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Alkane and Alkanethiol Passivation of Halogenated Ge Nanowires

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Abstract

The ambient stability and surface coverage of halogen (Cl, Br and I) passivated germanium nanowires were investigated by X-ray photoelectron and X-ray photoelectron emission spectroscopy. After exposure to air for 24 h, the stability of the halogen-terminated Ge nanowire surfaces towards re-oxidation was found to improve with the increasing size of the halogen atoms, *i.e.* I > Br > Cl. Halogen termination was effective in removing the native Ge oxide (GeO_x) and could also be utilized for further functionalization. Functionalization of the halogenated Ge nanowires was investigated using alkyl Grignard reagents and alkanethiols. The stability of the alkyl and alkanethiol passivation layers from the different halogen-terminated surfaces was investigated by X-ray photoelectron spectroscopy and attenuated

total reflectance infrared spectroscopy. Alkanethiol functionalized nanowires showed greater resistance against re-oxidation of the Ge surface compared to alkyl functionalization when exposed to ambient conditions for one week.

Keywords: Germanium nanowires, surface passivation, X-ray photoelectron spectroscopy.

Introduction

Group 14 semiconductor nanowires have been successfully fabricated via several different bottom-up and top-down strategies¹. Germanium (Ge) offers potential advantages over silicon (Si) for performance gains in high speed electronic devices due to greater free carrier mobility.²⁻³ There have been advances in Ge nanowire growth and several groups have already demonstrated the fabrication of single Ge nanowire devices such as field effect transistors (FETs) $^{4-6}$ and *p*-*n* junctions⁷. However, Ge possesses an unstable, non-uniform oxide surface on both bulk and nanowire surfaces which gives rise to a poor Ge/GeO_x interface characterized by a high density of surface states⁸⁻⁹. The negative influence of these surface states on the electrical properties of nanowires has been theoretically and experimentally studied¹⁰⁻¹³. The successful integration of Ge nanowires into many device applications consequently requires effective surface oxide removal and passivation. Literature studies have shown the oxidation of Ge surfaces to be a complex process depending on the conditions such as a wet or dry environment, illumination and crystal orientation¹⁴⁻¹⁵. Using high resolution photoelectron spectroscopy (XPS) Schmeisser et al.¹⁶ could resolve all four oxidation states in the Ge 3d spectrum and found a core level shift of 0.85 eV per oxidation state. Unlike Si, which exhibits only one stable oxide (SiO₂), Ge forms stable oxides in the 2+ (GeO) and 4+ (GeO₂) oxidation states, the latter being soluble in water. Prabhakaran and Ogino¹⁷ found that bulk single crystal surfaces of Ge oxidised in a dry O_2 environment, forming predominately Ge^{2+} , while exposure to ambient conditions led to a mixture of oxides, mainly Ge^{2+} and Ge^{4+} , proposing that atmospheric moisture plays a role in the formation of higher Ge oxidation states. Furthermore, the oxidation species observed on nanowire surfaces differs from those reported on bulk planar Ge surfaces, for example thermal annealing of air-oxidised bulk Ge favors the formation of the 2+ species¹⁶, while thermal annealing of water-oxidized Ge nanowires results predominately in the formation of the 1+ oxide¹⁸. This difference in oxidizing behavior is most likely due to the high curvature of the nanowire surfaces. Removal of the surface oxide, GeO_x, is typically achieved by treatment with aqueous HF solution, resulting in hydrogen terminated surfaces¹⁹. The stability of the H-passivation layer on Ge surfaces is limited to a few minutes when exposed to ambient conditions²⁰.

Termination of Ge surfaces with halogens was first achieved by Cullen *et al.*²¹ using hot gaseous HCl. Since then milder passivation methods including dilute halide acids (HCl, HBr, and HI) and electrochemical dissociation of silver halide salts under ultra high vacuum conditions have proved effective for Cl, Br and I passivation of $Ge^{22\cdot24}$. Sun and co-workers²⁵ detected the presence of both monochloride and dichloride species on HCl treated bulk Ge(100) surfaces, while the Ge(111) surface was found to be terminated only by the monochloride which was attributed to the Ge(111) surface having only one dangling bond. They further observed that HF treatment resulted in greater surface roughness compared to HCl treatment due to the greater Ge back-bond breaking that occurs with HF etching²⁵. In a later paper, Sun *et al.*¹⁵ investigated the oxidation behavior of Cl and Br-terminated surfaces and illustrated three important findings (i) Cl/Br-terminated surfaces displayed increased resistance to re-oxidation relative to H-surfaces under dry conditions, (ii) the presence of water vapor resulted in the halogen species being replaced by –OH groups, which allowed for

