Research Article
Exploring the Internal Radiative Efficiency of Selective Area Nanowires

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Nanowires are ideal building blocks for next-generation solar cell applications. Nanowires grown with the selective area (SA) approach, in particular, have demonstrated very high material quality, thanks to high growth temperature, defect-free crystalline structure, and absence of external catalysts, especially in the InP material system. A comprehensive study on the influence of growth conditions and device processing on optical emission is still necessary though. This article presents an investigation of the nanowire optical properties, performed in order to optimize the internal radiative efficiency. In an initial preamble, the motivation for this study is discussed, as well as the morphology and crystallinity of the nanowires. The effect on the nanowire photoluminescence of several intrinsic and extrinsic parameters and factors are then presented in three sections: first, the influence of basic growth conditions such as the temperature and the precursor ratio is studied. Subsequently, the effects of varying dopant molar flows are explored, keeping in mind the intended solar cell application. Third, the manner in which the processing and the passivation affect the nanowire optical emission is discussed. Precise control of the growth conditions allows maximizing the nanowire internal radiative efficiency and thus their performance in solar cells and other optoelectronic devices.

1. Introduction

Photovoltaic (PV) technology can provide a substantial portion of electricity generation, and great progress has been observed in the recent decades, with the field projected to have 1 TW of installed power by 2020. Its development is still curbed by the opposite thrusts of low cost and high efficiency though. Solar cells based on nanowire arrays hold promise to solve the current impasse, as they have many advantages: they allow fast growth [1] and lower material consumption by enabling substrate reuse [2] and thanks to high absorption-to-volume ratio through light-trapping [3, 4]. Furthermore, their tolerance to lattice mismatch allows unprecedented material integration [5], and their geometry allows unconventional current extraction designs [6, 7]. Finally, they have an enhanced performance compared to planar layers, as a result of geometry-related increased photon outcoupling and reduced bulk recombination due to their lower volume [8]. Most nanowire growth is carried out by using the vapor-liquid-solid method, which uses a catalyst particle to grow a pillar-shaped structure [9–11]. Selective area nanowires, instead, are obtained by using a substrate covered with a growth mask, from which nanostructures shaped as columns can be grown by carefully tuning the reactor conditions. They have been extensively studied in the last 10 years, with extremely promising results in lasing, transistor, and solar cell applications [12–14]. In particular, selective area InP nanowires have demonstrated very good PV device performance when measured in a single nanowire geometry [15, 16].

The performance can be further improved by careful examination of growth conditions and device processing, and their influence on the photoluminescence emitted by the nanowires. By optimizing the optical emission, the open-circuit voltage can be substantially enhanced [8]. The external radiative efficiency \( \eta_{\text{ext}} \) is in fact expressed as the ratio between the number of radiative recombination events that lead to photons emitted out of the device and the total...
recombination events. The \( \eta_{\text{ext}}^{\text{PL}} \) is closely related to the efficiency of a solar cell device, in particular to the open-circuit voltage \( V_{\text{oc}} \), as expressed clearly by the equation

\[
V_{\text{oc}} = V_{\text{oc}}^{\text{rad}} - \frac{k_B T}{q} \ln \left( \frac{\eta_{\text{ext}}^{\text{PL}}}{\eta_{\text{int}}^{\text{PL}}} \right),
\]

where \( V_{\text{oc}}^{\text{rad}} \) represents the \( V_{\text{oc}} \) in the radiative limit, meaning with external radiative efficiency of 1; \( k_B \) is Boltzmann’s constant; \( T \) is the device temperature; and \( q \) is the electronic charge. This equation can be rewritten in terms of the PL intensity of the sample, which, being directly proportional to the \( \eta_{\text{ext}}^{\text{PL}} \), becomes a direct relation between the PL and the \( V_{\text{oc}} \):

\[
I_{\text{PL}}(E) = a(E) I_{\text{bb}}(E) \left( \exp \left( \frac{q V_{\text{oc}}}{kT} \right) - 1 \right),
\]

where \( E \) is the energy of the emission, \( kT \) is the thermal energy (25.7 meV at room temperature), \( a(E) \) is the absorptivity, and \( I_{\text{bb}}(E) \) is the blackbody emission intensity. An increase in the radiative efficiency is then directly connected to an increase in the PL intensity, which thus corresponds to a \( V_{\text{oc}} \) enhancement.

Additionally, if the internal radiative efficiency \( \eta_{\text{int}}^{\text{PL}} \) is low (<0.1) as in the case of the nanowires considered here, the \( \eta_{\text{ext}}^{\text{PL}} \) can be defined as

\[
\eta_{\text{ext}}^{\text{PL}} = \eta_{\text{int}}^{\text{PL}} \times P_{\text{exc}},
\]

where \( P_{\text{exc}} \) is the average photon escape probability.

