A Non-Oxidative Approach Towards Hybrid Silicon Nanowire-Based Solar Cell Heterojunctions

Abstract
A general method for the non-oxidative termination of silicon nanowires (Si NWs) is reviewed. Oxide-free Si NW have been successfully alkylated in the lab using a two-step chlorination/alkylation process. The distinctive properties of the resulting Si NW have been taken advantage of by integrating the Si NWs into functional devices such as solar cells. Moreover, molecularly terminated Si NWs exhibit lower defect density emissions than unmodified Si NWs. This, in part, explains the better performance of the molecularly terminated Si NW-based solar cells. Solar cells that use organic-inorganic hybrid Si NWs as absorbers show an increased open-circuit voltage ($V_{oc}$), an increased short-circuit current ($I_{sc}$) and a higher fill factor (FF). The aim of chemical functionalization is to protect Si NWs from extensive oxidation, add functionality and to adjust surface electronic properties such as the work function, surface Fermi level and band bending. The stability of the terminated of Si NWs was found to be dependent on the molecular chain length, molecular coverage, interaction type (θ-h or e-e), surface energy and Si NW diameter.

Keywords
Silicon Nanowire • Oxide Free Silicon • Two-Step Process Chlorination/Aklylation Process • Hybrid Solar Cell • Oxidation Resistance • Photoemission • Heterojunction

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1. Introduction
In recent years, a wide variety of one-dimensional nanostructures requiring many different synthesis, modification and characterization methods, have been extensively studied [1,2]. Nanowires are considered to be an important category of one-dimensional nanostructures with promising characteristics for applications in electronics [3], optoelectronics [4], photovoltaics [2,5-15] and sensing [16-18].

Silicon nanowires (Si NWs) have received the attention of many scientists in the last decade as a means to decrease electronic device size while preserving the overall industrial facility production process, thus reducing process redesign costs [19-21]. Si NWs have been characterized through a wide variety of methods such as electron microscopy, x-ray photoelectron spectroscopy (XPS), electron beam induced current (EBIC)/electron backscattered diffraction (EBSD), photoluminescence (PL) and FT-IR spectroscopy, Raman spectroscopy, etc. Moreover, their successful integration into electronic and photovoltaic devices such as field effect transistors (FETs) [12,19,22,23] and solar cells [7,11,24-26] has revealed their potential to become the mainstream building blocks of future nanodevices.

The fabrication of hybrid Si NWs in combination with organic and inorganic materials via a bottom up route, allowing for fairly accurate control over the NW diameter, length and doping, is currently under research. The large surface area per unit volume of NWs makes them highly sensitive to surface characteristics such as surface morphology, topography and physical/chemical bonds with other atoms and molecules. Indeed, the termination of surface dangling bonds with chemical and/or biochemical moieties is expected to have a significant impact on the final physical and chemical properties of Si NWs. Therefore, the Si NW-based composites can be scaled down to the molecular level by applying surface functionalizations, which cover the Si NW surfaces with molecules bonded to individual surface atoms. The resulting Si NWs are known as “hybrid-Si NWs” [27,28].

The presence of oxide in Si NW surfaces is undesirable for subsequent applications and can have a large detrimental effect on Si NW device performance. When hybrid Si NW devices are exposed to air, oxidizing agents such as water and oxygen form an undesirable layer of oxide with high impurity levels, which can result in uncontrolled oxide/silicon interfaces. Thus, to be effective, Si NWs require a form of protection against oxidation. Surface bonds, which are formed during the functionalization of NWs, can enhance oxidation resistance. For example, etched Si NWs and their surface-bound hydrogen atoms exhibit low surface recombination velocities [29]. However, these surfaces tend to oxidize within a few minutes. Another method to functionalize the Si NW with organic molecule is through Si-C