easier oxidative attack by atmospheric O_2 and water vapor, due to the smaller size of the –OH groups and (iii) the rate of surface oxidation was greatly enhanced by the presence of UV light. While these and other reports^{20, 26-28} have been conducted on bulk single crystal Ge, Adhikari and co-workers²⁹ carried out XPS studies of HF and HCl-treated Ge nanowires with synchrotron radiation and found similar stability trends, *i.e.* chlorinated surfaces displayed an increased stability relative to H-terminated surfaces. To date, Ge nanowire passivation with heavier halogens (Br and I termination) has been reported by Jagannathan *et al.*³⁰

Hydrogen and halogenated surfaces can also be subsequently employed as further scaffolds for the attachment of organic ligands. Unsaturated hydrocarbons have previously been grafted onto H-terminated Si and Ge surfaces³¹⁻³⁴. These hydrosilylation and hydrogermylation reactions can be achieved by thermal activation, UV initiation or Lewis acid mediation³⁵⁻³⁸. The attachment of alkyl chains has been demonstrated on Cl-terminated surfaces using Grignard reagents. This chlorination/alkylation route has been effective for functionalizing both bulk³⁹⁻⁴⁰ and nanowire⁴¹⁻⁴² surfaces of Si and Ge. Ge surfaces passivated with alkyl chains show far greater stability compared to hydrogen or halogenated (Cl /Br /I) surfaces due to the strong Ge-C bond (460 kJ mol⁻¹) and the presence of a hydrophobic monolayer hindering the access of oxidising species towards the Ge surface. Functionalization of Ge with alkanethiols is typically achieved via hydrogen passivated surfaces⁴³⁻⁴⁵ but Bent and co-workers⁴⁶ found that alkanethiol passivation could also be achieved on planar Cl and Br-terminated surfaces.

Here we present a detailed investigation into the relative stability of Cl, Br and I-terminated Ge nanowires using XPS. While previous studies on Ge nanowire passivation have focused on nanowire bundles, we utilize X-ray photoelectron emission microscopy (XPEEM) to

analyze *individual* Br and I terminated Ge nanowire surfaces. We compare the reactivity of these halogenated surfaces towards further functionalization with Grignard reagents and alkanethiols. We further evaluate the effectiveness of alkane and alkanethiol passivation layers, obtained from different halogenated Ge surfaces, to prevent the re-oxidation of Ge nanowires.

Experimental

Ge Nanowire Synthesis and Passivation

The Ge nanowires used in this study were synthesized by the thermal decomposition of diphenylgermane (purchased from ABCR, Germany) in the presence of gold-coated silicon substrates in supercritical (sc) toluene. Details of the experimental set-up have been described elsewhere⁴⁷. The reactions were carried out a temperature and pressure of 400 °C and 24.1 MPa, respectively, yielding nanowires with a mean diameter of 80 nm. The nanowires displayed a predominately <111> growth direction with <110> and <112> growth directions also present.

Diethyl ether (Et₂O) was distilled from Na/benzophenone, anhydrous methanol (MeOH) and isopropyl alcohol (IPA) were purchased from Sigma-Aldrich. All other reagents were purchased from Sigma-Aldrich. Halogen termination of the Ge nanowires was carried out by immersing the nanowires into 10 % aqueous HCl, HBr and 5 % aqueous HI solutions for 10 min. The substrates were washed with deionized water, IPA and dried under N₂. The Ge nanowires were functionalized with alkyl chains via a halogenation/alkylation route using alkyl Grignard reagents. After halogen passivation the nanowires were immersed in 1 M dodecylmagnesium bromide (DD-MgBr) in Et₂O and heated to 45 °C for 24-72 h. The substrates were soaked in anhydrous Et₂O for 5 min and then rinsed with more Et₂O. This soaking/rinsing procedure was repeated 3 times. The nanowires were then rinsed with MeOH and dried under N₂. Ge nanowires were passivated with alkanethiols by immersion into 0.1 M dodecanethiol in anhydrous IPA. The nanowires were heated to 60 °C for 2-24 h under N₂. Following the passivation procedure the substrates were soaked in IPA for 5 min and rinsed with IPA (\times 3). The nanowires were then rinsed with chloroform, MeOH and dried with N₂.