In this article, the focus is on the maximization of the first term by changing the growth conditions and studying the influence of device processing, assuming all along that the second factor is constant.

2. Methods

A 50 nm thick silicon nitride layer is used as the growth mask, patterned by soft contact nanoimprint lithography [17] on a (111)A-oriented p-doped InP substrate (Zn doping carrier concentration \( 2 \times 10^{18} \) cm\(^{-3} \) from AXT, USA). InP nanowires are normally obtained at relatively high temperature (650–750°C) and low ratio between group V (PH\(_3\) for P) and group III (TMI for In) precursors (10–200), at a pressure of 100 mbar in an H\(_2\) flow of 151/min, in a laminar MOVPE reactor. Diethylzinc (DEZn) is used as the p-dopant, with molar fractions typically between 1 \( \times 10^{-6} \) and 7.4 \( \times 10^{-5} \). Di-tert-butyl silane (DTBSi) is used as n-dopant, with molar fractions within 5 \( \times 10^{-9} \) and 1.6 \( \times 10^{-6} \). The nanowire arrays are imaged on cleaved samples by using a Zeiss Sigma field emission scanning electron microscope (SEM) operating at 3 kV. Imaging is performed at either 30° or 90° tilt compared to the normal to the surface. TEM and STEM analyses were conducted using a JEOL JEM-ARM200F aberration-corrected TEM operated at 200 kV, while for the chemical analysis, energy-dispersive X-ray (EDX) measurements were carried out using a 100 mm\(^2\) Centurio silicon drift detector (SDD).

After growth, either the photoluminescence (PL) of nanowires is measured directly or nanowire array solar cell devices are fabricated using a process described previously [18, 19]. A custom-built PL setup is used to measure the optical emission of the nanowires. Microphotoluminescence (μPL) studies are carried out on nanowire arrays (either as-grown or processed). To image the specimen and to focus the laser beam, lenses or objectives (with 50x/100x magnification) are used. Nanowire samples are mounted on the cold finger of a flow cryostat, which allows controlling the temperature between helium cooled (∼4 K) and room temperature (293 K). All measurements in this article are performed at 300 K. The cryostat is mounted on an \( x \)–\( y \) translation stage for basic positioning, while fine aligning in \( x \), \( y \), and \( z \) directions is performed by piezoelectric actuators (with 30 nm resolution) controlling the objective. The optical emission of the NW is directed towards the entrance slit of a triple grating SP2500A (Princeton Instruments, 50 cm focal length). A white lamp coupled with a camera is used for controlling the location of the incident laser on the sample. Time-resolved PL measurements are done using a time-correlated single-photon counting module (TCSPC, PicoHarp 300), which measures the time delay between the sample excitation and the arrival of emitted photons at a single photon detector (SPD). After the measurement is repeated many times, the delay times are sorted in a histogram plotting the time distribution of the emission after the excitation pulse [20]. As the temperature has an effect on the PL, all measurements are performed at room temperature (∼300 K), which is the solar cell operating temperature.

3. Results and Discussion

3.1. Selective Area Nanowire Morphology and Crystallinity. In Figures 1(a) and 1(b) are presented two typical SEM images of undoped selective area nanowires, grown at 730°C for 20 min at a V/III ratio of 114 and TMi molar fraction of 4.7 \( \times 10^{-5} \). The morphology of the sample is extremely uniform, as all nanowires have a hexagonal cross section and have very homogeneous length and diameter. In Figures 1(c)–1(f), several TEM-acquired images are shown, displaying the crystalline properties of a nanowire with a \( p^+ \)–\( p \)–\( i \)–\( n \)–\( n^+ \) doping profile as depicted in panel (c). The doping transitions in the growth procedure are sharp, with doping molar fractions 7.4 \( \times 10^{-5} \)–0–4 \( \times 10^{-7} \)–1.6 \( \times 10^{-6} \), respectively. Apart from the very highly n-doped top part of the nanowire, which has a high density of stacking faults, the nanowires have a perfect wurtzite crystalline structure, showing no effect of doping on the crystallinity. Undoped nanowires have an identical crystalline structure. A sample grown with linearly increasing n-doping during growth, characterized by both atom probe tomography and TEM [21], reveals no stacking faults up to a doping molar flow of ∼10\(^{-6}\), further proving the purity of the crystalline structure.

