Copper Nanowires Synthesis and Self-Assembly for Interconnect Applications

Srikar Rao Darmakkolla
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Copper Nanowires Synthesis and Self-Assembly for Interconnect Applications

by

Srikar Rao Darmakkolla

A dissertation submitted in partial fulfillment of the requirements for the degree of

Doctor of Philosophy
in
Chemistry

Dissertation Committee:
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Portland State University
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ABSTRACT

One-dimensional (1D) nanomaterial self-assembly offers an excellent approach to the fabrication of highly complex nanodevices. Despite considerable effort and research, precisely controlling the orientation and positioning of nanowires (NWs) on a large-scale area and assembling into a functional device is still a state of the art problem. This thesis focuses on the dimensionally controlled copper nanowires (Cu NWs) synthesis, and magnetic field assisted self-assembly of cupronickel nanowires (Cu/Ni NWs) into interconnect structures on a carbon doped silicon dioxide (CDO) wafer.

CDO is a low dielectric constant (k) material used for copper interconnects in multilayered complex integrated circuits (ICs). Here, a strong affinity of copper (Cu) and nickel (Ni) to thiol (-SH) functional groups were exploited to strongly adhere the nanowires (Cu/Ni NWs) onto the CDO substrate. Thiol (-SH) functionalization of the CDO surface was achieved via a series of reactions involving (1) esterification of the surface exposed $\equiv$Si-OH functional group to its triflate ($\equiv$Si-O-Tf), (2) reduction of triflate to $\equiv$Si-H using DIBAL-H, and (3) hydrosilylation of $\equiv$Si-H using 2-propene thiol ($\equiv$Si-(CH$_2$)$_3$-SH) in a photochemical reaction. The thiol functionalization of CDO surface enhances the interaction of Cu/Ni NWs with strong chemical bonds. The same reaction scheme was also used in the functionalization of the hydrophilic (Si-OH) surface to the hydrophobic long alkyl chain derivatized ($\equiv$Si-CH$_2$-(CH$_2$)$_{16}$-CH$_3$) surface. This long alkyl chain modified surface acts as an excellent moisture resistant film, which helps to maintain the low-k value of the CDO.
The dimensionally controlled Cu NWs were synthesized by a wet chemical approach. Optimization of the reducing agent, hydrazine (N$_2$H$_4$), controlled the surface morphology of nanowires (NWs). Interestingly, the high concentration of reducing agent produced particle decorated and/or with a rough NW surface, and conversely decreasing its concentration resulted in a comparatively thin, particle-free and smooth surface. The reaction temperature affected the aspect ratio (Length/Diameter) of the NWs. As the reaction temperature increased from 60 to 90 °C, the aspect ratio decreased from 140 to 21.

Controlling the orientation of Cu NWs in a magnetic field was accomplished by coating them with a thin layer (~20 nm) of ferromagnetic nickel (Ni). This Ni-coated NWs showed an excellent degree of alignment (half-width ≈ 10 degrees) in the direction of an applied magnetic field over a large surface area at field strength as low as 2500 Gauss. Also, the Ni coating helped in protecting the copper core from oxidation resulting in better electrical wire-to-wire contacts.

A nanowire-based interconnect channel was fabricated by combining magnetic field assisted alignment and deposition of aligned NWs on a thiol-modified and photolithography patterned CDO substrate. The NWs, deposited in the trenches, strongly bonded to the thiol-derivatized CDO substrate while an acetone wash removed loosely bound NWs on the photoresist surface. In electrical characterization, the directionally well-aligned Cu/Ni NWs channel displayed surprisingly two-fold higher conductivity than randomly arranged NWs channel.
DEDICATION

This dissertation is dedicated to my parents and all my family members.
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I would like to express my deepest gratitude towards my research advisor Dr. Shankar B. Rananavare for his motivation, support, guidance, and encouragement during my studies at Portland State University. I am thankful for teaching me how to approach the difficult problems in research and allowing me to grow as a research scientist. I am blessed to have you as my supervisor for my Ph.D. study.

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<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Meaning</th>
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<tbody>
<tr>
<td>Cu NWs</td>
<td>Copper nanowires</td>
</tr>
<tr>
<td>Cu/Ni NWs</td>
<td>Nickel coated copper nanowires</td>
</tr>
<tr>
<td>SEM</td>
<td>Scanning Electron Microscopy</td>
</tr>
<tr>
<td>TEM</td>
<td>Transmission Electron Microscopy</td>
</tr>
<tr>
<td>CA</td>
<td>Contact angle</td>
</tr>
<tr>
<td>DI</td>
<td>Deionized</td>
</tr>
<tr>
<td>t-EBSD</td>
<td>Transmitted electron backscattered diffraction</td>
</tr>
<tr>
<td>EBSD</td>
<td>Electron backscattered diffraction</td>
</tr>
<tr>
<td>CDO</td>
<td>Carbon doped silicon dioxide</td>
</tr>
<tr>
<td>CMP</td>
<td>Chemical mechanical polishing</td>
</tr>
<tr>
<td>DD</td>
<td>Dual Damascene</td>
</tr>
<tr>
<td>EM</td>
<td>Electromigration</td>
</tr>
<tr>
<td>PET</td>
<td>Polyethylenetetraphthalate</td>
</tr>
<tr>
<td>DB</td>
<td>Diffusion barrier layer</td>
</tr>
<tr>
<td>IC</td>
<td>Integrated circuits</td>
</tr>
<tr>
<td>RC</td>
<td>Resistance and capacitance</td>
</tr>
<tr>
<td>FTIR</td>
<td>Fourier transform infrared spectroscopy</td>
</tr>
<tr>
<td>XPS</td>
<td>X-ray photoelectron spectroscopy</td>
</tr>
<tr>
<td>XRD</td>
<td>X-ray diffraction spectroscopy</td>
</tr>
<tr>
<td>EDS/EDX</td>
<td>Energy dispersive X-ray spectrum</td>
</tr>
<tr>
<td>EDA</td>
<td>Ethylenediamine</td>
</tr>
<tr>
<td>PECVD</td>
<td>Plasma enhanced chemical vapor deposition</td>
</tr>
<tr>
<td>OTS</td>
<td>Octadecyltrichlorosilane</td>
</tr>
<tr>
<td>DIBAL-H</td>
<td>Di isobutyl aluminum hydride</td>
</tr>
</tbody>
</table>
CHAPTER ONE
INTRODUCTION AND RATIONALE

Copper is one of the oldest and the third most industrially consumed metal. It has received a great deal of attention for its wide variety of applications, such as electrical conductor in electronics, architecture, plumbing, human health, and many other consumer products\(^1\). Copper has been the wiring metal of choice in electronic devices because of its ductility, malleability and excellent conductivity characteristics. Especially in chip fabrication, copper is being used as an interconnect material which plays an important role in the power supply and electric signal distribution\(^2\).

![Cross section SEM image of copper multilevel interconnect system with CDO as a low-k dielectric\(^3\) insulator.](image)

**Figure 1.1.** Cross section SEM image of copper multilevel interconnect system with CDO as a low-k dielectric\(^3\) insulator.

In 1997, IBM revolutionized the technology industry with its remarkable innovation of chips made of copper interconnects. Replacing the traditional aluminum with copper as an interconnect material, enabled smaller chip sizes, cheaper production, and faster
performance\textsuperscript{4-6}. Today, in the current ultra large-scale integration (ULSI) technology copper (Cu) and tungsten (W) are being used as interconnect materials. Tungsten is used for ground level interconnects which communicates to the transistors. Copper is being used for all other layers, which can have as many as 12 metallic interconnect levels. Copper metal lines in each level communicate to other levels through copper vias as shown in Figure 1.1.

Ultra large scale integrated (ULSI) circuit device manufacturing technology can be divided into two main stages. The first stage is transistor and its other component structures fabrication. At the second stage, huge numbers of transistors are connected to each other by three-dimensional multilayers of copper interconnect systems\textsuperscript{3, 7}. Transistors are the active components of a chip and are where signals are controlled and generated. Each transistor communicates to other active parts of a chip through electric signals that are carried by highly complex copper interconnect systems. Ultimate speed limit and chip performance are determined by the frequency at which transistors can turn on and off \textsuperscript{8}. Since clock frequency increases with decreasing transistor size, advances in chip speeds have been achieved by shrinking the feature sizes and increasing the count. Currently, the speed limit and performance of a chip is set by signal propagation delays and the time constant of the copper interconnects\textsuperscript{9}.

1.1. Review on traditional process of copper interconnects fabrication on chips

All metal lines and vias of integrated circuits (ICs) are fabricated by an industrially standard Dual Damascene (DD) process\textsuperscript{2}. DD involves defining the line and via pattern in
a dielectric using photolithography technique and backfilling the opening with copper by an electrochemical method. Finally, the excess copper removed by chemical mechanical planarization (CMP) process\textsuperscript{2, 10}.

**Dual Damascene (DD) process**

The Damascene process was initially developed for jewelry manufacturing\textsuperscript{2}. In a single Damascene process trenches and vias are filled with copper in two separate steps. In Dual Damascene (DD) process trench and vias are filled in a single step. In comparison to single Damascene, DD process lowered the number of processing steps and became the industrially standard interconnect fabrication technique. The advantage of this method is super filling, which results in a void-free and seamless filling of high aspect ratio trenches and vias\textsuperscript{10}.

In advanced IC integration, single Damascene is only used for Tungsten (W) local interconnects and all other copper interconnects are made by using Dual Damascene process. All high-density metal lines and vias are insulated by an inner layer dielectric (ILD) material. Copper is known to be a fast diffuser into silicon and its oxide dielectrics. This causes deep level defects which may lead to poor performance or chip failure. By using a Ti, TiN, Ta or TaN as a diffusion barrier, one can localize copper to where it is desired\textsuperscript{2}.
**Figure 1.2.** Schematic of DD process for making metal lines and vias. The process involves diffusion barrier layer (DB) deposition, copper seed layer deposition (CSL), copper filling by electroplating process and removal of excess copper by a chemical mechanical planarization process (CMP)\(^2\).

