other words, the exceptional elongation rate in VLS-HVPE necessarily requires the droplet to be Ga-rich, which yields the pure ZB phase due to the suppression of the TPL nucleation on surface energetic grounds. This conclusion is robust with respect to variations in temperature (700 °C–740 °C) and in the generally unknown pre-exponential factor \(A\).

As a summary, record pure cubic phase 20 nm long and 5 nm in radius GaAs NWs were grown by Au-assisted VLS-HVPE [94]. It was demonstrated that VLS growth took place at high liquid supersaturation, yielding a Ga concentration higher than 0.5 in the Au–Ga–As droplets. The high Ga concentration decreases the surface energy of the droplets, which inhibits nucleation at the TPL, thus preventing the formation of WZ structure regardless of the NW radius. These results confirmed the theoretical predictions of Glas, Dubrovskii et al. by providing experimental data with Ga-rich concentrations not readily accessible by the MBE and MOVPE processes. Recently, NW growth was followed in situ in transmission electron microscopes. Emphasis was placed on the values of the contact angle of the catalyst droplet that favor nucleation at the TPL (90°) yielding to the WZ phase, or favor the formation of a truncated facet at the liquid–solid interface which yields nucleation at a distance from the TPL (greater than 90°) and results in the ZB phase [102–104].

One of the major strategic challenge of the 21st century still remains the integration of III–V devices with silicon, driven by the demand for data treatment and transmission. Transitioning from electrical data transmission to optical data communication is one of the key points that focus attention. Obviously, the NW geometry allows to evacuate the elastic stress due to the large lattice mismatch between the III–V materials and Si (ranging from 4% to 11%) through the free lateral surfaces of the NWs. Nevertheless, nucleation and further growth of III–V materials on Si is not that easy. HVPE brings answers to issues encountered in MOVPE (nucleation issue) and in MBE (doping issue).

4. Integration of GaAs NWs on silicon substrate

4.1. Introduction

For Au-assisted VLS growth of GaAs NWs on GaAs substrates, MBE [105–107] and MOVPE [108, 109] operate at temperatures ranging from 400°C to 620°C. HVPE GaAs NWs are grown at temperatures higher than 700 °C. In this wide range of temperatures, the atomic interdiffusion between the substrate and the catalyst always serves as an additional supply of gallium and should be favorable for the NW nucleation on GaAs substrates [87, 105].

The picture becomes different when the VLS growth of GaAs NWs takes place on silicon substrates covered or not with an oxide layer. Yet, integrating III–V compounds or NWs with silicon is of prime interest, for it opens the way to the development of photonic integrated circuits on Si for data treatment. Implementing Au-assisted VLS growth on Si in the HVPE environment brings new insights to nucleation and Si doping issues of GaAs NWs.

4.2. Au-catalysed VLS-HVPE of GaAs NWs on Si: the nucleation issue

Au-assisted VLS growth of GaAs NWs requires the deposition of an ultra-thin gold film on Si. The further density of the NWs on unpatterned substrate depends on the dewetting of the Au film on the silicon substrate surface. The control of this step is crucial to adjust the density and the diameter of the NWs during VLS growth. Several papers have reported the dewetting of Au on GaAs substrate [110]. As for Si substrate, whether covered with its native oxide layer or covered with a controlled dioxide film, the kinetics of Au droplets formation is strongly modified. We have first studied the dependence of the diameter and spatial distribution of the droplets on the surface of Si (111) substrates with respect to the thickness of the Au film and the temperature of the substrate during deposition and post-deposition annealing step.

Au films of thickness ranging from 0.2 to 2 nm were deposited at RT in an UHV system, on Si (111) substrates covered with their native oxide. The films were subsequently annealed at 600 °C for 10 min in UHV. Figure 6 displays the scanning electron microscopy (SEM) images of the surfaces for four thicknesses of gold. The first point that stands out is the increase of the droplets size and the decrease of the droplets density with increasing the Au film thickness.