The growth of nanowires with a wurtzite crystalline structure on a zinc blende substrate is explained by the fact
Figure 1: SEM and TEM of selective area InP nanowires. (a) SEM image, inclined 30° respective to the normal, of SA i–InP. The scale bar is 1 μm. (b) Top view SEM image of i–InP nanowires taken normal to the substrate. The scale bar is 1 μm. (c) Overview of a bright-field TEM image of a representative nanowire with a p⁺–p–i–n–n⁺ doping profile. The nominal design of the doping profile is depicted in the image as well. (d) Selected area electron diffraction pattern, acquired from a 1.3 μm-long segment of the middle part of the nanowire, displaying the wurtzite structure. (e) BF-TEM image of the top of the wire (red arrow panel (c)), displaying the defected top n⁻-doped part and the defect-free rest of the NW. (f) HRTEM image of the center of the nanowire (blue arrow panel (c)), displaying the perfect wurtzite structure.
that in the typical SA growth conditions (high temperature and low V/III ratio), the desorption of P is significant, and thus the surface is terminated with In atoms. In this configuration, growth along the $1\over 111$A is only possible if additional In atoms are adsorbed: the WZ stacking though is favored as the $ABAB$ stacking is more energetically stable, due to the larger Coulomb interaction between In atoms and the second nearest pair of In and P [22].

3.2. $I_{PL}^{\text{peak}}$: Effect of Growth Temperature and V/III Ratio. The first two parameters taken into account are basic growth settings such as the temperature in the reactor and the V/III ratio, which are the two most elementary variables adjustable during MOVPE growth.

Temperature is a crucial parameter in nanowire growth, as it has been shown to change the crystalline structure of the nanowires, to influence the concentration of intrinsic and extrinsic defects, and in general to dramatically affect optoelectronic properties of nanowires [23–25]. In particular, for InP selective area growth, temperature increase beyond $\sim 675°C$ is demonstrated to lead to pure wurtzite crystalline structure and a strong enhancement of PL intensity [12]. In Figure 2(a), a series of PL spectra, acquired at room temperature for samples grown at increasing temperatures, up to $770°C$, is depicted. No growth is observed at temperatures of $780°C$ or higher. In all cases, irrespective of the incident laser power, the PL spectra show a peak with a shoulder at higher energy, representing, respectively, the band edge

![Figure 2](image-url)
emission and the split-off valence band (often labeled as A and B transitions) of WZ InP nanowires. The ratio between the two peak intensities is different among different samples, hinting at a complex underlying mechanism. While all measurements are performed as-grown, with nanowires standing on the (111)A InP substrate, no contribution to the PL from the bulk ZB InP, expected at an energy of 1.35 eV, is observed. The intensity, and thus the optical quality expressed as $n_{\text{PL}}$, increases for higher growth temperature up to 730°C, while it decreases at higher temperatures. 

The external radiative efficiency for the best sample, which is also one of the highest among all the samples following in this article (since both doping and processing decrease the PL performance), is estimated as $\sim 10^{-4}$ at 1-sun illumination intensity. This value is comparable to what has been measured for InP bulk layers grown by MOCVD, but lower than nanowires etched by ICP-RIE (respectively, $5.2 \times 10^{-4}$ and $7 \times 10^{-4}$) [26]. The second parameter to be investigated is the V/III ratio: a series of samples is realized by keeping the TMi molar fraction constant at $4.7 \times 10^{-3}$, and varying the PH3, such that the proportion between the two is adjusted between 41 and 164. A clear increase is witnessed up to a quotient of 114, while a further raise is detrimental (Figures 2(d)–2(e)). No influence of the V/III ratio on the shape of the PL peaks is observed in this case (Figure 2(f)).

The PL intensity in this case is influenced mainly by a modification in the density of point defects, such as the introduction of P vacancies [27], which are included because of the increased evaporation of precursors and change in the surface reconstruction of the nanowire top surface, which are both expected with rising temperature [28]. Radiative recombination is then reduced at extremely high temperature by the accumulation of point defects. This explanation is supported by the fact that at temperatures higher than 750°C the growth rate is lower, and at 780°C growth stops entirely: as P atoms evaporate progressively more quickly, crystalline growth is not supported anymore, and growth cannot occur. This mechanism also explains the V/III ratio series results, as the high temperature would result in increased precursor evaporation: In from the precursor has high vapor pressure already at lower temperatures, but as the temperature is increased, the phosphorous vapor pressure increases as well. As the group V flow is increased with augmenting V/III ratio though, the density of P-related point defects would be diminished, so that the PL intensity increases. Two other possible explanations that may partially explain the observed effect on PL, in conjunction with the dissociation of P, are the increased precursor pyrolysis at high temperature, which would reduce the incorporation of carbon in the nanowires [29], and an alteration of crystalline structure, as the density of stacking faults has been shown to be dependent on the V/III ratio [12, 30] and to influence the PL intensity [24].