* E-mail: Muhammad.Bashouti@mpl.mpg.de
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Organic Functionalization via Chlorination/Alkylation: One of the most significant methods employed for Si NWs surface functionalization, among those developed for planar silicon, is the versatile two-step chlorination/alkylation process wherein oxide-free silicon surfaces are first covered by chlorine atoms and later by silicon-alkyl surface bonds. The chlorination/alkylation process was carried out in a reflux system and the water temperature was kept to 5-10 ºC with a slow flow of ~10ml/min. The hot-plate temperature was kept at 80 ºC. (See Figure 2)

Scanning Electron Microscopy: To observe the geometry of as-grown Si NWs and to assure that no alteration was caused by H-termination and oxidation, SEM was occasionally performed on samples with a S-4800 Hitachi microscope. The voltage and current used for this purpose were 5.0 kV and 10.0 μA, respectively. The morphologically sensitive secondary electron (SE) detection mode was employed.

2. Experimental Procedure

Synthesis and Growth of Si NW: Si NWs were prepared by the vapor–liquid–solid (VLS) method using chemical vapor deposition (CVD) with silane protected on Si(111) substrates, (See Figure 1).

The VLS growth sequence can be summarized in the following steps: (a) Evaporation of growth species and their diffusion and dissolution into liquid droplets, (b) Diffusion and precipitation of saturated species at the liquid – substrate interface, (c) Nucleation and growth of desired material on the interface and (d) Separation of droplets from the substrate by further precipitation and the growth of NWs. To synthesize the Si NWs, substrates were etched in dilute HF to remove the native oxide and subsequently covered with a 2 nm thick Au film by sputtering and then transferred into the CVD chamber. The substrates were annealed at 580 ºC and a pressure of 5 × 10⁻² mbar for 10 min. The temperature was then reduced to 520 ºC and a mixture of 5–10 sccm (standard cm³min⁻¹) argon and 5 sccm silane (SiH₄) was introduced for 20 min at a pressure of 0.5–2 mbar to allow for the nanowire growth. The resulting Si NWs have a distribution of diameters and lengths which have been measured through Scanning Electron Microscopy (SEM), as depicted in Figure 1.

Termination with a Hydrogen Group: To evaluate the thermal stability of hydrogen atoms bonded to Si NW (H-Si NW) surfaces and determine the dominant oxidation mechanisms, H-Si NWs were annealed at atmospheric conditions at six distinct temperatures: 50°C, 75°C, 150°C, 200°C, 300°C, and 400°C, each for five different time-spans: 5, 10, 20, 30, and 60 min. The annealing and hydrogen-termination processes were gentle in the sense that the processes did not melt the Si NWs or change their diameters. It should be noted that the Si NW were immersed in 5% HF for 1min and in NH₄F for an additional 1min.
thicknesses were 1.6±0.4nm. The Si NW grew along the [112] axis showing a single (111) axial twin crossing the entire length.

Organic Functionalization via Chlorination/Alkylation: One of the most significant methods employed for Si NWs surface functionalization, among those developed for planar silicon, is the versatile two-step chlorination/alkylation process wherein oxide-free silicon surfaces are first covered by chlorine atoms and later by silicon-alkyl surface bonds [30,34]. The chlorination/alkylation process is schematically shown in Figure 5 and can be summarized in the following three steps: (i) termination with hydrogen bonds: Si NW samples are dipped sequentially in buffered HF, (ii) Chlorine termination (Chlorination): H-terminated Si NWs are immersed into a saturated PCl₅ solution, (iii) Organic termination (Alkylation): Cl-terminated Si NWs are immersed in alkyl Grignard (R-MgCl).

X-Ray photoelectron Spectroscopy: In this study, X-ray photoelectron spectroscopy (XPS) was extensively utilized to physically and chemically investigate Si NW surfaces. Outputs of XPS measurements are given as spectra in which electron intensities are a function of kinetic energy (or binding energy). Figure 3 depicts a schematic layout of an XPS system including all important components. Core level and valance band photoelectron spectra were excited by monochromatic Al K radiation (1,487 eV) and collected at a take-off angle of 35° by a hemispherical analyzer with an adjustable overall resolution between 0.8 and 1.2 eV. The surveys were conducted in various electron energy ranges, including the overall binding energy survey (0 to 1,000 eV) and individual spectra for Si 2p (95.0 to 110.0 eV), C 1s (282.0 to 287.0 eV) and O 1s (520 to 550 eV), which were monitored more accurately in a discrete number of scans. All spectra were taken at room temperature in a UHV chamber of about 10⁻⁹-10⁻¹⁰ Torr pressure. The resulting XPS spectra were analyzed and oxide levels were determined by spectral decomposition using the XPS peak software.