Major steps involved in DD process are illustrated in Figure 1.2. The DD process starts with depositing a thin layer of a diffusion barrier (DB) layer (Ta, TaN) commonly done by ionized metal plasma (IMP) method. It is followed by depositing a thin copper seed layer (CSL), usually by sputtering. Copper electroplating method emerged as the most promising and commercial viable technique for copper filling\(^8\). A schematic of electrochemical deposition set up is shown in Figure 1.3\(^8\). CSL coated wafer is then immersed in an electroplating solution and electrically connected and acts as a cathode. Copper sulfate (CuSO\(_4\)) and sulfuric acid (H\(_2\)SO\(_4\)) are the common components of
electrolytes. Cupric ions (Cu$^{2+}$) from an electrolyte solution get reduced and deposited on CSL surface which results in super filling of trenches and vias.

**Figure 1.3.** Schematic diagram of electroplating of copper on the copper seed layer (CSL) coated wafer$^8$.

After trenches and vias are filled with copper, the excess copper is removed by Chemical Mechanical Polishing (CMP). In CMP process, polishing with a slurry removes excess copper by the combination of chemical reactivity and mechanical abrasion$^2, 10$. Chemicals in the slurry react with copper and typically form copper oxides and are subsequently removed by mechanical abrasion. After polishing for an optimum period, the wafer surface becomes planar. Capping of Cu interconnects by silicon nitride, or silicon carbide (known as Etch Stop (ES) layer) completes one level of interconnect fabrication. All these above discussed processes will be repeated for multiple times for the fabrication of multilayered highly complex Cu interconnects.
1.2. Challenges in modern integrated circuits fabrication

Today, modern chips use several billion of transistors and other electronic components in an area of about 1 cm$^2$\textsuperscript{11}. As the transistor count increasing, copper wire dimensions are shrinking to nanometer scales to connect these individual transistors.

With the ever-decreasing copper wire dimensions, the resistance of the narrow conductive channel is increasing due to electron scattering by the grain boundaries and at wire surfaces. Increased resistance adversely contributes to the overall signal propagation delay. In the modern chips, total resistance (R) of the copper interconnect structure is a significant factor that affects the chips performance\textsuperscript{9,11}. Furthermore, as the area of copper interconnects increase, the capacitance (C) between the closely squeezed conductive channels also increases. Both effects significantly increase in the RC signal delay, which results in degraded performance of the chips\textsuperscript{9}. Hence, in recent years enormous efforts have been devoted to developing new process technologies to fabricate miniaturized crystalline copper interconnects and to prepare low-$k$ insulators to minimize the capacitance (C) and leakage current.

In the narrow copper wires, mechanical stress and current density significantly increase (>2 MA/cm$^2$) which ultimately leads to electromigration (EM) related interconnect failures\textsuperscript{12,13}. High current density in the narrow metal lines causes void nucleation, propagation along the metal lines and creates cracks\textsuperscript{14}. Electromigration failure mechanisms have been discussed in detail elsewhere\textsuperscript{13,15,16}. Also, the standard DD process requires deposition of diffusion barrier and copper seed layers in the narrow trenches and vias. But, as the feature sizes are shrinking to few nanometer scales, it is becoming very
challenging to have these layers and fill them with copper. Ultimately, the fabrication of interconnects in a DD process is reaching its fundamental limits. Therefore, innovative materials and process solutions are very critical to sustaining the miniaturization and performance trends.

1.3. Alternate materials and methods for interconnect fabrication

In the recent history, bottom-up or hybrid approaches of nanoscale circuit fabrication using nanomaterials as building blocks, have received considerable attention\textsuperscript{17-19}. This kind of bottom-up approach of integrated circuit fabrication involves the synthesis of nanocrystals with desired physical and electrical properties and their self-assembly into electronic circuits\textsuperscript{19}.

Desirable material properties for the fabrication of interconnects:

As the DD processed on-chip copper wire dimensions are reaching to nanoscales, they experience two principal issues. First, electron scattering induced degradation of electrical resistivity. Second, their reliability issues due to poor electromigration (EM) resistance\textsuperscript{17}. In addition to the above two issues, oxidation under ambient conditions and enhanced oxidation at high resistive self-heating are also the major growing concerns\textsuperscript{20}.

The ideal building blocks used for the interconnect fabrication should possess high conductivity, excellent electromigration (EM) resistance, high oxidation resistance and economical\textsuperscript{20}. Therefore, a quest for alternate interconnect materials which address the above issues have been under active search.
**Promising building blocks for the fabrication of interconnects:**

Nanomaterials show significantly improved electrical, mechanical and thermal properties compared to their bulk material counterparts. Especially 1D nanomaterials, such as pure metallic nanowires, core-shell nanowires and carbon nanotubes (CNT), etc., are very attractive building blocks for functional nanoscale electronic devices, owing to its unique anisotropic structure-dependent properties. Assembly of 1D nanomaterials in a bottom-up or hybrid approach offers an excellent alternative for the fabrication of miniaturized and high-performance interconnects\textsuperscript{17, 20-28}. Many kinds of 1D nanomaterials such as pure metallic (Cu\textsuperscript{17, 29}, Ag\textsuperscript{20}, Au\textsuperscript{25}, Pt\textsuperscript{30}) NWs, core-shell NWs (Graphene encapsulated Cu NWs\textsuperscript{31}) and carbon nanotubes\textsuperscript{32} (CNT), etc., have been explored as promising building blocks for interconnects\textsuperscript{17, 20-28}.

Among all 1D nanomaterials, copper nanowires (Cu NWs) have attracted considerable interest in recent years due to their intriguing electrical, thermal and mechanical properties\textsuperscript{17, 18, 33}. Copper has the second highest electrical conductivity among metals, highly abundant and low cost. Thus Cu NWs have been considered as promising building blocks for many applications, such as flexible & transparent electrodes\textsuperscript{33-35}, conductive inks\textsuperscript{18}, catalyst\textsuperscript{36} and interconnects\textsuperscript{17}. Also, copper is successfully being used as an interconnect material on-chip since 1997\textsuperscript{37}. Therefore, Cu NWs are the ideal candidates for many micro and functional nanoscale devices, and it is imperative to develop a simple and reliable technique to assemble the Cu NWs into a functional device.
Review of NW self-assembly techniques:

Self-assembly refers to the process in which nanowires or other discrete components spontaneously organize into ordered macroscopic structures owing to its direct specific interactions (e.g., interparticle forces), or indirectly using an externally applied field\textsuperscript{30, 38, 39}. The essential feature of nanowire self-assembly is an organization of NWs in an orientationally and positionally well-ordered manner\textsuperscript{30, 40}.

![Figure 1.4](image)

**Figure 1.4.** Schematic describing various kinds of nanowires self-assembly techniques that arranges disordered NWs into well-ordered manner\textsuperscript{41}.

In a bottom-up or hybrid approach, precisely controlling the NWs orientation and positioning are the first critical steps for the successful assembly of individual NWs into a functional nanodevice\textsuperscript{38, 42, 43}. Several approaches for controlling the orientation of NWs via a self-assembly process have been studied (See Figure 1.4), such as Langmuir-
Blodgett\textsuperscript{44}, microfluidic flow\textsuperscript{38}, evaporation induced\textsuperscript{38}, nematic liquid crystal phase\textsuperscript{40}, shear induced\textsuperscript{45}, mechanical force\textsuperscript{38}, electrospinning\textsuperscript{41, 46}, electric\textsuperscript{47} and magnetic field assisted techniques\textsuperscript{30, 38, 41-43, 45, 48, 49}.

The Langmuir-Blodgett technique involves compression of NW suspensions at air-water interface into dense films\textsuperscript{44}. During the compression, NWs organize themselves into the well-aligned and closely packed structure. And, dip coating process can then be used for transferring the well-aligned NWs on to a solid substrate. However, the primary challenge in using this approach is, during the dip coating process reorganization of the NWs results in overlapping features and gaps within the dense arrays of NWs\textsuperscript{38, 44}. Evaporation-induced and microfluidic channel assisted alignment techniques involves, reorientation of NWs into well-ordered dense arrays with the direction of fluid flow to minimize the fluid drag\textsuperscript{38}. Confining the fluid flow to the microfluidic channel enhances the shear force and results in more uniform alignment. However, it is hard to control the alignment uniformity in the case of the evaporation induced technique, and the requirement of fabrication of microfluidic channels limits the wafer-scale assembly of NWs using these techniques\textsuperscript{38}.

Nematic liquid crystal phase technique depends on the volume fraction of nanowires or nanorods in a given solvent. In a very dilute solution, NWs can rotate or translate freely under Brownian motion without interference from another NW/rod. But, as the concentration increases to a critical concentration ($\Phi_N$), a nematic liquid crystal phase forms and NWs organize themselves into a densely packed and orientationally ordered manner\textsuperscript{40}. However, the special requirements like monodispersity, high aspect ratio
nanowires and that NWs should not precipitate out from the solution, etc. limits this technique.

Shear force or mechanical force induced reorganization of NWs is one of the large-scale assembly techniques\textsuperscript{38,45}. Here, a shear force is created by the motion of a solid object or fluid against a second solid object or fluid. In either case, a force is generated parallel to the sliding direction and aligns the NWs in the direction of movement. This technique is inexpensive and can be applied to various substrates such as Si wafers and flexible or curved substrates. However, this process requires excellent skills and the challenging thing in this technique is controlling the viscosity of the NWs solution\textsuperscript{38}.

Electrospinning is a versatile technique, using electrostatic forces to form continuous chains of fibers, polymers or nanowires. In a typical electrospinning technique, NWs or a polymer suspension pumped through a thin nozzle with an inner diameter of \( \sim 100 \text{ \textmu m} \)\textsuperscript{41,46}. And, the nozzle simultaneously serves as an electrode, to which high electric field of 100-150 kV m\(^{-1}\) is applied. While the NWs solution is passing through the nozzle, under the influence of the strong electrostatic field, assembly of electrospinning fibers along the axial directions occurs. Very complex experimental setup and the requirement of high voltages are limits to the usefulness of this technique.

In the case of electric field assisted alignment technique, highly polarizable NWs polarize readily in the alternating electric field due to charge separation at the NWs surface and will experience the dielectrophoretic force that produces net movement to the NWs\textsuperscript{47}. However, NWs at various locations in between the electrodes experience an unequal dielectrophoretic force which results in poor alignment in the center regions and excellent
alignment in the regions close to the electrodes. However, the requirement of electrode fabrication for the NWs alignment and it is only applicable to the polarizable materials (e.g., CdSe, ZnO, Se NWs), etc., limits this technique to implementation for large-scale assembly processes.