3.3. Doping Effect on $n_{\text{PL}}$. Control of the presence, the concentration, and the profile of dopant atoms is fundamental for nanowire solar cells: the presence of impurities, charged in respect to their environment, creates the built-in voltage that separates photogenerated carriers and ultimately enables the operation of the device and the power generation. 

Dopants also influence the PL intensity of the material, which as just discussed impacts profoundly the $V_{\text{oc}}$: careful design of the doping is thus crucial. The PL spectra of doped semiconductors are affected both in shape and in intensity by doping. The Burstein-Moss (BM) shift, an effect also known as Pauli blocking, is caused by the occupation of higher energy states in the conduction band due to the Pauli exclusion rule between carriers. The BM shift hence results in an apparent increase of the bandgap by an amount defined as

$$\Delta E_{\text{BM}} = \frac{\hbar}{2m^*} (3\pi^2 n)^{2/3}, \quad (4)$$

where $\hbar$ is the reduced Planck constant, while $m^*$ and $n$ are the carrier effective mass and concentration, respectively. As it is dependent on the effective mass, this effect is much more pronounced for electrons than for holes. If generalized for the nonparabolicity of the conduction band, and substituting the values for electrons in bulk ZB InP, the formula then becomes [31]

$$\Delta E_{\text{BM}}(\text{eV}) = 4.73 \times 10^{-14} n_e^{2/3} - 13.83 \times 10^{-28} n_e^{4/3}, \quad (5)$$

An effect contrasting the BM shift is the bandgap renormalization (BGR), which leads to a narrowing of the bandgap and an increase in the FWHM. It is caused by many-body effects: the net balance between Coulomb and spin interaction among carriers is attractive, basically resulting in an increase in the valence band edge and a decrease in the conduction band edge. BGR is commonly described by an empirical relation where the coefficients are measured experimentally [31]:

$$\Delta E_{\text{BGR}}(\text{eV}) = An^{1/3} + Bn^{1/4} + Cn^{1/2}$$

$$= 17.2 \times 10^{-9} n^{1/3} + 2.62 \times 10^{-7} \times n^{1/4} + 17.3 \times 10^{-12} \times n^{1/2}. \quad (6)$$

Since electrons have a rather low effective mass in InP (~0.077), the effective optical gap at room temperature can be easily increased by degenerated doping. While this can be difficult to believe and to achieve for other material systems, it is instead very effective in InP. S doping of $3 \times 10^{19} \text{ cm}^{-3}$, for instance, results in a bandgap of 1.55 eV, an increase by 0.2 eV [32].

In this section, the PL intensity and the minority carrier recombination lifetime are examined for varying p- and n-doping molar flows.

3.3.1. n-Doping. Si is a common dopant element for InP: since it is a group IV material, it is in principle an amphoteric element, as it could function both as a donor and as an acceptor. In practice though, its amphoteric behavior is very limited (only $< 10^{-3}$ of all Si atoms behave as an acceptor) [33], so it is used efficiently as a dopant in thin film devices, often employing ditertiarybutyl silane (DTBSI) as a precursor [34, 35]. At growth temperatures above 630°C, Si incorporation in MOCVD growth is efficient and can result in
very high concentrations, up to $10^{20} \text{cm}^{-3}$. Comparable concentrations have been measured by atom probe tomography also in the SA-grown nanowires presented in this article [21], with a linear dependence on the incorporation of the dopant on the molar flow, up to $10^{19} \text{cm}^{-3}$, without any noticeable influence on the growth rate or the diameter of the nanowires.

A large effect on their optical properties is observed instead, as n-doping is known to influence the shape of the PL spectrum. The combined effect of the BM shift and BGR can be seen in Figure 3(a), where the normalized spectra of as-grown nanowires with several different doping molar fractions (from $5 \times 10^{-9}$ to $1.2 \times 10^{-6}$) are depicted. The PL peak shifts with the increasing doping concentration; at the same time, the FWHM of the peak broadens, as it has been observed also before with increasing the doping concentration [36, 37]. Another peak at $\sim 1.6 \text{eV}$ is evident in most