3. Results and Discussion

Although the term “hybrid material” is used to describe many diverse systems covering a widespread array of materials, we limit our definition to composite structures with physical or chemical interactions between their organic and inorganic units. In this respect, two classes can be recognized: Class (I) hybrid materials show weak interactions between the two phases, such as van der Waals, hydrogen bonding or weak electrostatic interactions; Class (II) hybrid materials show strong chemical interactions between the components, such as covalent bonding between silicon and organic molecules (Si-C bond). An example of a class II system is illustrated in Figure 4, where alkyl molecules are chemically attached to Si NW via Si-C bonds.

As shown in Figure 1 and 4, VLS grown Si NW with lengths of 3±1μm and diameters of 60±10nm were obtained, the VLS mechanism is described in Ref. [33]. The fabrication of Si NWs with a controlled diameter, length, and electronic properties is essential to technical applications. The SEM image evidences that Si NWs were grown in random orientations. The resulting NWs contain thin oxide layers formed during growth, SiO₂.

Figure 3. Schematic layout of an X-ray Photoelectron Spectroscopy (XPS) system including the analyzer, sample stage, X-ray anode, X-ray monochromator and electron gun.

Figure 4. Hybrid Si NW: The organic molecules (alkyl as example) are chemically connected to Si NW by covalent bonds Si-C. Reproduced with permission from ref [27].
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the respective activation energies \( E_{A}^{\text{ox}} \) based on the Arrhenius equation. In the high temperature region, Si 2D and Si NWs exhibit similar \( E_{A}^{\text{ox}} \) values of 46.35 meV and 48.22 meV, respectively. At low temperatures, \( E_{A}^{\text{ox}} \) values were estimated as 35.20 meV for Si 2D and 23.31 meV for Si NWs [39]. The lower \( E_{A}^{\text{ox}} \) of Si NWs may correspond to the strain induced by dangling bonds rebonding at the (110) and (111) adjoining surfaces as well as the higher ratio of the ambient exposure area to the oxide/silicon interfacial area [40]. The differences in the activation energies of Si NW in the high and low temperatures reveal different oxidation kinetic mechanisms. At low temperatures, increases in the total oxide intensity are accompanied by a rise in the suboxide intensity with amounts comparable to the intensity of SiO\(_2\). Backbond oxidation can be considered as the primary mechanism causing

**Figure 5.** Scheme illustrating the functionalization of Si NWs through the two-step chlorination/alkylation process.

**Figure 6.** Schematic diagram of SiOx/Si interface.

**Figure 7.** XPS spectrum of Si2p core level emission showing two silicon and four oxide peaks.

**Hydrogen Termination**

In order to understand the stability of the organic molecules on the Si NW surface, we terminate the Si NW with a hydrogen group (i.e. H-Si NW). This has three main advantages (i) it allows us to explore oxidation during very fast stages since hydrogen is only stable for a few minutes (less than 5min), (ii) hydrogen is considered as the smallest group, with a lower diameter than the distance between two adjust Si-Si atoms, which as a result, give us full termination i.e. all the Si surface atoms are attached to hydrogen, and (iii) H-terminated is the starting surface for the chlorination/alkylation step. The thermal stability of the hydrogen-terminated Si NWs was investigated by means of the surface sensitive XPS for a variety of temperatures and times. (See experimental section). The oxide growth on Si NWs comprises the formation of interfacial sub-stoichiometric oxides, termed as **transient oxides** including: Si\(_2\)O (n=1), SiO (n=2) and Si\(_2\)O\(_3\) (n=3), followed by the stoichiometric or full oxide SiO\(_2\) (n=4), [35,36] as schematically illustrated in Figure 6.