The following main behaviours have been established supported by in situ low energy electron microscopy coupled to low energy electron diffraction technique [111]. When Au is deposited at RT, there is no effect of the oxide thickness on the size distribution and the density of the droplets. Under the annealing step in the range of 600 °C–700 °C, and for thin SiO\(_2\) layers, the decomposition of the oxide is activated by the Au atoms. This process modifies the Au droplets growth. We have shown that large Au droplets decompose the SiO\(_2\) layer and unveil the Si (111) substrate while migrating perpendicular to the step edges. Small droplets are pinned and decay.

| \(h_x = 0.2\) nm | \(h_x = 0.5\) nm | \(h_x = 1\) nm | \(h_x = 2\) nm |
|-----------------|-----------------|-----------------|-----------------|
| (a1)            | (b1)            | (c1)            | (d1)            |
| Native Si\(_2\)O | Native Si\(_2\)O | Native Si\(_2\)O | Native Si\(_2\)O |

Figure 6. Distribution of Au droplets after deposition of (a1)–(a2) 0.2, (b1)–(b2) 0.5, (c1)–(c2) 1 and (d1)–(d2) 2 nm of Au on native Si\(_2\)O at RT followed by annealing at 600 °C for 10 min under UHV. The white and yellow scale bars represent 200 nm and 50 nm, respectively [111].
spontaneously. This study brings important information to NW growers using VPE methods. It allows to control the density and the diameters of the Au droplets for subsequent NW growth.

Contrary to MBE, in VPE methods, it is almost difficult to start growth of III–V materials on bare silicon surface. It is worth mentioning that the native SiO$_2$ is always removed prior to the growth in MBE chambers. In the HVPE process, the SiO$_2$ layer is always present at the surface of the substrate since HVPE operates at atmospheric pressure.

It has been shown that the presence of an oxide layer at the interface between gold and the silicon substrate blocks the inter-exchange between the two materials [112]. Therefore, the interdiffusion of Si atoms into the gold alloy droplets occurs only after decomposition of SiO$_2$ underneath the droplets, which starts for temperatures above 400 °C. The Si$_{6+}$ + SiO$_{26+}$ → 2 SiO$_{4+}$ reaction is enhanced in the presence of gold atoms that reach the SiO$_2$/Si interface by thermally activated diffusion into SiO$_2$. Obviously, this diffusion is easier through thinner SiO$_2$ layers. A similar enhancement is anticipated when the VLS growth on silicon is assisted by gallium droplets [101, 113–115]. Additionally, the SiO$_2$ layer can be decomposed at high enough temperatures via the reaction with gallium. A fraction of the gallium droplet is evaporated as Ga$_2$O and the remaining part takes silicon atoms away from the substrate, leading to enhanced silicon evaporation [116].

Therefore, in both gold-catalysed and self-assisted VLS growth of GaAs NWs on silicon substrates, the diffusion or re-adsorption of silicon atoms into the droplet prior to nucleation of GaAs NWs is unavoidable. The strength of this effect may vary depending on the deposition conditions and is anticipated to be larger for thinner oxide layers. Also, silicon should diffuse into the catalyst droplets more easily at higher temperatures and in chemical epitaxy techniques where different precursors are able to etch out silicon atoms from the substrate.

In order to quantify how the presence of Si atoms in the catalyst droplets affects the nucleation rate of GaAs NWs, V. G. Dubrovskii et al developed a model, based on HVPE data [117]. This approach is general as it treats chemical potentials in a quaternary Au–III–V–Si liquid alloy over a wide range of temperatures, and it is irrespective of a particular epitaxy technique employed. Therefore, any VLS growth of III–V NWs on silicon with different catalysts (gold, gallium, and indium) may be treated using the proposed scheme. Silicon is widely used as dopant for GaAs NWs. Here, we consider the influence of the silicon concentration in the droplet on the nucleation probability of pure GaAs NWs on different SiO$_x$/Si substrates, that is, not allowing for silicon to enter the solid state. The growth of Si-doped GaAs NWs and ultimately a ternary Si-GaAs solid alloy is presented in section 4.3.