Figure 3: Effect of Si and Zn doping on the photoluminescence of nanowires. (a) Series of spectra normalized to the maximum of each one, with different DTBSi-doping molar fractions, between $5 \times 10^{-9}$ and $1.2 \times 10^{-6}$, showing a peak shifting to higher energy and an increase in FWHM, with increasing Si. (b) Time-resolved photoluminescence of the same samples, measured at a laser intensity of 1.4 W/cm², highlighting the decreased minority carrier recombination lifetime with larger dopant molar flow. (c) PL intensity as a function of incident laser intensity, converted in multiples of AM1.5 sun intensity: by rising Si incorporation, the emission strength decreases sharply. (d) Series of spectra normalized to the maximum of each one, with different DEZn-doping molar fractions, from $10^{-6}$ to $7.4 \times 10^{-5}$, showing no dependence of the peak shape on the molar flow. (e) Time-resolved photoluminescence of the same samples, measured at a laser intensity of 1.4 W/cm², depicting the decrease in minority carrier recombination lifetime corresponding to a larger dopant molar flow. (f) PL intensity as a function of incident laser intensity, converted in multiples of AM1.5 sun intensity: as the DEZn fraction increases, the intensity is monotonically enhanced.
spectra and is explained by the mixing and superposition of the A and B transitions, due to the larger FWHM associated with a higher doping level. The doping concentration is determined both by the BM and BGR shift and by double-checking this with atom probe tomography (APT) calibration performed on analogous nanowires, grown with a step-like doping distribution from $5 \times 10^{-7}$ to $9 \times 10^{-6}$. At the highest DTBSi molar fraction presented here ($1.2 \times 10^{-6}$), the doping concentration is thus estimated to be $2 \times 10^{18}$ cm$^{-3}$. The dopant concentration appears to be linearly dependent on the molar fraction used, but at low molar fraction values, it is challenging to reliably estimate a concentration value by atom probe measurements. As the combined effect of BM and BGR shifts is very small at concentrations $<5 \times 10^{17}$ cm$^{-3}$, it is difficult to determine precisely the doping level if the molar fraction is $\lesssim 5 \times 10^{-7}$: a direct proportionality can then only be assumed, as is done later in this article.

Time-resolved PL measurements, performed with a time-correlated single-photon counting (TCSPC) module, presented in Figure 3(b), reveal that increasing the doping concentration impacts dramatically the minority-carrier recombination lifetime: starting from a reference level of 1.8 ns measured for nominally undoped InP nanowires grown in identical conditions, the lifetime decreases sharply with higher DTBSi, until the lifetime is shorter than the instrument response function (limiting the measured laser decay time to ~0.3 ns) at a molar fraction higher than $5 \times 10^{-7}$, which corresponds to a doping level of ~$10^{18}$ cm$^{-3}$, as just discussed.

Regarding the PL intensity, in Figure 3(c), a series of laser power-dependent PL for the different dopants is presented. The laser power is converted to sun intensity, with 1 sun corresponding to the power intensity of AM1.5 illumination (~100 mW/cm$^2$). It is clear that by increasing the dopant concentration by almost three orders of magnitude, the time-correlated single-photon counting (TCSPC) module, subsequent discussion, it is now crucial to distinguish three behaviors: this is shown further in Figure 4, where the peak energy (scatter dots) as a function of the dopant concentration impacts dramatically the minority-carrier recombination lifetime of the same samples instead, where the increase in doping concentration results in a shorter lifetime, from around 1 ns (for $10^{-6}$ fraction) to ~0.6 ns at the highest molar fraction (Figure 3(e)). Furthermore, the PL intensity of p-doped NW has a different trend than for n-doped, as can be seen in Figure 3(f). An increase in PL intensity is observed for increasing doping molar flow, in sharp contrast to the Si-doped case: the intensity is approximately a factor 15 higher for the largest molar fraction, which would result in an enhancement of $V_{oc}$ of ~70 mV.

3.3.3. p- and n-Doping PL Discussion. For the sake of the subsequent discussion, it is now crucial to distinguish three quantities related to doping. While valid also for the previous sections, they assume particular relevant in this examination:

1. The dopant molar fraction, expressed as a pure number (without unit), represents simply the nominal ratio between the amount of dopant precursor flowing in the reactor chamber and the total flow present, as can be calculated from the reactor source values (bubbler temperature, partial pressure, and dilution).

2. The dopant concentration in the semiconductor is expressed as a density (commonly in cm$^{-3}$) of the number of dopant atoms, or more generally impurity atoms, present in a material. They supply free carriers and can act as radiative or nonradiative recombination centers. Within a certain limit, which roughly speaking corresponds to a dopant concentration much lower than the solubility limit of the element in a material, they are dependent linearly on the molar fraction introduced. The atomic density measured by APT provides a very accurate estimation of this density.