Finally, the magnetic field assisted technique involves applying of an external magnetic field to directionally organize the magnetically active nanowires suspended in the solution\textsuperscript{30, 38, 41-43, 45, 48-50}. In the presence of an applied magnetic field, each NW will polarize into north and south poles and orient themselves parallel to the applied magnetic field direction\textsuperscript{30, 51}. The uniformity of the NWs alignment depends on the applied magnetic field strength. The added advantage of this technique is the assembled NWs forms chain-like structures by connecting each NW in a head to tail fashion, which gives a continuous network of NWs. These kinds of continuous networks are ideal for the best electrical performance. However, the main drawback of this technique is, it can only be applied to magnetically active NWs.

**Review on using nanowires self-assembly techniques to build functional nanodevices:**

The number of nanodevices, such as interconnects, FETs, bipolar transistors, light emitting diodes (LED) and logic gates have been assembled in a bottom-up approach from well-defined nanowires\textsuperscript{17, 29, 30}. Here discussed are few NWs self-assembly techniques used for assembling the NWs into functional devices.
Figure 1.5. Schematic diagram showing shear force induced self-assembly of Ge NWs into photolithography patterned channels (a), and optical micrographs of crossed arrays of Ge NWs shown at low (b) and high (c) magnifications. SEM images of single and multiple Ge NWs assembled into FET device (d)\textsuperscript{38,52}.

Figure 1.5 depicts the shear force assisted NWs self-assembly into crossed arrays and field effect transistor (FET) device. This process involves, mechanical movement of a donor substrate consisting of a lawn of vertically grown NWs, onto a photolithography...
patterned receiver substrate. This mechanically generated force can direct the NWs to align in a unidirectional manner, and the repetition of sliding in an orthogonal direction produces crossed arrays of NWs. The same shear force induced technique was used to fabricate single or multiple NWs based FET devices. This is an excellent wafer scale assembly technique; however, the limitation is this process requires vertically grown NWs which are usually prepared by complex CVD or template assisted processes\textsuperscript{53, 54}.

![Figure 1.6](image)

**Figure 1.6.** Schematic illustrating electric field assisted alignment of nanowires (a). Dense arrays of cadmium selenide nanowires assembled near the electrode (b). SEM images (c, d) zinc oxide nanowires assembled between electrodes. (e-g) SEM and optical microscope images of flexible selenium nanowires before and after (g) assembly\textsuperscript{38, 55}.

When a highly polarizable, but uncharged nanowire (e.g., CdSe, ZnO, Se NWs) is subjected to a reasonably strong ac electric field (> 1kV/cm), an induced dipole moment is created enabling the nanowire to respond to the electric field gradient\textsuperscript{55}. The NWs will reorient in the lowest energy configuration, such as parallel to the applied electric field. This process involves deposition of an NWs solution in between the electrodes and allowing it to dry in the presence of an applied electric field. Figure 1.6 shows optical and
SEM images of self-assembled nanowires near or in between the electrodes. However, this process is only applicable to polarizable NWs, and requirements such as microelectrode fabrication and high voltage limits this technique.

Figure 1.7. Schematic diagram illustrating the process of magnetic field assisted assembly of nickel (Ni) NW interconnect on Ni electrodes (a). SEM image of as-fabricated Ni NW based interconnect (b). Optical microscope images of Ni/Au/Ni NWs based interconnects.

Several efforts have been devoted to the magnetic field assisted self-assembly of magnetically active nanowires. The magnetic field assisted self-assembly involves directed deposition of well-aligned NWs on the electrodes. This process involves deposition of suspension of a magnetically active NWs solution on substrate and drying of the solution in the presence of applied magnetic field like the schematic shown in Figure 1.7. In the solution, the NWs align in the applied magnetic field direction, and the orientationally aligned NWs settles down on the electrodes. Self-assembly of single and multiple NWs based interconnects have been fabricated using pure Ni NWs,
Ni/Au/Ni NWs \(^{51}\), Au/Ni/Au \(^{50}\) nanowires. However, to date, the magnetic field assisted self-assembly technique is limited to assemble the single or few NWs into a nanodevice but, assembly of NWs on a large-scale has not yet been demonstrated.

This thesis work describes an investigation of magnetic field assisted self-assembly of high aspect ratio nickel coated copper nanowires (Cu/Ni NWs) into photolithography patterned channels on a large-scale area.

1.4. Thesis objectives and outlines

The overall objective of the thesis is to self-assemble one-dimensional (1D) nanocrystal-based interconnects. The central hypothesis of the project is “interconnects on IC surfaces can be self-assembled.” This hypothesis was tested by developing a magnetic field assisted technique to self-assemble the Cu/Ni NWs into photolithography patterned channels on the large-scale area on a carbon doped silicon dioxide (CDO) wafer, like the figure shown in Figure 1.8.

Figure 1.8. Schematic representing self-assembly of Cu/Ni NWs into photolithography patterned channels on CDO wafer.
Nanowires self-assembly was achieved via three specific aims. **Aim #1** is Cu NWs synthesis and electroless coating of nickel onto the Cu NWs surface. **Aim #2** is a preparation of the CDO surface to accept nickel coated copper nanowires (Cu/Ni NWs). **Aim #3** is magnetic field assisted self-assembly and deposition of NWs into photolithography patterned channels.

**Aim #1: Cu NWs synthesis and electroless coating of nickel onto the Cu NWs surface:**

In the fabrication of future bottom-up nanotechnology, the synthesis of copper nanowires (Cu NWs) has received considerable attention in recent years, and a number of methods have been successfully demonstrated which include template assisted processes\(^ {26, 27}\), chemical vapor depositions\(^ {53}\), hydrothermal/solvothermal routes\(^ {57}\), and solution-based methods\(^ {33, 35, 58}\). Among all the available synthesis routes, the solution-based approach is very attractive due to its process simplicity, low cost and easy to scale up for bulk scale production\(^ {58}\). However, a thorough understanding of the effect of reagent concentrations and synthesis conditions on the morphology, shape and aspect ratio of NWs are very critical. This study focuses on the effect of reducing agent concentration and synthesis temperature on the resulting surface morphology and aspect ratio of Cu NWs. The optimum reagent conditions to synthesize high aspect ratio Cu NWs with a smooth surface are to be established.

Although applications of Cu NWs have been demonstrated in a wide range of devices (e.g., transparent and flexible electrodes and conductive ink, etc.), oxidation of pure copper into copper oxides limit its reliable performance\(^ {59}\). Also, very limited success has been achieved in controlling the orientation and positioning of Cu NWs, which is very critical
for assembly of these NWs into complex functional devices. The above two challenges are addressed by coating a thin layer of oxidation resistant and ferromagnetic nickel onto the Cu NWs surface in an electroless coating approach. Nickel coating helped in protecting the copper core from oxidation and gave the magnetic response to Cu NWs. The orientation of Cu/Ni NWs is controlled by applying the uniform magnetic field to the NWs solution deposited onto the CDO surface.

**Aim #2: Preparation of CDO surface to accept nickel coated copper nanowires (Cu/Ni NWs).**

CDO is a low-$k$ dielectric material composed of Si, C, O, H elements and contains surface exposed hydrophilic Si-OH functional groups. A CDO material is chosen to demonstrate the NWs self-assembly process $^{60-62}$. Modern interconnects utilize this material to separate copper wiring in interconnects. It is well known that thiol (-SH) functional groups show strong interaction towards copper and nickel. Therefore, this work uses a thiol-functionalized CDO surface to attract and hold the Cu/Ni NWs strongly after deposition. Thiol functionalization of a CDO surface is achieved via a novel reaction scheme developed to selectively functionalize the surface exposed Si-OH functional groups to thiol-terminated alkyl groups, deploying the scheme shown in Scheme 1.1. The reaction process involves the conversion of hydrophilic Si-OH functional groups to its triflate ester (Si-O-Tf), and it is followed by reduction to hydrogen-terminated silicon (Si-H) using DIBAL-H reagent. Finally, hydrosilylation of Si-H using 2-propene thiol generates the thiol-functionalized CDO surface.
Scheme 1.1. Schematic representing selective functionalization of surface exposed Si-OH functional groups to thiol (-SH) terminated surface on CDO.

Aim #3: Magnetic field assisted self-assembly of NWs into photolithography patterned channels

Cu/Ni NWs self-assembly into a photolithography patterned channel is demonstrated by using the combination of photolithography, and magnetic field assisted NWs orientation controlling techniques as shown in Figure 1.8. Photolithography is used to define the circuit pattern (300 µm width and 4 mm length) on a thiol-functionalized CDO wafer. And, the orientation of Cu/Ni NWs is controlled by applying the uniform magnetic field to the Cu/Ni NWs dispersed aqueous solution. When the solution is dried, orientationally well aligned NWs will deposit on the photolithography patterned CDO substrate like the image shown in Figure 1.9. Finally, acetone solvent is used to dissolve the photoresist, and it simultaneously washes off the loosely bound NWs from the photoresist surface and ultimately leaves the self-assembled NWs channel on the CDO substrate. Electrical characterization is performed by measuring the resistance of the as-fabricated NWs based interconnect channel.
Figure 1.9. Schematic diagram showing a process flow of Cu/Ni NWs self-assembly into interconnect channels.

Thesis Outlines:

The following chapters provide the detailed information about results collected to support the above-discussed hypothesis. Chapter-2 discusses instrumentation, materials, and synthesis methods used in this study. The results of copper nanowires synthesis, an electroless coating of nickel onto Cu NWs surface and magnetic field assisted NWs orientation controlling technique details are discussed in Chapter-3. Selective functionalization of surface exposed hydrophilic Si-OH functional groups to thiol (-SH) terminated surface results are discussed in Chapter-4. And, Cu/Ni NWs self-assembly into photolithography patterned channels and electrical characterization results are discussed in Chapter-5. The thesis concludes with the discussion of summary, limitations, other applications and future scope of the developed techniques in Chapter-6.
CHAPTER TWO

INSTRUMENTATION, MATERIALS AND METHODS

2.1. INSTRUMENTATION

2.1.1. FTIR Spectroscopy

Structural and functional group information on Silica gel, SiO\textsubscript{x}, and CDO substrates were studied by Fourier transform infrared (FTIR) spectroscopy. The infrared spectrum is a nondestructive and powerful identification tool for functional groups in a wide range of samples (liquid, gel, solid substrates). FTIR data was collected on Thermoscientific, Nicolet\textsuperscript{TM} FTIR Spectrometer, in transmittance mode from 400 to 4000 cm\textsuperscript{-1} at a resolution of 4 cm\textsuperscript{-1}. For collecting FT-IR spectra of silica gel samples, a small amount of sample (~5 wt\%) was mixed with KBr salt and pressed into a thin pellet. CDO or SiO\textsubscript{x} wafers (dimension 1x2 cm) were directly placed in the path of IR beam and spectra collected in transmission mode.