HVPE growth of GaAs NWs were performed after deposition of gold on native SiO$_2$/Si (111) substrates (figure 7(a)) and on Si (111) substrates without native SiO$_2$ (figure 7(b)). NWs grew easily when the native oxide was kept prior to the growth, while only few parasitic islands grew on the bare Si (111) after HF etching of the oxide layer. Figure 7. Scanning electron microscopy (SEM) images of the GaAs structures grown at 715 °C after depositing 1 nm of gold on Si(111) (a) native SiO$_2$ and (b) without SiO$_2$. TEM images of the gold-catalysed GaAs structures with droplets on top, grown at 715 °C [117].

Matteini et al reported the influence of surface silicon oxides on the density of self-catalyzed grown GaAs NWs [118]. In agreement with this work, we demonstrated that in VLS growth with Au–Ga–As droplets, NW growth was enhanced as long as an oxide film is present on the substrate, which decreases the interdiffusion of Si atoms into the gold-alloy droplets [117]. Post-growth EDS quantitative analysis revealed that the silicon concentrations in the droplets were less than 2% on SiO$_2$/Si (111) substrates and around 30% on bare Si (111) substrates. On the post-growth cooling stage under arsenic flux, Ga atoms were consumed, leading to droplet shrinking and the corresponding diminishing of the NW top radius. Assuming that the number of Au and Si atoms present in the droplet during growth remained the same after growth, we assessed the concentrations of Si and Ga atoms from geometry, by comparing the initial volume of the droplet with Ga and its final volume without Ga, with the measured initial and final radii of the NW top [117]. In terms of Ga concentration, the calculations gave values close to 0.95.

A model based on the idea that adding silicon to the Au–Ga–As droplets had a drastic effect on chemical potentials, thus on the nucleation probability of GaAs NWs was developed. The model calls on a parameter $f$ that is written as the chemical potential difference per GaAs pair in the catalyst Au–Ga–As droplets with and without Si [117]. The variations of $f$ as a function of the Si concentration ($c_{Si}$) are plotted in figure 8 for various concentrations of Ga ($c_{Ga}$) in the catalyst droplet. The curves are read like this: if $f$ is negative, that is observed for Ga-rich droplets, then the chemical potential of the droplets containing Si is lower than the chemical potential in the droplets free of Si. It means that adding silicon to Ga-rich gold-gallium droplets makes the initial nucleation of GaAs NWs more difficult. The condition $c_{Si} < 0.3$ at $c_{Ga} > 0.65$ guarantees that the presence of silicon suppresses nucleation of GaAs NWs on Si(111). The limiting case of that is pure gallium-assisted growth ($c_{Ga}$ close to 1), which is the most sensitive to the presence of Si.

As a conclusion, Si suppresses nucleation from Ga-rich droplets as long as a certain critical concentration is not reached, with the most difficult nucleation for gold-free self-assisted VLS growth. As more Si atoms enter the droplets in MOVPE and HVPE due to enhanced silicon etching by the chemical precursors, we now explain the known issue of Ga-catalysed growth of GaAs NWs by VPE techniques.
Much less etching should be present during MBE, so the silicon concentration is lower in the catalyst droplets and Ga-catalysed GaAs NWs can easily be grown on Si by MBE [113, 114]. The model confirms that having more gold in the droplets helps to nucleate GaAs NWs on silicon. This property has been long known for MOVPE (see, for example [119, 120]).

### 4.3. Au-catalysed VLS growth of GaAs:Si NWs on Si: the doping issue

GaAs:Si NWS were grown on Si (111) substrates covered with a native SiO2 layer after deposition and annealing under UHV of a gold layer 1 nm thick, using SiH4 gas for doping. Various growth temperatures were tested [121].