Characterisation of Functionalized Ge Nanowires

Scanning electron microscopy (SEM) images were acquired on a FEI Inspect F, operating at 5 kV accelerating voltage. Transmission electron microscopy (TEM) images were acquired using Jeol 2010 at 200 kV accelerating voltage. Attenuated total reflectance Infra-red (ATR-IR) spectra were recorded on a PerkinElmer Spectrum 100 using 20 scans with 2 cm⁻¹ resolution. The nanowires were dispersed in tetrachloromethane and dropped onto the ATR crystal (ZnSe). The solvent was allowed to dry before the measurements were recorded. Xray photoelectron spectroscopy (XPS) analysis was conducted on a VSW Atom Tech System using achromatic Al X-rays from with a twin anode (Al/Mg) X-ray source. Survey spectra were captured at a pass energy of 100 eV, a step size of 0.7 eV and dwell time of 0.1 ms. The core level spectra obtained were averaged over 15 scans and captured at a pass energy of 50 eV, a step size of 0.2 eV and a dwell time of 0.1 ms. XPS data was also acquired using a KRATOS AXIS 165 monochromatized X-ray photoelectron spectrometer equipped with a dual anode (Mg/Al) source. Survey spectra were captured at as pass energy of 100 eV, step size 1 eV and dwell time of 50 ms. The core level spectra were an average of 10 scans captured at a PE of 25 eV, step size of 0.05 eV and dwell time of 100 ms. The spectra were corrected for charge shift to the C 1s line at a binding energy of 284.6 eV. A Shirley background correction was employed and the peaks were fitted to Voigt profiles. The Ge 3d signals were fitted to two peaks with a spin-orbit coupling of 0.58 eV and an intensity ratio of 3:2, corresponding to the Ge $3d_{5/2}$ and Ge $3d_{3/2}$, respectively. In figure 1, the Ge oxide peaks were plotted by adding separate peak contributions using Gaussian profiles in OriginPro 8. The peaks were centered at 30.5 eV, 31.35 eV, 32.2 eV, 33.05 eV, corresponding to Ge^{1+,} Ge²⁺, Ge3⁺ and Ge⁴⁺, respectively¹⁶. The S 2p doublet peaks were fitted to Voigt profiles with a spin-orbit splitting of 1.2 eV⁴⁸. XPEEM measurements were carried out at the nanospectroscopy beam line at the Elettra synchrotron facility in Trieste, Italy. A detailed description of the beamline set-up is described elsewhere ⁴⁹⁻⁵⁰. The passivated *i.e.* oxidised nanowires were dispersed in IPA and drop cast onto doped Si substrates. Ge nanowire passivation was carried out by immersion into aqueous halide acid solutions. The substrates were rinsed with deionised water, methanol and dried under Ar.

Results and Discussion

Relative Stability of Halogenated Ge Nanowires

Figure 1(a) shows a SEM image of Ge nanowires synthesized on a Au-coated Si substrate. Once removed from the reaction vessel the surface of the nanowires begins to oxidise immediately. The Ge 3d XPS core level spectra shown in figure 1(b) is comprised of an elemental Ge peak, which exhibits spin-orbit splitting of 0.585 eV, consistent with that of the Ge $3d_{5/2}$ and Ge $3d_{3/2}$ peaks, located at 28.6 eV and 29.2 eV, respectively¹⁶. In addition to bulk Ge, 4 chemically shifted satellite peaks at higher binding energies are also present, corresponding to the four Ge surface oxidation states, as illustrated in figure 1(b).

Oxide removal and halogen termination was achieved by treatment with aqueous halide solutions. Jagannathan *et al.*³⁰ previously used 20 % HBr and HI solutions for generating halogen-terminated Ge nanowires. However, we observed considerable roughening of the Ge nanowire surfaces at these concentrations and consequently 10 % HCl or HBr solutions were

used in our study. A 10 % HI solution was found to be particularly aggressive to our nanowires, etching both the surface oxide and the crystalline Ge (with a known etching rate of 0.6 nm min^{-1 51}), leading to very rough surfaces, as shown in figure S1 in the Supporting Information. Rough surfaces have been demonstrated to oxidise faster than smooth surfaces and consequently a 5 % HI solution was used in our studies²⁸.