2.1.2. NMR Spectroscopy

Nuclear magnetic resonance (NMR) spectroscopy was used in structural characterization and reaction progress study of organic model compounds. \textsuperscript{1}H and \textsuperscript{19}F-NMR spectra were collected on a 400 MHz Bruker NMR instrument at 256 scans. Reaction mixtures were dissolved in C\textsubscript{6}D\textsubscript{6} or DMSO-d\textsubscript{6} or CDCl\textsubscript{3} solvents. In all discussed reactions percentage yields were calculated from \textsuperscript{1}H or \textsuperscript{19}F-NMR spectra of crude mixtures.

2.1.3. X-ray Photoelectron Spectroscopy (XPS)
Surface chemical composition and bonding information were investigated using XPS. It is a qualitative and quantitative technique for analyzing surface chemistry of materials with extremely high sensitivity. XPS spectra were acquired on PHI Versa Probe II Scanning XPS equipped with Argon (Ar), C\textsubscript{60} sputtering gun and dual charge neutralizer. Survey spectrums were collected at 187.85 eV pass energy, 1 eV/step, 20 ms per step, 5 cycles and 5 sweeps. And, high-resolution spectra were collected at 23.5 eV pass energy, 0.1 eV/step, 20 ms per step, 5 cycles and 5 sweeps. PHI MultiPak data reduction program was used for the deconvolution and analysis of raw XPS data.

2.1.4. Contact angle measurements

The contact angle (CA) refers to the angle where a liquid droplet interacts with a flat and horizontal solid surface. It quantifies the wettability of the surface by the liquid droplet. A low contact angle (flat droplet) for a water drop indicates that the surface has high wettability (hydrophilic) and a high contact angle indicates the poor wettability (hydrophobic), see Figure 2.1b.

Water contact angle data was collected in a laboratory fabricated setup, like the schematic and digital photograph shown in Figure 2.1 a & c. For the data collection, a PC connected to a Kodak MDS 100 CCD camera was used to capture the droplet images on the surface of a wafer substrate. The contact angle data was analyzed by using Image-J software application (freeware available from NIH).
Figure 2.1. Schematic diagram depicting water contact angle measurement set up shown in a), and definition of the hydrophilic and hydrophobic surface shown in image b). Digital photograph in c), shows the laboratory fabricated CA measurement setup.

2.1.5. Electromagnet system

Unidirectional alignment of nickel coated copper nanowires (Cu/Ni NWs), experiments were carried out by applying a uniform magnetic field in electromagnet system, see Figure 2.2 a & b. GMW electromagnet system, Model-3470 with the dimensions of 45 and 30 mm of the pole diameter and pole gap respectively was used. Magnetic field strength can be varied by changing the current through magnet coil, Figure 2.2 c.
Figure 2.2. a) and b) showing digital photographs of electromagnet setup used for magnetic field assisted alignment of Cu/Ni NWs. Plot c) showing change in magnetic field strength as a function of applied current.

2.1.6. Scanning Electron Microscopy (SEM)

Morphology and dimensions of nanomaterials were investigated by scanning electron microscopy (SEM). FEI Sirion XL30 SEM equipped with field emission electron gun which operates from 200 V to 30 kV was used. Also, the microscope was equipped with Oxford energy-dispersive X-ray spectroscopy (EDS) and electron backscatter diffraction (EBSD) detectors. The EDS detector enabled the elemental composition and distribution analysis from photoelectrons generated from the specimen. Crystallographic orientation distribution was obtained from electron backscatter diffraction (EBSD) or transmission Kikuchi diffraction (TKD) patterning techniques as shown schematically in Figure 2.3. EBSD is a conventional technique which uses backscattered electrons, to study the orientation of ultrafine-grained metals and alloys, with grain sizes down to the
submicron scale. However, the spatial resolution of the EBSD technique, even in the advanced SEM equipped with FEG electron sources, is limited to 20 nm for dense materials, and 50 nm for lighter materials such as Mg and Al\textsuperscript{63,64}. Also, the resolution is inversely related to the sample tilt angle. Typically, EBSD sample holder is tilted to 70° with respect to the horizontal plane, which results in the poor (~3 times worse) spatial resolution. Hence, conventional EBSD in an SEM, cannot be applied as a routine characterization tool for nanostructured/nano-sized materials.

Figure 2.3. Schematic diagrams illustrating traditional EBSD a), and transmitted EBSD (t-EBSD or TKD) b), techniques used for crystallographic orientational mapping.

**Transmission Kikuchi Diffraction (TKD) pattern data collection:**

TKD is also known as transmitted electron backscatter diffraction (t-EBSD), and this offers significantly better spatial resolution than the conventional EBSD. TKD makes use of an EBSD detector and software application to capture and analyze the angular intensity
variation of transmitted electrons. Transmitted electron diffraction patterns originate from very close to the bottom surface of the sample. TKD working principles and data collection procedures are discussed in detail elsewhere\textsuperscript{64,65}.

The sample preparation for TKD studies involved drop casting of a dilute NWs solution (dispersed in ethanol) on a TEM grid. The TEM grid was mounted onto the custom-made sample holder (see Figure 2.4a). The TEM grid loaded sample holder was placed in the path of incident electron beam like the image shown in Figure 2.4b. Working distance, tilt angle and operating voltages had to be optimized to get high-resolution diffraction patterns.

![Figure 2.4](image)

**Figure 2.4.** Digital photographs showing custom-made sample holder for t-EBSD data collection of Cu NWs a), and SEM chamber highlighting (pointed with red arrows) the positions of the sample holder and EBSD detector.

2.1.7. Transmission Electron Microscopy (TEM)
High-resolution images of Cu NWs were collected in FEI Tecnai F-20 TEM/STEM equipped with Gatan dark and bright field detectors for STEM and Gatan ultra-scan CCD (2kx2k) camera. Sample preparation involved drop casting of a Cu NWs dilute (0.05 mg/mL) ethanol solution (20 µL) on the copper grid (200 mesh).

2.1.8. Optical Microscopy

Olympus BX51-P Optical microscopy was used to study the positioning and orientation of nanowires. It is equipped with a halogen lamp (100 Watts) and Canon high definition camera to capture the micrographs allowing a maximum lens magnification of 100X.

2.2. Materials and methods

2.2.1. Materials

Materials used for copper nanowires synthesis and nickel electroless coating:

Copper nitrate (Cu(NO$_3$)$_2$ 2.5 H$_2$O, ≥99.99%, 19004-19-4), Ethylenediamine (C$_2$H$_8$N$_2$, 99+ %, extra pure, 107-15-3), Hydrazine (N$_2$H$_4$, 35 wt % in H$_2$O, 302-01-2), Ethanol (C$_2$H$_5$OH, 99.99%, 64-17-5), Sodium hydroxide (NaOH, ≥97%, pellets, 1310-73-2), Ethylene Glycol (C$_2$H$_6$O$_2$, ≥99.8%, 107-21-1), and Nickel Nitrate (Ni(NO$_3$)$_2$ 6H$_2$O, 99.99%, 13478-00-7) were used as received from Sigma Aldrich, without further purification.

Materials used for surface functionalization of CDO wafer:

The materials tert-Butyldimethyl silanol (((CH$_3$)$_3$CSi(CH$_3$)$_2$OH, 99%, 18173-64-3), tert-butyl dimethyl silyltrifluoro methanesulfonate (CF$_3$SO$_3$Si(CH$_3$)$_2$C(CH$_3$)$_3$, 98%, 69739-34-0), trimethyl aluminum (Al(CH$_3$)$_3$, 97%, 75-24-1), trifluoro methane sulfonic anhydride ((CF$_3$SO$_2$)$_2$O, ≥99%, 358-23-6), ethoxytriphenylsilanol (C$_{20}$H$_{20}$OSi, ≥99%, 13478-00-7).
chloroform (CHCl$_3$, ≥99%, 67-66-3), anhydrous toluene (C$_6$H$_5$CH$_3$, 99.8%, 108-88-3) diisobutylaluminumhydride ((((CH$_3$)$_3$CHCH$_2$)$_2$AlH, 1191-15-7), 1-octadecene (C$_{18}$H$_{36}$, ≥95%, 112-88-9), octadecyltrichlorosilane (C$_{18}$H$_{38}$SiCl$_3$, ≥90%, 112-04-9) and 2-propene-1-thiol (CH$_2$=CHCH$_2$SH, ~60%, 870-23-5) were all reagent grade, purchased from Sigma-Aldrich and used as received. Sigma-Aldrich was the source of amorphous silica gel (high-purity grade (Davisil Grade 62), SiO$_2$, 60–200 mesh, Chromatography grade, Fisher Scientific, 7631-86-9). Intel Corporation provided CDO and SiO$_x$ coated silicon wafers.

Materials used for Photolithography patterning:

MEGAPOSIT™ SPR 220- 3.0 photoresist (Shipley), Sodium hydroxide (NaOH, ≥97%, pellets, 1310-73-2, Sigma Aldrich) and acetone (C$_3$H$_6$O, ≥99%, 67-64-1, Sigma Aldrich).

2.2.2. Copper nanowires synthesis method

Copper nanowires (Cu NWs) were synthesized in a solution based approach using ethylenediamine (EDA) as a growth directing agent and hydrazine as a reducing agent in an aqueous NaOH solution$^{33, 58}$. This work explored the effect of reducing agent concentration and synthesis temperature on dimensions and morphology of nanowires and established the optimized concentration of reagents for synthesizing surface morphology controlled, high-aspect-ratio (Length/diameter) Cu NWs. More synthesis details appear in Chapter-3.

2.2.3. Electroless coating of nickel on Cu NWs surface

Copper nanowires were coated with a thin layer (~20 nm) of nickel (Ni) in an electroless coating method. The nickel electroless coating procedure involves preparation
of a well-dispersed solution of Cu NWs, nickel nitrate (Ni(NO$_3$)$_2$) and reducing agent hydrazine in an ethylene glycol solution$^{50}$. After heating the as-prepared solution in the oil bath at 120 °C for 10 minutes produced floating of nickel coated Cu NWs at the air-liquid interface.