From figure 9, it comes out that the substrates contain high density of nanostructures. They exhibit large tapering at the base, except the ones grown at 580 °C (panel a) which have larger uniform diameters. The samples grown at 715 °C present the thinnest structures. HRTEM and FFT revealed pure ZB phase of representative NWs [121]. Micro-photoluminescence was performed on GaAs NWs with different Si doping concentrations. The blueshift of the spectra revealed the n-type behavior of GaAs NWs grown by HVPE.

Si is known for amphoteric behavior [122–125]. Replacing Ga with Si atoms creates As–Si pairs and results in n-type doping, while replacing As with Si atoms creates Ga–Si pairs and results in p-type doping. The replaced Ga or As atoms are dissolved in the liquid droplet. This property has previously been used to change the doping of GaAs layers from n-type to p-type in MBE by tuning the growth conditions. In the case of MBE, p-doping is achieved by lowering the As/Ga ratio and increasing the growth temperature [126]. V.G. Dubrovskii developed a model to understand why Si can act as a n-type or p-type dopant in GaAs NWs grown by VLS, while it remains a n-type dopant of GaAs grown by VS condensation [121].

Considering the VLS growth of GaAs:Si NWs from a quaternary Au–Ga–As–Si liquid alloy, we study the Si incorporation from liquid to solid under equilibrium conditions for all species containing Si [127] and is expected to be even more relevant for doping due to extremely low Si concentrations in both liquid and solid phases.
In conclusion, it has been shown that Si atoms are preferentially incorporated into As, rather than Ga sites of GaAs NWs, due to a much lower As/Ga ratio in the mother phase during the VLS growth compared to VS growth. This explains why most VLS GaAs:Si NWs are p-type, while the standard VS growth leads to n-type doping. However, n-type doping of GaAs NWs with Si is expected at higher As concentrations in the droplet, which occurs for HVPE growth on Si substrates. These results can be used for obtaining n-type or p-type Si doping in GaAs NWs by tuning the growth conditions, or even Si-doped p–n junctions within a single NW.

5. Self-catalysed growth of GaAs and InAs NWs on silicon substrate

5.1. Thermodynamic conditions to form Ga and In liquid droplets from chloride gaseous molecules

In order to grow self-catalysed III-As NWs, a thermodynamic analysis of the conditions that yield the deposition of liquid droplets of Ga or In in HVPE has to be investigated. Gallium or indium droplets should nucleate from III-Cl gaseous molecules. One should consider the chemical reactions involving any III-Cl$_{mg}$ species likely to be present in the vapor phase to form liquid droplets of element III on the substrate surface:

$$\text{IIICl}_{mg} + \frac{m}{2} \text{H}_{2g} \rightleftharpoons \text{III} + m \text{HCl}_{g}. $$

For instance, GaCl$_{3g}$ and GaCl$_{5g}$ molecules are found in the vapor phase at a temperature of 600 °C, and all the more for higher temperature, provided the Cl/H atomic ratio is low, which is always the case for HVPE carried out at atmospheric pressure under H$_2$ vector gas [128].

The equilibrium constants $K$ for the reactions producing liquid Ga or liquid In and HCl$_g$, from GaCl$_e$ or InCl$_e$ and H$_2g$ reactants were calculated using thermochemical data of [129]. They are plotted in figure 12 as a function of temperature $T$. Both equilibrium constants $K_{Ga}$ and $K_{In}$ rapidly decrease with increasing temperature and are much smaller than 1 for the
usual process temperature in HVPE \((T > 600 \, ^\circ C)\), indicating that the material balance at equilibrium is not in favor of liquid gallium or indium when trying to form droplets from III-Cl gaseous molecules. As a matter of fact, any attempt to form droplets on a Si surface from III-Cl gaseous species is doomed to failure. As a consequence, the stabilization of droplets of Ga or In requires the presence of both III and As species. Gaseous III-Cl and As precursors are to be supplied simultaneously at 600 \(^\circ C\). In order to prevent the two-dimensional crystallization of GaAs or InAs that would lead to the fast consumption of the droplets, the ratio of the III-Cl to the As input partial pressures should be tremendously high, to drive the supersaturation toward nucleation of liquid Ga or In on the substrate surface. As a conclusion, unusual III/V atomic flow ratios are necessary to keep the Ga or In-droplets that will subsequently catalyze the VLS nucleation and growth of III-\(V\) NWs in a HVPE reactor.