To quantitatively evaluate the oxidation process, Si 2p spectral decomposition was conducted on the spectra. Consequently, the Si 2p spectra includes two silicon spin-splitting peaks as Si 2p\(_{1/2}\) and Si 2p\(_{3/2}\) (in the regime of 99-100 eV). (See Figure 7)

The amount of each suboxide present, relative to the amount of intact silicon, can be calculated by dividing the integrated area under the suboxide’s peak \( A_{\text{subox}} \) by the sum of the integrated area under the Si 2p\(_{1/2}\) and Si 2p\(_{3/2}\) peaks \( A_{\text{Si}2p_{1/2}} + A_{\text{Si}2p_{3/2}} \). The resulting value is the suboxide intensity, shown by \( I_{\text{subox}} \). In addition, the total oxide intensity \( I_{\text{ox}} \) can be calculated as the sum of the four suboxide intensities \( I_{\text{ox}} = I_{\text{Si}2O} + I_{\text{Si}O} + I_{\text{Si}2O3} + I_{\text{Si}O2} \). The total oxide intensity and suboxide intensities and shifts for the Si NWs are illustrated in Figure 8 and compared to hydrogen terminated planar surface (100). Except for SiO\(_2\), all the suboxide intensities are comparable with more or less very slight variations over the annealing time. However, Bashouti and co-authors observed different mechanisms at low temperatures (from 25°C to 150°C), in which suboxides hold a larger share of the total oxide intensity than at high temperatures (200°C to 400°C) where SiO\(_2\) is the main contributor to the overall detected oxide amount [37].

To understand the thermal stability of hydrogen bonds in the two different temperature regimes, logarithm of the oxidation rate as a function of the reciprocal temperatures in the Si 2D and H-Si NW specimens during the early stages (0-10min) was plotted (Figure 8) [38]. The near-linear behavior allows us to calculate the respective activation energies \( E_{A}^{\text{ox}} \) based on the Arrhenius equation. In the high temperature region, Si 2D and Si NWs exhibit similar \( E_{A}^{\text{ox}} \) of 46.35 meV and 48.22 meV, respectively. At low temperatures, \( E_{A}^{\text{ox}} \) values were estimated as 35.20 meV for Si 2D and 23.31 meV for Si NWs [39]. The lower \( E_{A}^{\text{ox}} \) of Si NWs may correspond to the strain induced by dangling bonds rebonding at the (110) and (111) adjoining surfaces as well as the higher ratio of the ambient exposure area to the oxide/silicon interfacial area [40]. The differences in the activation energies of Si NW in the high and low temperatures reveal different oxidation kinetic mechanisms. At low temperatures, increases in the total oxide intensity are accompanied by a rise in the suboxide intensity with amounts comparable to the intensity of SiO\(_2\). Backbond oxidation can be considered as the primary mechanism causing
the formation of Si-O-Si bonds below the surface-terminated Si-H bonds. The backbonds can be oxidized in different oxidation states and can finally form the full oxide layer atop. Compared to planar samples, Si NWs exhibit a more rapid backbond oxidation, indicating the effect of circumferential tensile stresses on the stability of Si-Si bonds. For longer oxidation times, an increased number of oxidized backbonds form and isolated Si-OH bonds begin to form upon interaction of the Si-H and Si-O bonds with oxidized backbonds [41]. In this regime, the surface energy also takes on a role, which may explain the suppression of the oxidation level compared to the 2D case. By completion of the backbond oxidation, remaining Si-H surface bonds start to rupture and hydrogen propagation begins. The low-temperature oxidation mechanism is summarized in the scheme illustrated in Figure 9. It should be noted that the hydroxyl groups shown in Figure 9 represent both the isolated hydroxyl groups formed throughout the oxidation and the isolated hydroxyl groups formed after the completion of oxidation, as is frequent for SiO₂ in H₂O-containing environments [42].

At high temperatures, the retarded oxidation in Si NWs, in analogy with planar silicon, can be attributed to the self-limited oxidation caused by the function of the initially-formed oxide layer as a diffusion barrier. The two mechanisms are summarized in Figure 10.