2.2.4. Unidirectional alignment of Cu/Ni NWs

Unidirectional alignment or self-assembly of NWs into interconnect channels was carried out by placing a plane or lithography patterned substrate in between the electromagnet poles presented in digital photographs shown in Figure 2.5 a & b. The Cu/Ni NWs aqueous suspension was deposited on top the substrate and allowed to evaporate under ambient conditions in the presence of magnetic field. Figure 2.5 c & d depict the schematic of Cu/Ni NWs arrangement after the solvent was allowed to dry in the absence and the presence of magnetic field respectively. In the absence of magnetic field, all the NWs oriented in random directions. Whereas in a high magnetic field, all the NWs oriented in the applied magnetic field direction.
Figure 2.5. Digital photographs of GMW electromagnet setup in top view a) and side view b). Schematic is shown in image c) and d) depicts an arrangement of Cu/Ni NWs in absence and presence of magnetic field.

Also, the uniform alignment of NWs on a given substrate requires a uniform magnetic field. Therefore, for all the NWs alignment experiments, substrates were placed in between the poles of the electromagnets. In comparison to the diameter of electromagnet poles, the substrates used were two-fold smaller in dimension.

2.2.5. Common synthetic procedure for selective functionalization of Si-OH moieties

The following common protocol was used to selectively functionalize the Si-OH functional groups on organic molecules (t-Butyldimethylsilanol and Triphenylsilanol), silica gel, SiOₓ and CDO wafer substrates.
**Scheme 2.1.** Schematic showing common reaction pathway for selective functionalization of Si-OH functional groups to Octadecyl and thiol-derivatized silicon.

The first step in the reaction scheme employed 0.1 M triflic anhydride ((CF$_3$SO$_2$)$_2$O) in anhydrous toluene at 50-60 °C for 48 hours to esterify the Si-OH functional group to yield silyl trifluoromethane sulfonate (Si-O-Tf, where Tf indicates the SO$_2$CF$_3$ group). In the second step, 0.1 M DIBAL-H (di-isobutylaluminum hydride, (CH$_3$)$_2$CHCH$_2$)$_2$AlH) reagent at 50-60 °C for 48 hours reduced the triflate intermediate (Si-O-(SO$_2$)CF$_3$) to silicon hydride. Because hydrogen-terminated silicon is unstable, Si-H was further treated with 1-octadecene (CH$_3$(CH$_2$)$_{15}$CH=CH$_2$) in the presence of UV light (254 nm) for 2 hours to produce Si-R (Si-C$_{18}$H$_{37}$). For thiol (-SH) functionalization, hydrogen-terminated silicon was treated with 2-propene thiol (CH$_2$=CH-CH$_2$-SH) under similar conditions. All reactions were carried out in dry N$_2$ within a glovebox. Between each set of reactions, silica gel, SiO$_x$ wafer, and CDO wafers were thoroughly washed with anhydrous toluene.

**2.2.6. RCA cleaning procedure for SiO$_x$ and CDO wafers**

SiO$_x$ and CDO coupons were cleaned with a standard RCA-1 solution. The cleaning protocol involved immersion of 1x2 cm coupons into RCA-1 solution (a mixture of 1:1:5
volume ratios of NH₄OH : H₂O₂ : DI water) at 70 °C for 10 minutes. RCA-1 solution is a strong oxidizer. It will effectively remove organic contamination from the wafer surface. After RCA cleaning, wafers were thoroughly washed in DI water, chloroform and, then anhydrous toluene solvent before proceeding to the functionalization reaction.

2.2.7. Photolithography

Photolithography or UV lithography refers to the process of using UV light to transfer the geometric pattern from a photomask to the thin film of photoresist. The schematic diagram shown in Figure 2.6, outlines the major steps in the photolithography process.

![Schematic diagram showing photolithography process flow.](image)

Figure 2.6. Schematic diagram showing photolithography process flow.

The procedure involved, spin coating of a thin film of positive tone photoresist (SPR 220 3.0, Shipley) on to the thiol-functionalized CDO substrate. The thickness of the as-
coated photoresist film was measured to be $3.0 \pm 0.2 \mu m$. The as-coated wafer was prebaked for 90 secs at 110 °C, which removed volatile organic solvents from the photoresist film. The circuit pattern (4mm long, 300 µm wide channel was used for self-assembly of NWs) was transferred into the photoresist film by exposing the photoresist coated wafer to UV light (365 nm) through a photomask in a benchtop projection mask aligner (Tamarack scientific Co., Model-162), see Figure 2.7. After UV exposure, the substrate was post-baked at 110 °C, to improve the solubility of UV exposed the photoresist. Finally, the post-baked substrate was developed in a dilute alkaline medium, which dissolved the UV exposed photoresist leaving the unexposed regions on the CDO substrate. The exact process conditions of the Photolithography patterning are summarized in table 2.1 below.

<table>
<thead>
<tr>
<th>Photolithography processing steps</th>
<th>Conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Photoresist</td>
<td>SPR 220 3.0 (Shipley), Positive tone resist</td>
</tr>
<tr>
<td>Spin coating speed</td>
<td>3500 RPM for 4 minutes</td>
</tr>
<tr>
<td>Pre-exposure bake</td>
<td>110 °C for 90 Sec</td>
</tr>
<tr>
<td>UV exposure duration</td>
<td>5 minutes</td>
</tr>
<tr>
<td>Post-exposure bake</td>
<td>110 °C for 90 Sec</td>
</tr>
<tr>
<td>Developer solution</td>
<td>0.1M NaOH aqueous solution</td>
</tr>
<tr>
<td>Development time</td>
<td>40 Sec</td>
</tr>
</tbody>
</table>

*Table 2.1. Photolithography process steps and conditions are presented here.*
Figure 2.7. Digital photograph showing projection mask UV aligner (Tamarack scientific Co., Model-162) used for photolithography patterning (capable of patterning resolvable feature size of 1 µm).

2.2.8. Electrical characterization of self-assembled interconnect channels

The self-assembly of nanowires into photolithography patterned channel protocol is discussed in detail in Chapter-5. Electrical characterization of the as assembled NWs channel is carried out by fabricating the silver contact pads (High purity silver paint, SPI supplies, CAS # 05002-AB, lot no # 1160331) on both ends of the channel and measured the resistance values using Keithley programmable multimeter (Model-617).
CHAPTER THREE
COPPER NANOWIRES SYNTHESIS AND MAGNETIC FIELD ASSISTED ALIGNMENT

In recent years, the synthesis of one dimensional (1D) nanostructures such as metallic NWs (Ag NWs, Cu NWs, Au NWs, and Pt NWs) or carbon nanotubes (CNT) and their assembly into functional devices have received special interest owing to its size and structure-dependent unique properties. Among all metallic nanowires, copper nanowires (Cu NWs) have attracted considerable attention lately due to their excellent electrical conductivity (only 7% less conductive than silver), thermal, mechanical properties, and cheap precursor cost (100 times cheaper than silver). In literature, various Cu NWs synthesis methods were discussed such as chemical vapor deposition (CVD) \(^{53}\), porous templates \(^{26, 54}\), hydrothermal/solvothermal \(^{57}\), and solution-based approaches \(^{33, 35, 58}\). However, in all the above methods, the ability to precisely control the morphology and dimensions of NWs was not possible. This thesis work adopted a solution based synthesis approach developed by Zeng et al. \(^{58}\) and Wiley et al. \(^{33, 66}\). This thesis studied the effects of reagent concentrations and synthesis temperature on morphology and aspect ratio controlled Cu NWs synthesis.

The bottom-up or hybrid approaches of NWs assembly into nanodevices requires simple, scalable and robust techniques to precisely control the orientation and positioning of NWs. For Cu NWs, very limited success has been achieved in controlling the orientation via electrochemical deposition of copper into a porous template \(^{26, 54}\) the thermal assisted growth of freestanding NWs on a copper substrate \(^{67}\) and vertically aligned NWs in a CVD
technique. The challenges in these techniques include high process complexity, poor yield, and the as-grown NWs stuck to the substrate (hard to manipulate the location); so Cu NWs are very challenging to transfer onto another desired substrate. To date, a simple and robust technique to control the orientation of Cu NWs has not been demonstrated. Here, for the first time, present work illustrates the magnetic field assisted unidirectional alignment of Cu NWs by coating them with a thin layer (~20 nm) of ferromagnetic nickel (Ni). Upon applying of the magnetic field, Cu/Ni NWs in the solution phase align in the applied magnetic field direction and as the solvent dries aligned NWs deposit on the substrate. In the subsequent sections, copper nanowires synthesis, an electroless coating of nickel onto Cu NWs surface (Cu/Ni NWs) and magnetic field assisted alignment of Cu/Ni NWs have been discussed in detail.

3.1. Copper nanowires synthesis

Copper nanowires were synthesized in a solution based approach, using copper nitrate (Cu(NO$_3$)$_2$ 2.5H$_2$O) as a precursor, ethylenediamine (EDA, C$_2$H$_4$(NH$_2$)$_2$) as a growth directing agent and hydrazine (NH$_2$-NH$_2$) as a reducing agent. The synthesis approach followed the method published in the literature$^{33, 58, 68, 69}$.

3.1.1. Cu NWs synthesis procedure: In brief, the procedure involved the mixing of a copper nitrate solution (0.1M in DI water, 2 mL) to a freshly prepared aqueous NaOH (15 M, 40 mL) solution. Ethylenediamine (0.1 M, 266 µL) and hydrazine (9.66×10$^{-3}$ M (35 µL) to 4.09×10$^{-3}$ M (15 µL)) reagents were added followed by a thorough mixing for 3 and 2 min respectively, after adding each reagent. Upon addition of hydrazine, the solution turned clear suggesting the reduction of Cu$^{2+}$ ions into Cu$^{+1}$. After 30 minutes, a reddish-brown
cake formed at the air-water interface leaving the clear solution at the bottom, see Figure 3.1c. NWs were collected and washed several times with DI water and ethanol in a centrifugation process, and finally stored in absolute ethanol solvent in a sealed glass vial.

![Image](image-url)

**Figure 3.1.** Digital images of Cu NWs reaction solution before adding hydrazine (a) and after adding hydrazine (b). After incubation for 30 minutes in hot water bath reddish-brown Cu NWs cake floating at the air-water interface.