3. The carrier concentration in the semiconductor, also expressed as a density in cm$^{-3}$, is the number of electrically active (free) carriers diffusing in a semiconductor, or more often in a region of it. The thermally excited carrier density is altered by the presence of dopant atoms and by photogenerated carriers created by incident light. They are influenced by several factors such as temperature, the presence of other impurities, the doping profile design, and bias voltages.

As it is clear from the previous two sections, p- and n-doping of SA nanowires have dramatically different behaviors: this is shown further in Figure 4, where the peak energy (scatter dots) as a function of the dopant molar fractions are directly compared.

It appears clear how n-doping strongly influences the position of the peak, effectively leading to a shift in the
emission energy of up to 40 meV, corresponding to a doping of \( \approx 2 \times 10^{18} \text{ cm}^{-3} \) as confirmed by APT. p-Doping does not affect the peak energy, so that the emission peak is always centered at \( \approx 1.42 \text{ eV} \), hinting that the carrier concentration in n-doped samples is much higher than in p-doped samples. Zn is known to be more difficult to incorporate if the growth temperature is raised over \( \approx 700\,\text{°C} \) [39, 40]; recent results published in literature also support this conclusion [41, 42]. This inference is further confirmed by measurements performed with APT of nanowires identical to those presented here: at the highest Zn molar fraction (\( 7.4 \times 10^{-6} \)), the Zn concentration is \( \approx 10^{17} \text{ cm}^{-3} \), which is difficult to discern from the ppm noise level of the instrument (1 ppm is \( 4 \times 10^{16} \text{ cm}^{-3} \) in InP). At lower molar fractions, thus, the dopant concentration cannot be estimated.

p- and n-doping influences the total intensity of PL in opposite ways: increasing Si doping leads to a decrease in emission strength (about a factor 3), while augmenting the DEZn molar fraction appears to be beneficial, leading to a 10-fold increase in PL intensity (Figure 5). It is interesting to note that the PL intensity of p-doped nanowires is always much smaller than that of n-doped though (between factors of 3 and 1000, depending on the values used to compare). Furthermore, increasing the dopant molar fraction results in a shorter minority recombination lifetime in both cases, as shown in Figure 5(b).

Two key equations must be introduced in order to explain this phenomenon. First, it must be considered that, under any kind of excitation, the radiative recombination rate is defined as

\[
R_{\text{ext}} \propto (n_0 + \Delta n)(p_0 + \Delta p), \tag{7}
\]

in which \( R_{\text{ext}} \) is the radiative recombination rate, \( n_0 \) (\( p_0 \)) are the equilibrium concentrations of electrons (holes), and \( \Delta n \) (\( \Delta p \)) are the density of excited carriers (in this case, photoexcited). In particular, it is important to note how, following this equation, an increase in the doping atom concentration (\( n_0 \) or \( p_0 \)) should result in an increase in \( R_{\text{ext}} \), which results in a decrease in minority carrier recombination lifetime. An increase in the recombination rate would then correspond to an increase in the PL intensity (when normalized to the integration time) and thus in an increase in the \( \eta_{\text{pl}}^{\text{ext}} \), which is, if the \( P_{\text{exc}} \) is constant, means that the \( \eta_{\text{pl}}^{\text{ext}} \) is enhanced. The contribution of nonradiative recombination must also be taken into account though, as expressed by the relation

\[
\eta_{\text{pl}}^{\text{ext}} = \frac{r_{\text{r}}^{-1}}{r_{\text{r}}^{-1} + r_{\text{rad}}^{-1}}, \tag{8}
\]

where \( r_{\text{r}}^{-1} \) is directly related to the radiative recombination rate \( R_{\text{ext}} \) just introduced [8, 44].

It is thus clear that two mechanisms take place when the dopant concentration is increased: as the equilibrium carrier concentration is larger, leading to an increase in the \( R_{\text{ext}} \), the \( r_{\text{rad}}^{-1} \) is also increased at the same time, as the number of impurity-related nonradiative recombination centers is increased. The different behaviors of undoped and p- and n-doped nanowires are then explained by comparing their dependence on carrier concentration of the PL intensity (\( \text{cc} \eta_{\text{pl}}^{\text{ext}} \)) and the minority carrier recombination lifetime (\( \text{cc} R_{\text{ext}} \)) (Figure 5):

(i) n-Doped NW: at the lowest DTBSi molar fraction considered, the dopant atom concentration is already relatively high (\( \approx 8 \times 10^{16} \text{ cm}^{-3} \)), so that as the dopant concentration is increased, the contribution to \( \eta_{\text{pl}}^{\text{ext}} \) of nonradiative recombination is rising more quickly than the radiative recombination is, as is expected in this doping range: the PL intensity is thus lowered [32]. This is further proved by the fact that with increasing carrier concentration (assumed to be linearly dependent on the dopant molar fraction), the radiative recombination rate does not increase linearly, but appears to saturate very quickly (Figures 5(b) and 5(c)). The saturation in the recombination rate is then attributed to the increased contribution from nonradiative recombination; as the PL intensity is nevertheless larger than in the p-doped case, it would be explained by a nonradiative recombination rate lower on absolute terms.