Figure 1. (a) SEM image of Ge nanowires grown in sc-toluene at a temperature of 400 $^{\circ}$ C and pressure of 24.1 MPa from a Au-coated Si substrate and (b) the corresponding Ge *3d* XPS core level spectra of the Ge nanowires acquired 1 week after synthesis.

Figure 2 compares the Ge *3d* XPS core level spectra of halogen-terminated nanowires after immediate treatment and after ambient exposure (~ 20 °C, 70 % relative humidity). All halide solutions effectively removed the surface oxide as illustrated by the absence of oxide associated peaks in the spectra immediately after acid treatment (blue spectra). After 24 h exposure to air (red spectra), the Cl-terminated nanowires showed the greatest degree of Ge re-oxidation, the HBr treated nanowires displayed only minor oxide formation and the HI treated nanowires showed no re-oxidation of the surface. The I-terminated surfaces exhibited a small oxide peak after 48 h ambient exposure.

increased with the increasing size of the halogen species (Cl < Br < I), as the larger halogen atoms serve as a better steric barrier to prevent re-oxidation of the surface. Differences in the electronegativity of the halogen species also influences the reactivity of the halogenterminated Ge surfaces⁵². Electronegativity values decrease down Group 17 in the periodic table and consequently the electronegativity difference between Ge and the halogen species reduces from Ge-Cl (1.5) to Ge-Br (0.95) to Ge-I (0.65)⁵³, leading to a higher degree of covalent bonding. Furthermore, the increasing orbital size of Cl (*2p*), Br (*3p*) and I (*4p*) means that the length of the Ge-X bond increases from Cl < Br < I. The combined effect of a weaker bond, longer bond length and smaller electronegativity difference reduces the Ge-X bond polarization with increasing halogen size, *i.e.* X = Cl > Br > I⁵². Less polarised Ge-X bonds are thus more resistant to an oxidative nucleophilic attack from O₂ and H₂O species, thereby giving larger halogen species greater stability on the Ge surface.

XPEEM Analysis of Individual Bromine and Iodine Passivated Ge Nanowires

Figure 3(a)-(h) illustrates XPEEM images and the corresponding background-subtracted, normalized Ge 3d and Br 3d spectra. The presence of Br species on the nanowire surface is clearly observed in the Br 3d PEEM image, figure 3(c). Furthermore, the Ge 3d spectra Ge nanowire exhibits a shoulder peak shifted to a lower kinetic energy, which is consistent with Ge bonding to the more electronegative Br atom⁴⁶. The Ge 3d spectra of the iodine-terminated nanowire, shown in figure 3(e) is best fitted with two peaks, one corresponding to Ge and the other chemically shifted by 0.4 eV due to the presence of iodine species. The presence of an iodine-terminated surface is also indicated by the I 4d XPEEM image and corresponding I 4d spectra shown in figures 3(g) and (h), respectively.



Figure 2. Ge *3d* XPS core level spectra showing (a) Ge nanowires immediately after treatment with HCl, HBr and HI and (b) iodide-terminated Ge nanowires after 48 h ambient exposure, and chlorine, bromine and iodine-terminated Ge nanowires after 24 h ambient exposure.

The halogen surface coverage can be estimated from the integral intensities of the Ge and halogen XPEEM spectra. The intensities were corrected for spectra that were collected at different photon energies. A detailed description of the XPEEM data analysis is described elsewhere⁵⁴⁻⁵⁵. The monolayer surface coverage (θ_x) was estimated from equation (1):

$$\theta_x = \frac{I_x / \sigma_x}{(I_x / \sigma_x) + (I_{Ge} / \sigma_{Ge})} \tag{1}$$