The chemical reactions involved in Cu NWs synthesis are shown below\(^3\)\(^3\),\(^5\),\(^8\). Cupric (Cu\(^{2+}\)) ions react with hydroxyl (OH\(^-\)) ions and form cupric hydroxide complex ions (see eq-1). The reducing agent, hydrazine, supplies electrons that are required for the reduction of cupric ions to copper (Cu(0)). In the first step of the reduction process, cupric hydroxide (Cu\(^{2+}\)) complex ions are reduced to cuprous (Cu\(^+\)) hydroxyl complex ions, (see eq-3) (solution color turns from blue to clear, See Figure 3.1 a, b). And, in the second step, as cuprous ion complexes (Cu\(^+\)) are reduced to metallic copper (Cu\(^0\)), (see eq-4). The capping agent ethylenediamine (EDA), directs the copper atoms into unidirectional growth to form nanowire morphology.
Cu$^{2+}$ + 4OH$^-$ → Cu(OH)$_4^{2-}$ (blue solution)  

(1)

N$_2$H$_4$ + 4OH$^-$ → N$_2$(g) + 4H$_2$O + 4e$^-$  

(2)

2Cu(OH)$_4^{2-}$ + 2e$^-$ → 2Cu(OH)$_2^{-}$ (clear solution)  

(3)

2Cu(OH)$_2^{-}$ + 2e$^-$ → 2Cu(0) + 2OH$^-$  

(4)

The amine functional groups in EDA capping agent act as a ligand to form the coordinate covalent bond and interact with unsaturated surface atoms in the nanocrystals$^{70}$. The binding affinities of EDA are determined by the distinct atom geometries in various crystal planes. The capping agents on the nanocrystal act as a physical barrier to restrict the access of reagents in the capped facets and allowing the active growth process to occur in the uncapped regions$^{70}$.

**Copper nanowire growth mechanism:** The growth mechanism of copper nanowire in an EDA and hydrazine assisted method have been discussed in detail by Wiley et al.$^{35,71}$ In brief, the mechanism involves during the initial stages of reduction process Cu(OH)$_4^{2-}$ ions are reduced to Cu(OH)$_2^{-}$ ions and Cu$_2$O nanoparticles (the resultant solution appears as translucent, see Figure 3.1b). The capping agent EDA helps in preventing the precipitation of these Cu$_2$O nanoparticles by capping the surface. And, keeping these nanoparticles suspended in the solution are very crucial for the initiation of nanowire growth process. The EDA coated Cu$_2$O nanoparticles are further reduced by hydrazine to form irregular shaped polycrystalline metallic copper aggregates with the size ranging from 200 to 500 nm in diameter$^{70}$. The surface of these copper aggregates consists some of the pentagonal twinned secondary particles (~40 nm) which will serve as a seed to sprout nanowire$^{71}$. And,
during the nanowire growth process, EDA preferentially caps the higher surface energy 
\{100\} crystal facets and leaving the lower energy \{111\} crystal facets (nanowire tip). The 
Cu(OH)$_2^{−}$ ions are reduced to metallic copper (Cu(0)) by hydrazine at nanowire tip which 
possesses \{111\} crystal facet and promotes the anisotropic growth of nanowire 
morphology\textsuperscript{35}.

In this synthesis, NWs dimensions such as aspect ratio (length/diameter) and 
surface morphology depended critically on the reagent concentrations and synthesis 
conditions. Here, systematic studies of the effect of synthesis temperature and 
concentration of reducing agent on NWs dimensions and morphology were investigated.

### 3.1.2. Effect of synthesis temperature

The effect of synthesis temperature was studied at 60, 70, 80 and 90 °C respectively. 
Except for the temperature, all other reaction parameters were kept constant. Figure 3.2 
shows SEM images of Cu NWs synthesized at various temperatures, which indicates the 
decrease in aspect ratio as the synthesis temperature increase. Image-J software analyzed 
aspect ratios of Cu NWs synthesized at various temperatures are shown in Table 3.1. As 
the synthesis temperature increased to more than 90 °C, all the copper atoms agglomerated 
and formed micrometer-sized (200 nm to 3 µm) sized copper aggregates. It was suggested 
that the formation Cu-NWs is controlled by EDA which caps the side-wall surface of NWs. 
The potential reason for the formation of micron size aggregates at a higher temperature 
could be due to desorption of EDA from the NW surface, coupled with lower super-
saturation of Cu atoms at temperatures above 90 °C.
Figure 3.2. SEM images of copper nanowires synthesized at 60 °C (a), 70 °C (b), 80 °C (c), and 90 °C (d) respectively.

<table>
<thead>
<tr>
<th>Temp. (°C)</th>
<th>Average Length (µm)</th>
<th>Average Diameter (nm)</th>
<th>Aspect ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>60</td>
<td>32 ± 7</td>
<td>160 ± 30</td>
<td>200</td>
</tr>
<tr>
<td>70</td>
<td>25 ± 6</td>
<td>170 ± 50</td>
<td>147</td>
</tr>
<tr>
<td>75</td>
<td>15 ± 4</td>
<td>210 ± 60</td>
<td>71</td>
</tr>
<tr>
<td>80</td>
<td>7 ± 2</td>
<td>220 ± 30</td>
<td>32</td>
</tr>
</tbody>
</table>

Table 3.1. Average lengths, diameters and aspect ratios of copper nanowires synthesized at 60, 70, 80, and 90 °C respectively.
3.1.3. Effect of reducing agent concentration

Nanowire anisotropic growth and thickening mechanisms were discussed by Wiley et al., and highlight the importance of reducing agent (hydrazine) concentration\(^ {33, 35, 68} \). Similarly, the present investigation focused on the effect of hydrazine concentration on surface morphology and the aspect ratio of Cu NWs.

Figure 3.3. SEM images of Cu NWs synthesized using high concentration (35 µL) of reducing agent. Images shown at low a) and high b) magnification scales. TKD overlaid SEM image shown in c), and out-of-the-page (Z) direction crystallographic orientation is shown in d), highlight the thick and highly particle decorated NWs. e) crystal plane to color mapping scale.
In these studies, except for hydrazine, all other reagent concentrations and conditions were maintained constant. See experimental section (3.1.1.) for more information about the synthesis procedure. Figure 3.3a & b shows SEM images of Cu NWs synthesized using a high concentration (9.66×10⁻³ M, 35 µL) of reducing agent hydrazine. These images revealed that as-synthesized NWs featured low aspect ratio (L/D ~20) with rough surface morphology suggestive of Cu nanoparticle decoration on the NW surface. To characterize the nanoparticle decoration, this study examined the crystallographic orientational mapping of these particulates on the NW surface using SEM. Transmitted electron backscattered diffraction (t-EBSD) orientational mapping images shown in Figure 3.3 c & d displayed various crystallographic orientations of these (adsorbed) nanoparticles.

**Figure 3.4.** SEM images of copper nanowires synthesized using an intermediate concentration of hydrazine (8.28×10⁻³ M, 30 µL).

At an intermediate concentration of hydrazine (8.28×10⁻³ M, 30 µL), the synthesis yielded nanowires surface decorated with comparatively small particles or spikes; See
Figure 3.4. As the hydrazine concentration was further reduced to 5.73x10^{-3} M (21 µL), the resulting NW exhibited high aspect ratio (~250) and a particle-free and comparatively smooth surface, see SEM images in Figure 3.5a & b. The average length and the diameter of NWs was ~30±4 µm and 120 ± 20 nm, respectively. The t-EBSD pattern of the same batch of NWs, shown in Figure 3.5c & d, suggested a comparatively smooth and single crystalline surface. The dark regions in the orientational mapping images represented poor electron transmission.
Figure 3.5. SEM images of Cu NWs synthesized using a low concentration of reducing agent shown at low a) and high b) magnification scales. TKD c), and out-of-the-page (Z) direction crystallographic orientation in d), showing particle free thin Cu NWs. And, e) showing a color map for crystallographic planes. Histograms in f) and g) represents the length and diameter distributions of as-synthesized Cu NWs extracted from samples containing >200 NWs.
Low magnification TEM image of the same batch of NWs (hydrazine concentration 5.73x10^{-3} M, 21 µL) shown in Figure 3.6 revealed the thinner and particle free nature of the Cu NW surface. As the hydrazine concentration was further decreased to 4.09x10^{-3} M (15 µL), it resulted in a very poor yield of Cu-NWs with low aspect ratios. Also, a large portion of the reaction mixture remained in blue, which indicated that the hydrazine concentration was not sufficient to reduce all the copper ions.

![TEM image of Cu NW](image)

**Figure 3.6.** TEM image of Cu NW synthesized using 5.73x10^{-3}M (21 µL) concentration of hydrazine highlighting the particle-free nature of the surface.

The following three models could explain the formation of these particle or spike decoration on the NWs surface. First one is based on the screw dislocation-driven growth model discussed by Song Jin et al. In the reaction, copper ions (Cu^{2+}) form complexes with OH\(^-\) ions (Cu(OH)\(_4\)^{2-}) and ethylenediamine (EDA) ([Cu(en)](OH)\(_2\)) and presence of 

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these complex ions cause a decrease in the supersaturation of nanowire anisotropic growth. When an excess of the reducing agent results in a reduction of these copper complex ions into copper atoms all at once, a decrease in these complex ion concentrations causes an increase in the supersaturation at the tip of the nanowires. Hence, NWs stop growing in the axial direction and promotes the growth of nanowire in the lateral axis and may form particles with various crystal planes which are grown at the screw dislocation sites on the nanowire surface. Whereas at moderate hydrazine concentration, the presence of copper complex ions decreases the supersaturation and hence causes the growth of NWs in longitudinal axis leading to high aspect ratio NWs.

The second hypothesis is based on the EDA as a facet selective promoter of NW growth mechanism which is discussed by Wiley et al. Presence of excess hydrazine keeps the NW surface free of Cu oxide and promotes deposition of copper on the NW surface which results in thicker NWs. Whereas at a moderate concentration of hydrazine, the NW surface remains oxidized due to the presence of high OH- ions and prevents the growth in NW lateral axis direction and promotes the longitudinal axis ((100) facet) growth which ultimately results in high aspect ratio NWs.