(ii) p-Doped NW: the introduction of Zn increases the recombination rate as \( p_0 \) is increased (as in equation (7)), and consequently the radiative lifetime is reduced. The PL intensity increase with higher DEZn is explained by the fact that since the doping is still rather low (\( < 10^{17} \text{ cm}^{-3} \)), the dominant variation in \( \eta_{\text{pl}}^{\text{ext}} \) would be the increased \( r_{\text{r}}^{-1} \). In comparison to the undoped case, however, with a small amount of Zn atoms, the \( r_{\text{rad}}^{-1} \) is suggested to be much higher, as the PL intensity is much lower (cf. equation (8)). Raising the DEZn molar fraction and thus the dopant atom concentration, which is the concentration of impurities, does not appear to augment \( r_{\text{rad}}^{-1} \) further in a substantial manner instead. In the range available, in fact, if a similar...

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**Figure 4**: Influence of varying n- (in red) and p- (blue) doping molar fraction on the energy of the PL peak. The x-axis is not in scale, for image clarity.
doping atom concentration is considered, p-doped NW have a lower lifetime and a correspondent higher recombination rate than n-doped NW (Figures 5(b) and 5(c)), so no saturation is observed, in contrast with the n-doped case. Several publications have already demonstrated how the internal radiative efficiency of p-doped InP is much lower than n-doped: this is often attributed to a large density of deep trap levels related to Zn-induced neutral acceptor centers, which result in nonradiative recombination dominance [45, 46]. The PL intensity has an ambiguous behavior though, as it is very high for undoped NW, is heavily reduced for the lowest DEZn molar fraction, and increases slightly again as DEZn becomes larger. A possible explanation is that since the molar fraction of DEZn is rather high (comparable or even larger than the TMI), it is densely present in the chamber. Even if it is not incorporated in the bulk of the nanowires, it can still accumulate at the surface of the nanowire lateral facets, where it would heavily increase the surface recombination. A possible explanation for the different incorporation of Zn on the top surface (where the axial growth takes place) and the lateral facets (where no radial growth is observed) is explained by the fact that lateral facets have different crystal-line orientations, which result in a dissimilar Zn atom adsorption [28]. The adsorption is not influenced by an increased DEZn molar fraction, as even at the lowest flow there would be an excess of Zn atoms. Zn is often deemed responsible for stimulating point defect constitution, for example, following the introduction of interstitial Zn or a phosphorus divacancy linked to Zn (\textit{V_p} – Zn\textit{i} – \textit{V_p}) [47]. The incorporation in the core part of the nanowires is instead dependent on the molar fraction, resulting in the strange behavior of PL intensity. Atoms present at the surface are difficult to measure by APT or other techniques though, so it is difficult to prove unequivocally that Zn deposited at the surface is responsible for this phenomenon.
Figures 6(a) and 6(b), several different cleaning solutions are thus evaluated by measuring the integrated intensity of PL emission of intrinsic nanowires. Three cleaning solutions, all in combination with ALD deposition, which as previously said have been demonstrated to substantially increase the nanowire optical properties [48].

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3.4. Effect of Device Processing. As the intended solar cell application of the nanowires presented in this study requires several processing steps, it is important to study the effect of device fabrication on the nanowire PL. The procedure, discussed previously in detail [18], starts with a cleaning and passivation step and continues with nanowire array planarization and front-side contacting.

Nanowires possess a rather high surface-to-volume ratio, which is around 5 times larger than the surface of a bulk layer if cylindrical NW are considered, with 2 μm length, 500 nm pitch, and 180 nm diameter. Surface protection and passivation is thus crucial: in the case of VLS InP nanowires, it has been demonstrated that during growth the surface easily incorporates defects, which cannot be removed by the in situ HCl etching. These imperfections are eliminated by removing the outer part of the nanowires with a short postgrowth ex situ Piranha etch (25 s in H₂SO₄ : H₂O₂ : H₂O₃ : 1 : 1), which allows improving the efficiency of the device from ~1% up to 11.1% [18, 19]. For selective area nanowires, both a dielectric coating and a lattice-mismatched window layer have been proposed as performance-enhancing passivations [14, 41], together with ALD deposition, which as previously said have been demonstrated to substantially increase the nanowire optical properties [48].