It is crucial to study the oxidative behavior of organic molecules to deduce their impact on Si NW oxidation resistance. Specifically, low temperatures (below 200 °C) are the main interest of this study, since almost all electronic devices are operated in this range.

Si₂p emission

Before chemical treatment, Si NWs show a native oxide as illustrated previously in Figure 6. The oxide layer can be detected by XPS of the oxide core level spectrum. The Si₂p of the oxide peak appears in the 101-104 eV binding energy range. In order to
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thus, nearly full surface coverage is possible (See Figure 13).

It should be noted that the Si2p emission spectra of SiO2-Si NW is shifted towards higher binding energy, which will be explained in the solar cell section.

Carbon 1S emission:
The emission of the C1s regime before and after termination is compared in order to describe the covalent attachment of the molecules to the surfaces. When comparing the two emissions, an additional peak must be introduced for molecule-terminated Si NW surfaces to obtain an appropriate deconvolution (Figure 12). Figure 12 depicts a XPS spectrum of C1s emission region for the methyl-terminated Si NWs. Before termination, the C1s spectrum was fitted to two C-C at 285.20±0.02 eV, and C-O at 286.69±0.02 eV. It is reasonable to assume that these adventitious hydrocarbons (C-O) could stem from wet chemical processing with a THF/methanol rinse before functionalization, and/or carbonaceous materials present in the laboratory environment. After termination, the alkyl-Si NW was fitted to three peaks: C-Si at 284.11±0.02 eV, C-C at 285.20±0.02 eV and C-O at 286.69±0.02 eV. The new peak at C-Si at 284.11±0.02 eV confirms the chemical bonding of organic molecules onto the Si NW surface. Peaks were typically adjusted to produce fits that minimized the difference between the full widths at half-maximum. The center-to-center distances were fixed at 1.10±0.10 eV between the C-Si and C-C emissions and at 2.60±0.10 eV between the C-O and C-Si emissions [27].

How to Calculates the Coverage of the molecules on the Si NW Surface?
Methyl (i.e. CH3) is the smallest organic alkyl molecule with a van der Waals diameter (VDW) of 2.5 Å, which is smaller than the internuclear distance between adjacent Si toms (3.8 Å), and thus, nearly full surface coverage is possible (See Figure 13).

To obtain the molecular coverage of surface atomic sites, the area under the C-Si peak is divided by the area under the Si2p peak (sum of Si2p1/2 and Si2p 3/2) and normalized by the scan time. The coverage of the alkyl functionalities was compared with the highest value obtained for methyl (C1) functionality; in this instance “highest” means nearly full coverage of the Si atop sites, achieved after alkylation times of >24 h. This comparison is expressed throughout the text as “(C-Si/Si2p)alkyl/(C-Si/Si2p)max.methyl”.

Termination the Si NW with different Alkyl molecules
The concentration of the molecules on the Si NW (i.e. coverage) was found to be dependent on steric effects caused by the lateral interactions between the terminating molecules on the surfaces. Steric effects and respective VDW diameter can hinder the formation of dense packing alkyls longer than methyl (i.e. C1).
not observed in the case of Si NWs. (3) The experimental error and/or accuracy of the peak fittings are relatively small (4-11%). It is reasonable to consider the passivation of Si NWs by alkyl molecules to be a function of two main factors: (I) molecule-molecule lateral interaction and (II) molecule-substrate vertical interaction. For short alkyl chains (C1-C5), which exhibit liquid-like behavior and thermal fluctuations, the determining factor is the vertical interaction [43]. Increasing the chain length to C6-C10 forms a solid-like phase, where the lateral interactions become dominant in the consequent coverage of the Si NW surfaces. It is worth pointing out that the lateral interactions between long alkyl chains might be formed during the physisorption stage, before the covalent bindings between the carbon and silicon atoms (or chemisorption) arise [44].

Stability of functionalized Si NW
The stability of Si NWs functionalized by organic molecules is affected by the molecular chain length, coverage level, surface energy and diameter. The stability plays a large role in the oxidation process. We will discuss these parameters here.