Where I_x and I_{Ge} are the integrated intensities of the Ge and halogen species, respectively, and σ_x , σ_{Ge} are the corresponding photoionization cross sections taken from literature values⁵⁶. The estimated values for θ_{Br} and θ_I were found to be 1.04 and 0.91, respectively. The surface

coverage values were an average of 6 Br-terminated and 5 I-terminated nanowires with a standard deviation of 0.09 and 0.13, respectively. It must be noted that errors such as non-linear background and approximations in photoemission cross-sections introduce errors into the surface coverage calculations, estimated to be \pm 0.2. The surface halogen coverage suggest complete termination after HBr/HI treatment, which corresponds to literature studies on planar Ge surfaces^{15, 46}. However, it is difficult to draw conclusions by comparison with studies on planar surfaces as the Ge nanowires possess a predominate <111> growth direction, having {110} surface facets. In contrast to Ge(100) and Ge(111) planar surfaces the Ge(110) surface has been much less investigated. The halogen coverage may also be less then unity if dihalide species are present on the nanowire surfaces.

Alkylation and Thiolation of Halogen-Terminated Ge Nanowires

Figure 4 illustrates XPS spectra of Cl-terminated Ge nanowires as well as alkane and alkanethiol functionalized nanowires, obtained via chlorinated surfaces. After alkylation with DD-MgBr there was an increase in the intensity of the C *Is* peak relative to the chlorinated nanowires. The presence of carbon in the untreated and halogen-terminated nanowires can be attributed to adventitious hydrocarbons adsorbed onto their surfaces and residual carbon contamination from the nanowire synthesis. Furthermore, the absence of the Mg 2p (50 eV) and Br 3d (70 eV) peaks in the spectra is suggestive that the alkyl chains are covalently attached and not merely adsorbed onto the Ge nanowire surfaces. After reacting the Cl-terminated Ge nanowires with Grignard reagents for 24 h, Cl species were still observed in the XPS survey, as shown by the presence of the Cl 2s and Cl 2p peaks at binding energies of 269 and 200 eV respectively (figure 4(a)). After 48 h, there was a reduction in the intensity of the Cl 2s peak, however complete removal of Cl species on the alkylated surfaces was not achieved. The high resolution Cl 2s spectrum shown in figure 4(b), taken after a reaction

time of 72 h, indicates that some Cl atoms still remain on the nanowire surfaces. There was no change in the intensity of Ge 3d:Cl 2s XPS peaks after alkylation times > 72 h. In comparison to alkylation, thiolation reactions on chlorinated Ge surfaces showed no Cl species in the XPS analysis after a reaction time of 4 h, as shown in the high resolution Cl 2sspectrum in figure 4(c). Figure 4(d) displays the high resolution S 2p XPS core level spectrum of the thiolated nanowires centred at 162.7 eV, which is in good agreement of with binding energies reported for thiolated monolayers⁵⁷⁻⁵⁹.

Figure 4(e) illustrates the O *1s* XPS core level spectra for oxidized, chlorinated, alkylated and thiolated nanowires. After HCl treatment, the intensity of the oxide peak reduced considerably due to the removal of GeO_x, but a small oxide signal remained which can be mainly attributed to the presence of adsorbed molecules after aqueous HCl treatment and from the MeOH rinse. Although the Ge *3d* spectra indicated an oxide free surface, reports on planar surfaces have shown that trace amounts of oxide are not always observed in the Ge *3d* spectra can be detected in the Ge *2p* spectra, which is more surface sensitive¹⁷. After alkylation and thiolation there is a slight increase in the intensity of the O (*1s*) peak, most likely attributed to oxygen functionalities in the solvents used for the functionalized reactions (IPA, Et₂O). Reactions of thiols at the Ge surface are more favorable than alcohols due to the lower S-H bond dissociation energy compared to that of the O-H bond⁵².



Figure 3. XPEEM images and spectra of (a-d) bromine-terminated nanowires, illustrating the Ge 3d and Br 3d spectra and (e-h) iodine-terminated Ge nanowires, illustrating the Ge 3d and I 4p spectra.



Figure 4. (a) XPS survey scans of Cl-terminated Ge nanowires and alkylation and thiolation functionalization via chlorinated surfaces, (b) Cl 2s core-level spectrum showing the presence of Cl after an alkylation reaction time of 72 h, (c) Cl 2s core-level spectrum after thiol functionalization, (d) S 2p core-level spectrum after thiolation reaction, (e) O 1s core level spectra of Ge nanowires before and after surface functionalization. The asterisks mark signals from the Si wafer.