5.2. Self-catalysed growth of GaAs NWs on Si

Gold-free growth techniques such as VLS catalysed by the group III metal rather than gold, are developed to save one step before growth and facilitate the integration of III–V devices with the silicon platform. Ga-catalysed VLS growth of GaAs NWs by MBE was introduced by Colombo et al [130]. Since then, remarkable progress has been achieved towards MBE synthesis, theoretical understanding and tailoring of physical properties, and device implementation of such NWs [118, 131–133]. In particular, one can take advantage of well-developed techniques of SiO\(_2\)/Si preparation and processing that influence the NW morphology to obtain high crystal phase purity (ZB) and modulate the NW diameter by tuning the As/Ga flux ratio [90, 132, 134].

Furthermore, gallium droplets can self-equilibrate to a certain stationary size within a range of As/Ga flux ratios and this effect leads to a narrowing of the radius distribution of NWs [101, 114, 135]. The elongation rate of Ga-catalysed GaAs NWs is determined by the arsenic vapor flux and is radius-independent for sufficiently large droplets [101, 130], which improves the length uniformity within the NW ensembles [31]. All these features allow for MBE growth of GaAs NWs on silicon with the highest crystal quality, spatial uniformity and tunable morphology.

Ga-catalysed VPE growth of GaAs NWs, namely by the MOVPE technique, remains a very challenging task and does not produce structures with morphological and crystal phase perfections comparable to those obtained by MBE. The main problem is due to the high temperature (at least 600 \(^\circ C\)) required for cracking organo-gallium precursors combined with arsenic-rich atmosphere in the reactor. Such conditions do not favor both initial nucleation and further stability of liquid gallium droplets [136]. Given these limitations, it is extremely difficult to find appropriate conditions for Ga-catalysed VLS growth, and the NWs obtained so far are limited in length (typically less than 1 \(\mu m\)).

First Ga-catalysed VLS growth of GaAs NWs by HVPE was carried out on partially etched native SiO\(_2\)/Si (111) substrate, at 600 \(^\circ C\). Hexagon bottle-like GaAs NWs, 2 to 10 \(\mu m\)-long, were grown in a 10 min-step. The density of NWs was very low (see figure 13(g)) [137]. This low density is related to the lack of liquid Ga droplets on the surface (see section 5.1). Actually, the oxide thickness has a lower influence in HVPE than in MBE. In order to increase the NWs density, one should deposit \(ex \, situ\) Ga droplets and further keep the substrate under controlled atmosphere to prevent the oxidation of the droplets.

HRTEM images of typical GaAs NWs mechanically separated from the substrate and deposited on a holey carbon grid are shown in figure 13. The NW shown in figures 13(a)–(f) grown at 600 \(^\circ C\) presents a very thick \(300 \, nm\) at the base) tapered trunk with a pure ZB crystal structure (figure 13(b)) and no evidence of twins or stacking disorder as often observed in MBE-grown self-catalysed GaAs NWs [90]. A similar NW observed perpendicular to the growth axis reveals a base of hexagonal shape (figure 13(f)). Elemental maps of Ga and As obtained from EDS analysis show that the NW has a homogeneous composition of gallium and arsenic (figures 13(d) and (e)), and in particular no Ga droplets or Ga-rich regions are found at the NW tip. As a matter of fact, all the excess gallium should be completely consumed in the cool-down stage under As\(_2\)/As\(_4\) atmosphere. So we assume that the NWs were grown by Ga-catalyzed VLS growth, according to four steps described in the discussion below.