Effect of coverage and chain length: The atomic sites not connected to the alkyl chains, which are known as atomic pinholes, determine the susceptibility of the surfaces to oxidation. Noting that the peak intensity is a function of the VDW diameter, which increases with the length of the alkyl chains, it can be inferred that the coverage obtained for > C6 chain lengths is artificial and/or not significant owing to problems in fitting the XPS data. In contrast, the absolute coverage of C10 on Si NWs, as shown in Figure 14, has been shown to be consistent with the similar decay of the coverage reported by others on flat Si(111) surfaces for longer chains than in the present work, with no increase in coverage above 7 carbon atoms. Based on these inconsistencies it can be inferred that the higher coverage obtained for > C6 chain lengths is artificial and/or not significant owing to problems in fitting the XPS data. In contrast, a few experimental observations suggest that the coverage behavior of chains longer than C6 on Si NWs might not be artificial: (1) ToF-SIMS experiments (which eliminate potential artificial observations) have shown higher absolute coverage on long chain lengths, in good consistency with XPS observations. (2) In the case of 2D Si(100) surface, the results of ToF-SIMS have shown (more or less) the expected coverage versus chain length behavior; at the time, similar behavior was

**Figure 13.** Schematic diagram of methyl connected to adjacent Si atoms.

**Figure 14.** T\textsubscript{\textit{max-alkyl}} versus alkyl chain length on Si NWs and, for comparison, on 2D Si(100) surfaces. Note: T\textsubscript{\textit{max-alkyl}} (C-Si/Si2p)\textsubscript{\textit{max-alkyl}}/(C-Si/Si2p)\textsubscript{\textit{max-C1}}. Reproduced with permission from ref [43].
interaction with oxidants (O$_2$ and/or H$_2$O). To investigate the effect of the coverage and chain length, freshly-prepared alkylated Si NW functionalities were exposed to ambient air for several hundred hours, as shown in Figure 15. All the alkylated Si NW show high oxide resistivity in the first 2 to 3 days of the experiment, but the oxide intensity became considerable after 8 days exposure to air, depended on the chain length and coverage level. For example, after 336 hours of exposure, the $C_2$ (showing 100% coverage) oxide intensity reached 0.03. However, for the same oxidation period, when the coverage decreased to 50% as in case of $C_2$ to $C_5$, the oxide level rose to ~0.13. This implies that Si NWs terminated with $C_2-C_5$ alkyl molecules exhibit less stability against oxidation compared to $C_6$, which show about 2-fold higher oxidation resistance than that of $C_2-C_5$ Si NWs. Nevertheless, when the molecules have the same coverage, as in case of $C_2-C_5$, the observations can be explained by the fact that shorter molecules have a lower oxidation resistance. Decreasing the coverage below 50%, (as in case on $C_2$), decreases the oxidation stability one fold relative to $C_2-C_5$ molecules and 3-4 fold relative to $C_6$ [28].

Effect of surface energy and diameter: To understand the effect of the surface energy and diameter, methyl groups were terminated onto four surfaces: 2D (100), 2D (111), Si NW 50nm in diameter (Si NW$_{50nm}$) and Si NW 25nm in diameter (Si NW$_{25nm}$). Then, all samples were exposed to ambient air for long periods of time. As shown in Figure 16, the stability of methyl groups on Si NW is higher than the stability on planar Si of either the (100) or (111) configuration. After 336 h, NWs show ca.3-fold lower oxidation intensity than the equivalent planar Si surfaces, despite similar initial coverage levels. These observations can be attributed to the stronger Si-C bonds on Si NW surfaces and are supported by the shift in the Si-C bond binding energy from 284.33±0.02 eV (Si NW$_{25nm}$), 284.22±0.02 eV (Si NW$_{50nm}$) to 284.11±0.02 eV for planar 2D Si. The stronger bond in Si NW is attributed to the higher surface energy in Si NW relative to 2D surfaces. The ~0.11 ±0.02 eV higher binding energy observed for CH$_3$-Si NW, compared to equivalent planar CH$_3$-Si(100) and (111) surfaces, can further be ascribed to the higher reactivity of atop sites. Interestingly, CH$_3$-Si NW$_{50nm}$ surfaces exhibits a higher stability against oxidation than equivalent CH$_3$-Si NW$_{25nm}$ surfaces, even though it has a lower binding energy. The higher oxide intensity can be attributed to the diffusion of gold nanoparticles along and inside the Si NW$_{50nm}$. The Au nanoparticles from the catalyst are very mobile and diffuse over the Si NW surfaces (and inside the bulk regime). This effect is enhanced in thin NWs and, as described in our previous report, nanoparticles can accelerate the oxidation on NWs [45]. The higher stability of the 2D(111) relative to the 2D(100) structure is understandable since it naturally has a 15–20% higher coverage than the 2D(100) case [46].