Figure 5(a) shows XPS survey scans of brominated and iodated Ge nanowires as well as functionalization of these halogenated surfaces with Grignards and alkanethiols. Bromination of the Ge surface can be seen from the presence of the Br 3d peak located at a binding energy of 69.1 eV. Alkylation and thiolation on Br and I surfaces are both accompanied by an

increase in their respective C *Is* signal. After alkylation there is a reduction, but not a complete disappearance of the halogen species similar to the trend observed on CI-terminated surfaces. The position of the Br *3d* peak at 69.1 eV in figure 5(b) is consistent with Br bonded to a Ge surface; if the Br peak was due to unreacted Grignard regent, *i.e.* DD-MgBr, the Br *3d* peak would be observed at a lower binding energy, as Br bonded to a more electropositive Mg atom (electronegativity (en.) = 1.3) would undergo a larger chemical shift, relative to Ge (en. = 2.0)⁵³. Figure 5(c) shows the I *3d*_{5/2} peak at a binding energy of 632 eV, after HI treatment as well as an I signal after alkyl functionalization. Thiolation of Br and I terminated surfaces was accompanied by the appearance of the S *2p* peak shown in figure 5(c) and the absence any Br and I peaks in the XPS spectra.

The absence of halogen species in the XPS survey spectra, after thiol functionalization indicates that alkanethiols are more effective in replacing surface halogen species compared to alkyl Grignard reagents. After ~72 h immersion in the Grignard solution there is negligible change in the intensity of the Ge:halogen XPS peaks, indicating that further reaction with the remaining halogen species is unfavorable. The mechanism for the covalent attachment of Grignard reagents to Ge surfaces is illustrated in equation (2).

$$R-Mg-X^{1} + Ge-X^{2} \rightarrow Ge-R + MgX^{1}X^{2} \qquad (X^{1}, X^{2} = Cl/Br/I, R = alkyl) \qquad (2)$$



Figure 5. (a) XPS survey spectra of Ge nanowire functionalization on Br and I-terminated surfaces, (b) Br 3d spectra after alkylation. (c) The I 3d spectra before and after iodations. (d) The S 2p spectra after thiolation of Br and I-terminated surfaces.

Grignard reagents are extremely reactive species due to a highly nucleophilic carbon atom adjacent to the Mg atom, consequently the reduced reactivity towards halogen-terminated Ge surfaces may be due to steric constraints. Both DDT and DD-MgBr have similar chain lengths (~ 18 Å) and only differ in the nature of their functional head groups. While Grignard reagents are commonly noted as 'R-MgX' (X = Cl, Br, I), their actual structure in solution is

described by the Schlenk equilibrium which involves the co-ordination of solvent molecules to the Mg atom⁶⁰. Furthermore, ethereal solutions of Grignard reagents in a concentration range of 0.5-1 M exist as dimeric complexes, as illustrated in figure 6^{61} .



Figure 6. Schematic illustrating the co-ordination of solvent molecules to Grignard reagents.

The mechanism for thiolation involves abstraction of the surface halogen to form a corresponding hydrogen halide, as shown in equation⁴⁶ (3):

$$R-S-H + Ge-X \rightarrow Ge-S-R + H-X \qquad (X = Cl/Br/I, R = alkyl)$$
(3)

Although alkylation is carried out at a lower temperature than thiolation (45 °C versus 60 °C) which can be expected to influence the reaction kinetics, increased steric effects experienced by Grignard reagents due to solvent co-ordination may also hinder the ability to access the halogenated species on the nanowire surface consequently resulting in unreacted residual halogen species detected by XPS analysis.

ATR-IR and TEM Analysis of Functionalized Ge Nanowires

Figure 7 shows ATR-IR spectra of DD- (red spectra) and DDT (blue spectra) functionalized Ge nanowires via modification of the initially Cl-, Br- and I-terminated nanowire surfaces.