All NWs are oriented along the (111)\(B\) direction. NWs have different lengths on a single sample. The average integrated growth rate (i.e. the mean NW length divided by the total growth time of 10 min) can be estimated to 30 \(\mu m\, h^{-1}\). Figure 14(a) displays the variation of the mean NW length as a function of the arsenic partial pressure at a fixed GaCl

![Figure 13. GaAs NW grown at 600 °C with III/V = 55. (a) Contact part of GaAs NWs and Si(111). (b) Corresponding HRTEM image of the bottom part, the FFT above reveals the defect free cubic ZB structure of the NW. (c) TEM image of a GaAs NW with a thick trunk (base dimension = 300 nm). (d), (e) Corresponding gallium and arsenic elemental maps from EDS. (f) TEM image of a NW view along the growth axis evidencing the hexagonal shape of the base. (g) SEM image of one typical GaAs NW, showing the particular bottle shape [137].](image)
partial pressure: the NW length decreases with increasing the As partial pressure until complete supression of the NW growth. This behavior is unusual compared with MBE, where the axial NW growth rate increases with the arsenic flux \[90, 132, 138\]. Figure 14(b) shows the non-monotonic dependence of the mean NW length with increasing the GaCl partial pressure \[128\]. In the case of MBE, the axial NW growth rate is not influenced by the gallium flux that only changes the droplet size \[90, 138\]. Indeed, Ga-catalysed VLS-HYPE involves four steps. To explain the low density of NWs, one should work on the very first step which is the nucleation step. V.G. Dubrovskii demonstrated that initial nucleation requires that the surface coverage in Ga atoms reaches locally a critical value, upon vapor flux and surface diffusion after adsorption on the substrate \[137\]. We showed in section 5.1 that it was particularly difficult to get liquid Ga adsorbed on the substrate surface from gaseous GaCl molecules. As a result, very few NWs can be grown. Second step is the inflation of the droplet which is rapid once the seed was nucleated. The third step is the shrinking of the droplet because of a high input flux of As, followed by a stationary regime (step 4) once the balance between the Ga flux and the As flux is reached. The four steps were modelled as a function of the adsorption flux of Ga, the net adsorption flux of As as we consider the possible desorption of the As element, and geometrical parameters related to the droplet (radius, height, contact angle) \[137\].

These results, namely very low density and bottle-like shape NWs, clearly demonstrate that HVPE is not suitable for self-catalyzed growth of GaAs NWs on non patterned substrates. One should look to direct condensation of the III–V on Si, as initiated with InAs.

5.3. Growth of InAs NWs on Si

InAs has a direct and narrow band gap, high electron mobility and small electron effective mass. The integration of InAs with Si is of great interest for ultra-high speed and low-power components \[139\]. The planar heteroepitaxy of high quality InAs materials on Si is even more difficult than it is for GaAs due to the larger difference in the lattice constants between InAs and Si (11.4%). Lapierre et al. have demonstrated the high potential of MBE to grow InAs NWs with a special focus on infrared detectors applications \[140–142\]. Yet, while indium-catalysed VLS growth of InAs NWs have been successfully demonstrated by both MBE and MOVPE, to our knowledge no demonstration of InAs NWs growth was given by HVPE.

In self-catalyzed MBE, the indium catalyst is obtained either by \textit{in situ} deposition prior to growth \[143\] or by initiating growth under indium rich conditions \[144\]. The NW morphology is very sensitive to the growth parameters such as the temperature and the vapor phase composition. For instance, if highly arsenic rich conditions are used, growth leads to InAs NWs with a highly defective wurtzite structure \[145\]. Nevertheless, InAs-based NW heterostructures have been successfully demonstrated by MBE \[144, 146\].