Effect of Bonds type: π-π vs. σ-σ Interactions: The stability of the Si NW surfaces has been found to be affected by the π-π vs. σ-σ interaction. Si NW were embedded with methyl CH$_3$ and propenyl (CH$_3$-CH=CH$_2$). As shown in Figure 17, the oxidation of CH$_3$-CH=CH-Si and CH$_3$-Si NWs began after only ~100 h of exposure and the process was more stable in the presence of propynyl.

After 180hr, CH$_3$-CH=CH-Si show an oxide intensity of 0.015±0.005 that was almost half of the 0.036± 0.012 oxide intensity of SiO$_2$/Si2p, which obtained after 720 h. In contrast, the oxidation of CH$_2$-Si NWs continuously increased after 100 h of exposure, reaching 0.11±0.017 after ~720 h, without showing any indication of termination. The high stability of the CH$_3$-CH=CH-Si NW can be attributed to the π-π interactions between the adjacent molecules [47-49].

Integration of hybrid Si NW into Solar Cells

Analysis suggests that extrinsic effects (i.e. surface states and surface charge) accompanied by the adsorption of molecular layers significantly affect solar cell characteristics. Three different surface terminations of Si NWs are presented to the discussion: (i) H-Si NWs for which the oxide shell was removed by a standard HF dip, (ii) SiO$_2$-Si NWs with a single monolayer of oxide grown after the HF dip (14 days); (iii) CH$_3$-Si NW samples, which were further processed by methyl functionalization immediately after

Figure 15. Observed oxidation intensity (SiO$_2$/Si2p peak ratio) of alkyl-terminated Si NWs at different exposure times to ambient air. Reproduced with permission from ref [28].

Figure 16. Ratio of the oxidized Si2p peak area to the bulk Si2p peak area for the methyl modification of NW$_{25nm}$ and NW$_{50nm}$ and 2D Si(111), exposed to air over extended time periods.
This Si NW-based cell concept has the following advantages:

(i) efficient light absorption
(ii) short diffusion distance of carriers to the wrapped around heterojunction [51,52];
(iii) use of the air-stable and robust polymer, PEDOT:PSS, as an efficient hole conductor [9];
(iv) the Si NW/polymer solar cell design utilizes only 1% of the Si used in other thin film cells of identical thickness. However, for photovoltaic applications, full coverage of Si NWs by PEDOT:PSS is desirable but has not, as of yet, been achieved. Figure 20 shows the current-voltage (I-V) characteristics of CH3-SiNW/PEDOT:PSS and SiO2-Si NW/PEDOT:PSS solar cells under AM1.5 illumination. SiO2-SiNW/PEDOT:PSS samples can withstand a short circuit current (Jsc) of 1.6mA/cm2, an open circuit voltage (Voc) of 320 mV, a fill factor (FF) of 0.53 and a conversion efficiency (µ) of 0.28 %. CH3-SiNW/PEDOT:PSS devices exhibit improved performance with Jsc, Voc, FF and µ magnitudes of 7.0 mA/cm2, 399 mV, 0.44 and 1.2 %, respectively. The low values of FF and Jsc in both

Figure 17. Ratio of the SiO2 to Si2p peak areas for the different surface modifications of Si NWs, exposed to air over extended time periods. Reproduced with permission from ref [47].