(ii) Undoped NW: the radiative lifetime is relatively long (~1.8 ns) compared to doped nanowires, so the recombination rate must be small. Nevertheless, since the PL and thus the internal radiative efficiency are high, the nonradiative recombination rate must be very low. It is reasonable, since no intentional impurities are introduced, so the imperfection and trap density should be small. Surface recombination can be expected to be the major source of nonradiative recombination.

In the case of bulk InP, the reported minority carrier radiative recombination lifetime (a few tens of ns) is much longer than for nanowires discussed in the previous section though (~1 ns): surface recombination is undoubtedly responsible for a large part of this decrease, so an adequate passivation design should be implemented in nanowires to improve their performance. Atomic layer deposition (ALD) of PO₃ₓAl₂O₃ thin layers, deposited on the same undoped nanowires previously mentioned, has been demonstrated to greatly increase radiative lifetime. A 3-fold enhancement of radiative lifetime (up to >5 ns) is achieved, which results in a ∼τ_{rad} enhancement of more than an order of magnitude, since τ_{rad} is increased at the same time as τ_{rad} is decreased [48].
H₃PO₄ and HF instead, in combination with SiO₂ deposition, both increase significantly the PL intensity: this is attributed to the gentle oxide etch performed by both solutions. HF yields a slightly better performance as a result of the different chemistry of the postcleaning process. In fact, H₃PO₄ cleaning results in a hydrophobic surface which is terminated by hydrogen atoms bound to phosphorous sites, while HF produces a hydrophilic fluorine-terminated surface on the indium sites [50]. When heated during the SiO₂ deposition under N₂ atmosphere, the F-termination is removed quickly, so that SiO₂ would be deposited directly on the nanowires. The H atoms resulting from H₃PO₄, instead, are not removed and thus alter the surface of the nanowires and the interface with the deposited SiO₂ [50]. Additionally, a significant improvement has also been demonstrated by HF passivation in single-nanowire solar cells [16]. Dielectric coating is also expected to increase the absorbance of the nanowires, but it cannot explain a ~3 – fold increase in PL intensity, which is thus attributed to the beneficial passivation. A further demonstration of the enhanced performance is in Figure 6(c), where time-resolved photoluminescence of p – n nanowires grown at 730°C with a V/III ratio of 114 is compared between an as-grown sample and one with the best passivation scheme HF + SiO₂. A clear improvement in the minority carrier radiative recombination lifetime is observed, from 0.5 ns to 0.8 ns, which suggests a reduction of surface recombination thanks to the passivation. The shorter lifetime compared with the intrinsic case presented in the previous section is caused by the doping (molar fractions are 4 × 10⁻⁸ for DTBSi, 7.4 × 10⁻⁹ for DEZn), which as discussed previously decreases the minority carrier recombination lifetime. p – n nanowires have also been tested at all further stages of device fabrication: no significant effect on the PL intensity is observed at different steps. This is expected since no further material passivation is performed, and all processing is performed at low temperature, so that the nanowires are not affected by it. The variation between samples is in the order of ±10%, which is the same range of deviation as between different identical samples and regions of the same sample.

4. Conclusion

The internal radiative efficiency of the nanowires is studied and optimized by varying growth conditions, such as the temperature and the doping molar flows. It is found that the ideal parameters are a temperature of 730°C and a V/III ratio of 114, while a complex influence of dopants is determined. Both dopants result in a decrease in the minority carrier recombination lifetime and a reduction in the PL intensity compared to undoped nanowires. In particular, the introduction of Zn, the p-dopant, results in a strong decrease in the PL intensity of the nanowires, caused by the inclusion of nonradiative recombination centers. Furthermore, by investigating the effect of device fabrication steps, it is found that with a cleaning and passivation procedure based on diluted HF and SiO₂ deposition, the nanowire optical properties are improved.

Data Availability

The photoluminescence data used to support the findings of this study are available from the corresponding author A. Cavalli upon request (at a.cavalli.phd@gmail.com).

Conflicts of Interest

The authors declare no conflicts of interest.

Authors’ Contributions

A.C. conceived and designed the experiment, as well as fabricated and measured the samples, and wrote the manuscript. J.E.M.H. and E.P.A.M.B. provided the theoretical background to the study and the result interpretation, as well as supervised the study and strongly supported and encouraged its development. All authors reviewed, discussed, and approved the results and conclusions of this article.

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