Dick et al. investigated self-catalyzed MOVPE growth in the range of [400–500]°C on InAs and InP substrates \[147\]. The same group was pioneer in the growth of InAs NWs on silicon substrates \[148\]. Recent MOVPE work involved Si wafers precoated by metallic masks, to avoid the use of gold catalyst and reduce process costs \[149\]. Light emitting InAs NWs were grown directly on flexible plastic substrates by In-seeded MOVPE \[150\].

In MBE and CBE, several groups showed that VS growth of InAs NWs was mainly limited by the surface diffusion of adatoms, which does not favor a pure crystalline quality of NWs that exhibit a high density of twin planes, stacking faults and alternating ZB/WZ segments along the growth axis \[143, 144, 151\]. MOVPE provided also very promising results in the case of SAG, even though the wires did not exhibit a pure crystalline quality all along the NWs length limited to few microns \[152, 153\].

As a conclusion, when InAs NWs are grown by self-assisted VLS or VS epitaxy, their structural quality is clearly inferior to that of NWs grown by Au-assisted VLS epitaxy. It is highly relevant to develop a better understanding of the emergence of polytypism in NWs and to grow defect free NWs. The question is what the contribution of HVPE with its specific thermodynamic and kinetic properties would be in this context.

As described in section 5.1, the material balance at equilibrium is not in favor of the formation of liquid indium droplet in the growth temperature range (580°C–720°C) for InAs. Thus HVPE growth of InAs should proceed by condensation. Figure 15(a) displays the SEM image of InAs NWs...
grown on Si (111) at 640 °C. In this work, no surface treatment has been done prior to growth, except HF etching of silicon substrates prior to their introduction into the reactor. Consequently, the NWs were grown on silicon surfaces presenting random re-oxidation and random holes in the native silicon oxide. The NWs were grown predominantly along the vertical (111)B direction whatever the growth temperature. Very few tilted NWs were observed. We attribute the existence of the two populations to the non-polar nature of the Si (111) substrate. It is clearly seen that the NWs exhibit a hexagonal shape delimited by six equivalent {111} side facets and the (111)B facet at the top.

Alternate stacking of ZB segments with inclusions of WZ and twin planes are visible in figure 15(b). These observations have been confirmed whatever the HVPE growth conditions of temperature and III/V ratio. In MBE, it is assumed that the appearance of such structural defects is linked to the surface reconstruction of the (111)B face [153], during InAs growth implemented under As-rich conditions. With partial pressures of As$_4$ ranging from $3.5 \times 10^{-4}$ atm (35 Pa) to $1 \times 10^{-3}$ atm (101 Pa), it is obvious that HVPE falls into such a regime.

The average axial growth rate of the InAs NWs is plotted as a function of the growth temperature $T$ in figure 16. It ranges from some $\mu$m h$^{-1}$ to more than 50 $\mu$m h$^{-1}$ and exhibits a bell curve as a function of $T$. This strong dependence with the temperature demonstrates that growth is governed by the surface kinetics and the thermally-activated steps, as described in section 2. Alike planar growth, NW axial growth is limited by inhibited dechlorination of AsInCl ad-species at low $T$ and normal lack of material adsorption at high $T$. Having demonstrated the potential of HVPE for the easy nucleation and condensation growth of InAs NWs on unprepared and unpatterned Si substrate, efforts are now made to lower the As partial pressure to get pure phase InAs NWs. SAG on patterned Si substrates is also ongoing to fabricate regular and periodic arrays of InAs NWs of which length can be tuned at will by HVPE in short process time.