Stretching vibrational modes associated with aliphatic alkyl chains are visible for all samples (2800-3000 cm⁻¹), further indicating that the Grignard reagents and alkanethiols have reacted with the halogen terminated Ge surfaces. The ATR-IR spectra of alkanethiol passivated nanowires, figure 7(b), show essentially identical absorbance frequencies for Cl, Br and Iterminated surfaces. The C-H asymmetric and symmetric stretches are observed at 2921 and 2851 cm⁻¹, respectively, while the asymmetric CH₃ absorption peak is observed at 2955 cm⁻¹. These peak positions are in good agreement with IR absorbance frequencies reported by Kosuri et al.44 for alkanethiol functionalized bulk Ge surfaces. The peak positions of the C-H stretching modes occur at lower frequencies relative to isotropic liquid DDT, indicating that a degree of crystalline order is present in the alkanethiol passivation layer^{38, 62-63}. For Ge nanowires alkylated from Cl- and Br- terminated surfaces, the symmetric and asymmetric CH₂ stretching modes occur at 2921 cm⁻¹ and 2852 cm⁻¹, similar to those on thiolated surfaces. The asymmetric CH₂ stretching mode of highly crystalline hydrocarbons typically appears at 2918 cm⁻¹ ⁶², suggesting that some disorder is present in the alkyl and thiol functionalization layers. The vibrational modes for alkylation via I-terminated surfaces exhibited the highest absorption frequencies (v_{as}CH₂: 2924 cm⁻¹, v_aCH₂ 2853 cm⁻¹, v_aCH₃: 2959 cm⁻¹) indicating the most disordered passivation layer was achieved via a iodination/alkylation route. The presence of unreacted halogen species or defects at the nanowire surface would be expected to disrupt the ordering and assembly of the passivating ligands.



Figure 7. ATR-FTIR spectra of (a) dodecyl and (b) dodecanethiol-functionalized Ge nanowires from Cl, Br and I-terminated surfaces.

Figures 8(a)-(c) display SEM images of Ge nanowires before and after surface passivation, and show that the morphologies of the nanowires were not altered by the functionalization procedures. Figures 8(d)-(f) show TEM images of Ge nanowires before and after functionalization. The native Ge oxide (GeO_x), typically ~ 2-4 nm, shown in figure 8(d) has been replaced by a thin passivation layer (~1.8 nm) comprised of the alkane and alkanethiol ligands, figure 8(e) and (f), respectively. TEM analysis showed little difference between the thicknesses of the passivation layer formed from different halogen-terminated surfaces.



Figure 8. SEM and TEM images of Ge nanowires (a) and (c) post functionalization, (b) and (e) alkylated with DD-MgBr, (c) and (f) thiolated with DDT.

Stability of Alkyl and Alkanethiol Passivation Layers: Influence of Halogen Species

The degree of re-oxidation of the Ge surface provides insight into the quality of the passivation monolayers attained from the halogenated surfaces. Figures 9(a) and (b) illustrate the XPS Ge 3d peaks for Cl/Br/I surfaces functionalized with alkyl and alkanethiols, respectively, after one week exposure to ambient conditions. Overall, Ge nanowires functionalized with DD chains via Grignard reagents (red spectra) display a higher degree of

re-oxidation compared to DDT (blue spectra) passivated surfaces as indicated by the greater oxide component in the Ge 3d spectra. A comparison of the spectra within figure 8(a) shows that alkyl passivation via chlorinated surfaces exhibit the most oxidation, while passivation via iododated surfaces shows the least. The opposite trend is observed for alkanethiol passivation, with Ge nanowires thiolated from Cl and Br-terminated surfaces showing no oxidation after ambient exposure for 1 week, while thiolation via iodated surfaces do exhibit some re-oxidation. The oxide shifted peak in the Ge 3d XPS core level spectrum is small, indicating only minor oxidation of the surface.



Figure 9. Ge *3d* XPS core level spectra of (a) dodecyl functionalized Ge nanowires and (b) dodecanethiol functionalized Ge nanowires after exposure to ambient conditions for 1 week.

The stability trend observed for alkylated (Cl < Br < I) and thiolated (I < Br \approx Cl) Ge nanowire surfaces can be explained as follows. Thiolation of halogenated surfaces results in

the complete removal of the halogen atoms but alkylation via Grignard reagents does not, therefore the stability of the halogen species must also be considered for the alkylated surfaces. The residual halogen atoms on the surface are more susceptible to oxidation than alkylated surfaces, which are more hydrophobic. In addition, the long hydrocarbon chain length serves as a better steric barrier from atmospheric O₂/H₂O compared to single halogen species and the covalent character of the bond prevents bond cleavage. Furthermore, the Ge-C bond strength is 460 kJ mol⁻¹, which is greater than Ge-X (where X = Cl (356 kJ mol⁻¹) / Br (276 kJ mol⁻¹) / I (213 kJ mol⁻¹))⁵³. As illustrated in figure 2, the stability of halogen termination increases with the heavier atom, consequently the observed trend in alkyl stability parallels that of the halogen stability *i.e.* Cl < Br < I. The residual halogen species to readily gain access to the nanowire surface and consequently alkylated surfaces are considerable more oxidised after 1 week then the thiolated samples.