The third hypothesis is an excess of reducing agent concentration results in a reduction of copper ion complexes to copper atoms all at once, and these highly concentrated copper atoms agglomerate to form multiply twinned nanoparticles. TEM images shown in Figure 3.7 a & b shows the presence of nanoparticles in the solution along with NWs, and perhaps during the nanowire growth, some of the particles may stick onto the nanowire surface leading to NW thickening. See Figure 3.7c. The mechanism reported
by Wiley et al. supports this hypotheses. The exact reason behind the particle decoration is not completely understood for now. Nevertheless, experimental studies show how the concentration of reducing agent affects the morphology and allows the determination of the optimized concentration to synthesize high aspect ratio and particle free nanowires.

![Figure 3.7](image)

**Figure 3.7.** Low magnification bright field a), and high magnification dark field b), TEM image showing the presence of nanoparticles. Bright field image c), showing nanoparticles sticking to the surface of the nanowire at multiple sites (pointed out with arrows).

3.2. Electroless plating of nickel (Ni) on Cu NWs surface

3.2.1. Nickel electroless plating procedure: Electroless plating of nickel onto the Cu NWs surface was conducted by a slightly modified version of the method discussed by Wiley et al. In brief, the procedure involved mixing of Ni(NO₃)₂ (0.1 M in DI water, 0.4 mL) and hydrazine (0.16 M (600 µL)) reagents with a 10 mg of Cu NWs (dry mass) suspension in ethylene glycol (20 mL). The above solution was thoroughly vortex mixed for 5 minutes and heated in an oil bath at 120 °C. After heating for 10 minutes, the solution turned clear and dark-colored nickel coated Cu NWs (Cu/Ni NWs) floated at the air-water interface due to trapped N₂ gas bubbles generated in the reaction, See Figure 3.8.
Figure 3.8. Digital photographs are showing nickel electroless plating mother solution (a mixture of Cu NWs / Ni(NO$_3$)$_2$ / Hydrazine in ethylene glycol solvent) a). After heating at 120 °C for 10 minutes, floating of dark-colored nickel coated Cu NWs at the air-water interface can be observed in b), and magnetic response of Cu/Ni NWs towards neodymium disc magnets is shown in c).

When a disc magnet was placed next to the freshly prepared Cu/Ni NWs solution, it attracted the mass of dark cupronickel nanowires, see Figure 3.8c. The nickel coating thickness on Cu NWs surface was studied by using EDS elemental mapping, See Figure 3.9, which revealed nickel layer on the CuNW surface of ~20 nm thickness. Also, the presence of oxygen at the interface of Cu and Ni indicated oxidized copper NW surface. But the nickel surface did not show significant oxidation implying metallic surface.

Electroless coating of nickel would be advantageous in a few aspects, such as protecting the core copper from further oxidation and for reducing the contact resistance between nanowires due to strong binding resulting from the ferromagnetic interaction. Furthermore, the ferromagnetic nature of the nickel allowed a directional control over the orientation of the NW by applying an external magnetic field.
Figure 3.9. SEM image of nickel coated copper nanowire (Cu/Ni NW) shown in a), and elemental mapping image showing the distribution of copper core and nickel coating on the surface b). In between the copper and nickel layers copper oxide (green color) can be observed.

3.3. Magnetic field assisted alignment

Figure 3.10. Digital photographs of electromagnet setup (GMW, Model-3470) used for controlling the orientation of Cu/Ni NWs are shown in a) and b). Digital photograph of well-aligned NWs in solution at 2500 G of applied magnetic field and after turning off the magnetic field to H = 0G, no rearrangement of NWs chains can be observed in d).
Controlling the orientation of nanowires is very critical for the assembly of NWs into complex nano or microelectronic devices. Here the magnetic field assisted alignment was explored to systematically control the orientation of Cu/Ni NWs on a centimeter scale area. The GMW electromagnet system provided the uniform magnetic field in between its poles (see Figure 3.10). The alignment procedure involved deposition of the Cu/Ni NWs aqueous solution on a given substrate as shown in Figure 3.10 to align NWs aligns in the applied magnetic field direction. Each NW effectively acted like a nanobar magnet in the field and formed NWs chains by connecting with each other in a head to tail fashion. And, surprisingly even after turning off the magnetic field (H = 0G) NWs chains remained in a well-organized manner, see Figure 3.10 d. This suggests owing to the ferromagnetic, (not paramagnetic) nature of nickel all the NWs remained magnetized. This led to a strong binding to each other even after turning off the magnetic field. These types of well-aligned nanowire arrays are very attractive for nanowire-based devices owing to its excellent percolation network formation, uniformity and comparatively lower NW-NW junction resistance\textsuperscript{45}.

Figure 3.11 shows the optical microscopic images of Cu/Ni NWs alignment in the presence and the absence of an applied magnetic field. When the NWs solution was allowed to dry under ambient conditions in the absence of a magnetic field, typically all the NWs arranged in random directions forming agglomerations due to Van der Waal forces of attractions. However, when the external magnetic field was applied, all the NWs acted like nanobar-magnets and aligned in the applied magnetic field direction, See Figure 3.11b. Furthermore, strings of NWs formed suggesting end-to-end attachment of NWs due
to favorable magnetic interaction (i.e., the attraction between north and south poles of the neighboring NWs). Following two steps of deposition and alignment in orthogonal directions produced an NW mesh or crossed arrays (Figure 3.11c) useful for fabricating flexible and transparent electrode preparation.$^{45,49}$

Systematic studies of alignment phenomenon were undertaken. In the absence of a magnetic field, all the NWs arranged in the random directions with an average angle of 45° with respect to the hypothesized axis and the resulting deposits showed agglomerations. Whereas, with an applied magnetic field (2500 G), the majority of the NWs aligned with an angle less than 10° with respect to the applied magnetic field direction (see angular distribution in Figure 3.11d). The NWs alignment angle distribution with respect to the applied magnetic field direction (in case of a magnetic field applied) and hypothesized axis (in case of no magnetic field applied) were determined by using Image-J application, see Appendix-B for a detailed protocol. Also, the average angle with respect to the magnetic field direction depended on the field strength. Therefore, the alignment distribution function was determined as a function of the magnetic field strength that varied from 0 G to 2500 G.
Figure 3.11. Optical microscope images of Cu/Ni NWs when they are allowed drying in the absence (a), and in the presence of applied magnetic field (2500 Gauss) (b). Image (c) shows the NWs mesh prepared by depositing a 2nd layer of NWs orthogonally. The histogram in (c) highlights the NWs orientation distribution with respect to the applied magnetic field direction.

3.3.1. Effect of magnetic field strength on NWs alignment order

As discussed in the previous sections, each nickel coated Cu NW (Cu/Ni NW) acted like a nano bar-magnet and responded to the externally applied magnetic field. Alignment strength of these bar magnets was explored by depositing 100 µL of Cu/Ni NWs solution (0.5 mg/mL) on the 1x1cm sized wafer substrate and the solvent was allowed to evaporate in the presence of a magnetic field with strength varying from 0 to 2500 G. See Figure 3.12. When there was no magnetic field (0 G), all the NWs aligned in the random directions
and formed agglomerations. As the magnetic field strength increased, Cu/Ni NWs began to form chains by connecting the NWs in an end-to-end fashion. As the field strength increased, NWs chain lengths tended to increase.

![Optical micrographs showing alignment order of Cu/Ni NWs at various applied magnetic field strength from 0 G to 2500 G.](image)

**Figure 3.12.** Optical micrographs showing alignment order of Cu/Ni NWs at various applied magnetic field strength from 0 G to 2500 G.

Excellent alignment with an average alignment angle of ~12° was achieved at 1500G. As the field strength is further increased to the 2500G, the average alignment angle decreased to ~8°. The observation suggested that at least a 1500 G magnetic field strength is required to get a reasonably good alignment. The average alignment angle tended to be further reduced and reached saturation at higher field strengths, See Figure 3.13. The distribution function for the corresponding NWs is shown below along with a summary of parameters in the accompanying table.
**Figure 3.13.** Plot showing average NWs alignment angle with respective to the applied magnetic field direction, at various field strength from 0 G to 2500 G. The inset image describes the alignment angle definition.

**Figure 3.14.** Gaussian peak fitting plots are showing NWs alignment distribution with respect to the applied magnetic field strength from 0 to 2500 Gauss.

Orientation distribution histograms of Cu/Ni NWs measured as a function of magnetic field strengths were fitted to a Gaussian equation below.
\[ N(\theta) = Ae^{\left(\frac{(\theta-\langle\theta\rangle)^2}{\Delta\theta}\right)} + B \]

A and B are a normalization constant and a constant background fitting parameter. \( N(\theta) \) is a number of nanowires oriented at angle \( \theta \) in a bin size of 0.2 radians. \( <\theta> \) and \( \Delta\theta \) are the average absolute value of the NW orientational angle and the width of the distribution respectively. The best-fit parameters and respective curve fitting uncertainties appear in Table 3.2.

<table>
<thead>
<tr>
<th>Field Strength (gauss)</th>
<th>( \theta ) (radian)</th>
<th>( \Delta\theta ) (radian)</th>
</tr>
</thead>
<tbody>
<tr>
<td>000</td>
<td>0.7±0.1</td>
<td>0.4±0.2</td>
</tr>
<tr>
<td>500</td>
<td>0.75±0.05</td>
<td>0.25±0.08</td>
</tr>
<tr>
<td>1000</td>
<td>0.54±0.03</td>
<td>0.21±0.03</td>
</tr>
<tr>
<td>1500</td>
<td>0.23±0.07</td>
<td>0.41±0.08</td>
</tr>
<tr>
<td>2000</td>
<td>0.13±0.09</td>
<td>0.37±0.09</td>
</tr>
<tr>
<td>2500</td>
<td>0.3±0.7</td>
<td>0.5±0.3</td>
</tr>
</tbody>
</table>

**Table 3.2:** List of field dependent parameters.

### 3.4. Summary of copper nanowires synthesis and magnetic field assisted alignment

Control over nanowire surface roughness, aspect ratio, and alignment are very important for their device application. In this thesis research, a solution based synthesis method was optimized with respect to the reducing agent concentration to synthesize the high aspect ratio NWs with a smooth surface. Also, the effect of synthesis temperature on the aspect ratio of nanowires was investigated. As the synthesis temperature increased, the aspect ratio of NWs decreased. In the case of controlling the alignment of Cu NWs, the previous researcher had reported very limited success via CVD, electrochemical deposition in a porous template and thermal assisted growth of vertically aligned NWs. For the first-time simple magnetic field based alignment of NWs on macroscopic wafer-scale was realized.
by coating Cu NWs with a thin layer of ferromagnetic nickel. The nickel coating would be further advantageous in protecting the copper core from oxidation.
CHAPTER FOUR
SURFACE DERIVATIZATION OF CARBON DOPED SILICON DIOXIDE (CDO) WAFER

Carbon doped silicon dioxide (CDO), a highly nanoporous low-\( k \) dielectric, was chosen as a substrate for the fabrication of nanowires based interconnects. The self-assembly of NWs into interconnect structures on CDO wafer required surface preparation of CDO to attract and strongly hold the nanowires after deposition. In this work, thiol (-SH) functionalization of the CDO surface and strong interaction of thiol to copper and nickel was explored. Thiol functionalization of the CDO surface was achieved via a novel approach developed to selectively functionalize the surface exposed silicon hydroxide (Si-OH) functional groups\(^{60} \). The hydrophilic Si-OH functional groups are generated during various steps in chip integration processing and cause undesirable moisture uptake which adversely increases the dielectric constant (\( k \)) of CDO. We addressed this issue by selectively derivatizing the surface exposed hydrophilic Si-OH functional groups to highly hydrophobic long alkyl chain (Si-\( R \), \( R = \) octadecyl chain). The dense layer of octadecyl chains then acts as a moisture repellent thin film. This chapter provides the details of CDO film derivatization by octadecyl and thiol chains along with CDO surface characterization by XPS, IR, capacitance-voltage and contact angle techniques.