Figure 18. Photoelectron yield Y(hν) spectral and spectral density of states of SiO2-Si NW, H-Si NW and CH3-Si NW. Reproduced with permission from ref [38].
types of samples are attributed to high contact resistances ($R_s \approx 3000 \Omega$). It should be noted that by decreasing the resistance, the efficiency of the solar cell can be improved. In spite of notably low efficiencies of Si NW relative to the values commonly reported for efficient solar cells, the comparative increase in efficiency (by about a factor of 4) upon methylation proves that this kind of surface functionalization has very promising prospective.

The improved performance of the CH$_3$-Si NWs is attributed to the removal of the tunneling oxide and the reduction of defect states related to PY, which is caused by a reduced defect density at the heterojunction interface. Using one monolayer of CH$_3$ at the interface allows for a more efficient charge coupling between silicon and the polymer. By reducing the defect density of CH$_3$-Si NWs, the surface recombination is decreased, causing an increase in $V_{oc}$. According to the Shockley diode equation $V_{oc} = kBT/q \ln(J_{sc}/J_0)$, where $J_0$ is the saturation current and the other variables have their usual meanings. Since an increase of $J_{sc}$ influences $V_{oc}$ as well, it must be mentioned that the $V_{oc}$ gain observed cannot be explained by this effect alone. Assuming a similar $J_0$, which correspond to the removal of the tunneling oxide without changing the interface properties, the increase of $J_{sc}$ would lead to a $V_{oc}$ gain of $\Delta V_{oc} = kBT/q \ln(J_{sc,methyl}/J_{sc,oxide}) = kBT/q \ln(7.0/1.6) = 0.037$ V. Regarding the observed gain of $\Delta V_{oc} = 0.079$ V, only a reduced surface recombination's (as measured by PY) and/or a favorable barrier formations (surface dipole) can establish consistency with the experimental data. The well passivation of methyl groups showed an impact in different devices as reported by Lewis as well [22,53,54].

4. Conclusions

Si NWs were functionalized by different organic compounds through a two-step chlorination/alkylation process. Methyl provided the highest coverage of all of the alkyl molecules. Increasing the chain length decreased the coverage to 50–70%. Molecules with double bonds such as propenyl (–CH=CH-CH$_3$) showed excellent surface passivation, with a very small amount of silicon oxide forming after more than two months of exposure to ambient air, which was attributed to the $\pi$-$\pi$ interaction. The thermal stabilities of the organically-tailored Si NWs are dependent on the molecular coverage, chain length, types of bond interaction, $\pi$-$\pi$ or $\sigma$-$\sigma$ interactions, surface energy and Si NW diameter. Studies on the H-terminated Si NW oxidation kinetics revealed that their thermal stability relies strongly on the temperature. At lower temperatures, initially Si-Si backbond oxidation and later Si-H bond propagation are the rate-determining steps. At higher temperatures, oxygen diffusion is considered to be the initial rate-determining step, as it controls the growth site concentration.

The oxide free surfaces are important and necessary in many technological aspects, e.g. radial epitaxy on NWs to realize vertical P-N junctions in photovoltaics or radial Si super-lattices.
in optoelectronics. Hybrid Si NWs have already been integrated in solar cells in order to utilize their advantages. Several fundamental surface properties relevant to solar cells have been found to be strongly dependent on the type of functionalization used. Thus, the right choice of surface modification is crucial to solar cell performance. For instance, Si NW attached to CH$_3$, H or oxide groups produce a core-level shift and band bending in the Si2p emission derived from the Si surface atoms at 50 meV, 100 meV and 170 meV, respectively. The good passivation of or oxide groups produce a core-level shift and band bending in surface density. As a result, solar cells with CH$_3$-based radial heterojunctions demonstrate better performance and efficiencies for Jsc, Voc, FF and $\mu$ of 7.0 mA/cm$^2$, 399 mV, 0.44 and 1.2 %, respectively.

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