6. Summary

GaAs NWs grown by Au catalyst-assisted VLS-HVPE on GaAs (001) feature an untapered morphology with a constant diameter. They show very high aspect ratio, ranging from 800 to 1000 for radii ranging from 50 to 130 nm. They are grown at a high axial growth rate greater than 100 $\mu$m h$^{-1}$. GaAs NWs exhibit defect-free and stacking fault-free zinc blende structure. The occurrence of polytypism-free ZB phase in Au-catalysed GaAs NWs of record radii lower than 15 nm down to 5 nm was demonstrated for the first time. Theoretical modeling was proposed to elucidate the unexpected ZB structure and its purity for such small dimensions.

Nanowire growth by VLS-HVPE only proceeds by introduction of precursors in the catalyst droplets from the vapor phase. This promotes almost pure axial growth leading to NWs with a constant cylinder shape over unusual length. We demonstrate that VLS-HVPE takes place at high liquid chemical potential that originates from very high influxes of both As and Ga. The high Ga concentration in the Au–Ga–As droplets decreases the surface energy of the droplets, which disables nucleation at the TPL thus preventing the formation of WZ structure regardless of the NW radius. This conclusion supported theoretical and experimental data obtained by MBE, where it was demonstrated that high Ga concentration was needed to yield ZB nucleation for Au-assisted VLS-MBE, and that one should switch to Ga-catalyst to guarantee pure ZB GaAs NWs.

The implementation of Au-assisted VLS-HVPE on Si (111) revealed very interesting nucleation property and solved the enigma related to the almost impossible VLS growth of GaAs NWs on Si by VPE methods. It was demonstrated that the presence of silicon in Au–Ga–As droplets suppresses nucleation as long as a certain critical Ga concentration is not reached. We speculate that more silicon atoms enter the droplets in MOVPE and HVPE due to enhanced silicon etching by chemical precursors, thus inhibiting nucleation of NWs.

Si atoms play another big role as being doping element for GaAs. Si-doped GaAs NWs were grown by Au-assisted VLS-HVPE and a model for Si doping was developed. Si atoms are preferentially incorporated into As rather than Ga sites, because of a much lower As/Ga ratio prevailing in the mother phase during VLS growth compared to VS growth. This explains why most VLS Si-doped GaAs NWs are p-type, whereas the standard planar VS growth leads to n-type doping. N-type doping of GaAs NWs with Si is expected at higher As concentrations in the droplet, which occurs for VLS-HVPE growth on Si substrates. These results can be used for obtaining n-type or p-type Si doping in GaAs NWs by tuning the growth conditions, or even Si-doped p–n junctions within a single NW.

The development of low cost gold-free growth protocols by VPE techniques still remains an issue. Defect-free GaAs NWs were successfully grown by Ga-assisted VLS-HVPE on Si (111) substrates. However, as it is also observed in MOVPE, the NW density was drastically low. Actually, metal droplets can hardly be formed and stabilised from chloride gaseous precursors at temperature involved in HVPE. The presence of Si atoms in the droplets also hinders NW...
nucleation as already discussed, which affects both HVPE and MOVPE processes.

For InAs NWs, the HVPE environment is more favorable. Recently and for the first time, direct growth of vertically aligned InAs NWs on silicon substrate was demonstrated by HVPE. The NWs grow along the (111)B direction with a constant diameter. The growth occurs through pure VS condensation. It is mainly governed by the surface kinetics of adsorption, desorption and decomposition (dechlorination) of adsorbed species. However, the NWs still exhibit polytypism. Further work is in progress to improve the crystalline quality of the NWs versus the growth conditions.

HVPE is the third epitaxial technique to be used for the synthesis of NW arrays either by VLS, VS or SAG on patterned substrates. Because HVPE operates at conditions of temperature and vapor phase composition different from those of MBE and MOVPE, HVPE experimental results helped in solving issues encountered by MBE and MOVPE, and strengthened assertions upon growth and nucleation mechanisms. The low cost HVPE technique is now engaged and strengthened assertions upon growth and nucleation mechanisms, especially where long NWs are needed.

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