Thiolation on all halogen terminated Ge surfaces provide relatively good stability (over 1 week), with minor oxidation observed only on nanowires thiolated from iodated surfaces. The Ge-I bond is much less polarized than the Ge-Cl/Br bonds, giving rise to a higher activation energy barrier for alkanethiol attachment, consquently^{46, 59} alkanethiol formation from iodated surfaces is less favorable than chlorinated and brominated Ge surfaces.

Although thiolation on halogenated Ge nanowires has not been reported, several studies of thiolation via hydrogen terminated surfaces have found thiols to provide better protection against surface re-oxidation compared to alkyl chains^{18, 64-65}. In this study, it is evident from figure 9 that thiol functionalization of halogenated surfaces also display greater stability than alkyl functionalization. Interestingly, literature reports on planar Ge surfaces found that alkyl

ligands (Ge-C) impart greater passivation than alkanethiol ligands (Ge-S)⁴⁵. Although there has been no literature studies into the origin of this trend, high surface curvature, surface roughness and the presence of defects are all likely to have some influence on the stability of passivation layer on nanowire surfaces compared to planar surfaces.

Conclusions

In summary, the surface halogenation of Ge nanowires with Cl, Br and I atoms, followed by surface functionalization with alkanes and alkanethiols was investigated. The stability of the halogenated surfaces of the Ge nanowires increased from Cl < Br < I due to the increasing size of the heavier halogen atom which provided a greater steric barrier to oxidative attack. Attachment of dodecyl chains via Grignard reagents did not result in complete removal of the surface halogens, even after long reaction times. Conversely, after thiolation of the nanowire surfaces no halogen species were detected by XPS. Greater steric constraints due to dimerization and solvent co-ordination associated with alkyl Grignard reagents attribute to the reduced reactivity of alkyl functionalization compared to alkanethiols. Incomplete surface functionalization via Grignard reagents was also reflected in stability studies of the alkane and alkanethiol functionalized nanowires. After exposure to ambient conditions for 1 week the alkylated nanowires showed a greater degree of re-oxidation relative to thiolated nanowire surfaces. Furthermore, nanowires alkylated via chlorinated surfaces displayed the greatest degree of Ge oxidation while alkylation via iodated surfaces exhibited the least, a trend which reflects the stability of the residual halogen species on the nanowire surface upon alkylation. On the other hand, alkanethiol passivation layers showed excellent ambient stability; functionalization from Cl and Br surfaces showed no re-oxidation of the surface after 1 week, while those formed from iodated surfaces only exhibited minor oxidation. Overall the results show that alkanethiol functionalization of Ge nanowires can be achieved from halogenated surfaces and that the stability of these passivation layers exceeds that of alkyl layers formed from Grignard reagents.

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Supporting Information Available

TEM image illustrating the surface roughness of Ge nanowires as a result of etching with 10 % HI. Estimation of surface coverage on alkylated Ge nanowires from XPS data.

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Table of Contents Figure and Graphic

Gillian Collins, Peter	Halogen-terminated Ge	ا عط _{5/2} کو ک
Fleming, Sven Barth,	nanowires have been studied by	≥ Solo Br, I
Colm O'Dwyer, John	XPS and XPEEM. Halogen	X-X
Boland, Michael Morris	terminated Ge nanowire	~ -X
and Justin Holmes.	surfaces have been further	50 55 60 -X
	functionalized by alkyl Grignard	Kinetic Energy (eV)
Alkane and Alkanethiol	reagents and alkanethiols.	Ge
Passivation of	Passivation by alkanethiols was	S S S
Halogenated Ge	found to impart greater ambient	C ₁₂ H ₂₅ C ₁₂ H ₂₅ C ₁₂ H ₂₅
Nanowires	stability relative to alkane	Record
	functionalization.	