4.1. Introduction

Some of the biggest achievements in the past half-century involved the development of the microprocessor industry. To address a large number of transistors integrated into microprocessors, a sophisticated multilayer wiring, the so-called, interconnect architecture,
was essential. In the beginning of the Integrated Circuit (IC) era, aluminum alloys, tungsten, and SiO₂ insulators were the materials of choice for interconnects systems. In the early 1990s, researchers had realized the concept of improving the IC performance by increasing the transistor count and making changes to the traditional wiring and insulator materials. As the ICs feature sizes were downscaling, Al/W/SiO₂ interconnect system became the limiting factor due to high power consumption and low electromigration resistance. Owing to its advantageous high conductivity, abundance, and ductility characteristics, copper (Cu) as a interconnect material was extensively explored for many years. In 1997, IBM made a remarkable contribution to IC industry, by developing chips with copper interconnects, that could significantly enhance the chip performance. Multilayered copper interconnect chip developed by IBM in 1997, is shown in Figure 4.1.

In comparison to aluminum, copper offers low electrical resistivity and high electromigration resistance. Using low resistivity interconnect material like copper decreases the RC signal delay and ultimately increases the IC speed.
**Figure 4.1.** Image of the multilayered integrated circuit (IC) using Cu/W/ SiO$_2$ dielectric materials, developed by IBM in 1997.

As the transistor count and interconnect density increase, the RC signal delay, and power dissipation also increase because of increased capacitive coupling between densely packed metal lines. Since the RC signal delay and power consumption all are directly related to dielectric constant ($k$) of the inter-metal dielectric material, many efforts have been devoted to replacing the standard silicon dioxide ($k = 3.9$ to $4.2$) with an ultra-low $k$ material\textsuperscript{76}. The Dielectric constant ($k$) is defined as the ratio between permittivity of a substance to the permittivity of vacuum as shown in the below equation. In Table, 4.1 $k$ values of common materials are shown.

$$\text{Dielectric constant } (k) = \frac{\text{Permitivity of substance}}{\text{Permitivity of vacuum}}$$

Due to low $k$ value, many organic compounds are considered as a good low $k$ dielectric material, but organics have limitations associated with poor thermal and mechanical stability. Furthermore, inorganic materials show good mechanical and thermal...
stabilities, but they possess higher $k$ and more serious moisture sensitivity, requiring an additional liner and capping layers\textsuperscript{77,78}. Hybrid films, with organic and inorganic materials, have been a promising approach for low-$k$ materials with stable properties. Among hybrid type materials, carbon doped silicon oxide (CDO) material has attracted more attention owing to its low-$k$, and compatible mechanical and thermal properties\textsuperscript{3,79-82}. Hence, in modern ultra-large-scale integrated circuit (ULSI) devices CDO has been a choice as a low-$k$ insulator.

<table>
<thead>
<tr>
<th>Substance</th>
<th>$k$ - value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vacuum</td>
<td>1</td>
</tr>
<tr>
<td>Polyethylene</td>
<td>2.25</td>
</tr>
<tr>
<td>Carbon Doped Silicon Oxide (CDO)</td>
<td>2.2 to 3.0</td>
</tr>
<tr>
<td>SiO2</td>
<td>3.9</td>
</tr>
<tr>
<td>Silicon</td>
<td>11.68</td>
</tr>
<tr>
<td>Water</td>
<td>~ 80</td>
</tr>
</tbody>
</table>

Table 4.1. Dielectric constant ($k$) values of various substances.

4.2. CDO as a low-$k$ material

Over the last few years, the classic SiO$_2$ insulator used to separate metallic interconnect lines are being replaced by materials of lower $k$. Lower $k$ improves signal propagation delay. The lower the $k$ value shorter the propagation delay. Strategies to realize lower $k$ materials include (1) Introduction of carbon in SiO$_x$ network structure, by using so-called CDO materials (reviewed in this work). CDO film preparation and chemical structures etc. are discussed in detail in the following sections. (2) Incorporating porosity (note: air has lowest $k$ \~1). Porosity is inversely related to $k$-value; as the porosity
increases, $k$ values decreases. In the as deposited CDO film, nanopores can be introduced by plasma curing processes. The plasma burns off organic moieties and leaves nanopores in the film.

4.3. Review on CDO film preparation and chemical structure

Carbon doped silicon oxide (CDO), composed of Si, C, O, and H (SiCOH), is deposited on Si (100) surface by plasma enhanced chemical vapor deposition (PECVD) from mixtures of silicon and an organic precursor. In a PECVD reactor, the plasma is sustained by 13.56 MHz rf power. The silicon and organic precursors are carried by He gas into the plasma chamber. SiCOH film preparation employs various silicon precursors such as tetramethyldisiloxane, hexamethyldisiloxane, tetramethyltetrasiloxane. In the presence of plasma, precursor molecules break down into fragments and deposit on the Si wafer. The deposited films are annealed at 400 °C. The annealing step removes thermally unstable organic fragments and creates porosity in the material which enables the adjustment of the $k$ between 2.8 to 2.05. Post thermal treatment also causes extensive cross-linking of Si, C, and O to form Si-Si, Si-O-Si, Si-CH₂-Si and Si-CH₂-O-Si moieties. The annealing process depending on the duration and temperature causes the loss of $-\text{CH}_n$ groups and results in the formation of $-\text{Si-C}$- linkages and nanopores in the film.

An alternative method to thermal annealing is plasma curing which can reduce $k$ ($k = 2.0$ to 2.2). The plasma curing process creates nanopores and improves the crosslinking which helps in improving the mechanical strength of the film. The plasma curing process is an industry preferred method, owing to its advantageous lower operating temperature,
uniformity, cost-effectiveness, shorter processing time, and produces improved mechanical strength of the film.

4.4. Review on CDO damage and repairing methodologies

4.4.1. Review on CDO damage by plasma processing

Interconnects in modern ICs integrate multilayers of copper wiring and insulating low k materials (See Figure 4.1). They are fabricated using a combination of optical lithography and copper electroplating in a process known as the Dual Damascene (DD) method\textsuperscript{37}. In optical lithography, a photoresist film made up of organic polymers, and photosensitive compounds serve as a patternable and sacrificial stencil. An optical image of the desired circuit pattern of trenches and vias are projected on the photoresist film coated on top of the CDO layer. Photochemical reactions in the light exposed regions alter the solubility of the photoresist film regio-specifically. In the developer solution, the film undergoes preferential dissolution (positive tone) or dissolution inhibition (negative tone) to expose the underlying CDO transferring an image of the desired circuit pattern on the chip.
Figure 4.2. a) Chemical structure of porous cage networked structure of as-prepared CDO film from TMCTS precursor82, and b) chemical structure of plasma damaged film (R, represents an extension of Si-O-Si backbone structure)85.

This pattern is etched into the exposed CDO through plasma etching which opens up the electrical contacts in the lower layer. Subsequently, metallic copper is selectively and electrochemically plated in the etched vias/trenches, while avoiding its deposition on top of the undissolved electrically insulating photoresist film. After electroplating, the remaining sacrificial photoresist film is subjected to oxygen plasma ashing to remove organics as CO/CO₂ and water as ultimate byproducts86. Furthermore, oxygen radicals generated during the plasma processing react with the functional groups of the CDO film, breaking Si-CH₃, Si-H bonds, and creates dangling bonds of silicon. These dangling bonds can easily react with hydroxyl ions in the plasma environment to form Si-OH bonds (see Figure 4.2))85. The resulting increase in the number of hydrophilic Si-OH groups in the CDO films makes the film more polar, hydrophilic and allows moisture uptake (water k = ~80). Absorbed moisture increases the k value (note the dielectric constant of water is 80!)

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and leakage current (due to protonic conduction) thereby compromising the quality of the CDO$^{87}$.

### 4.4.2. Review on CDO repairing methodologies

Several approaches have been investigated to minimize the damage of porous CDO ultra-low $k$ film. One method involves protecting the CDO film from oxygen plasma damage by depositing a thin inert SiN$_x$ passivation layer. The SiN$_x$ film is generated from a thin layer of hybrid-organic-siloxane-polymer (HOSP) subjected to NH$_3$ plasma treatment$^{88}$. This inert layer can enhance the resistance of the HOSP film to moisture uptake and O$_2$ plasma attack. Helium (He) plasma treatment as a post CDO deposition has also been investigated to reduce the O$_2$ plasma damage on the CDO films$^{89}$. Treating the O$_2$ plasma damaged CDO with a hydrogen (H$_2$) plasma to repair the damage by reducing silanol to silicon hydride has also been explored$^{89}$. However, CDO repairs by He, NH$_3$ and H$_2$ plasma treatments cause a loss of CDO film thickness (by reducing porosity) and irreversibly modify the network of the SiCOH chemical structure. Etching of plasma damaged low $k$ CDO layer by dilute HF solution has been investigated, but the concentration of HF solution and duration of treatments are very critical to prevent further etching of the undamaged CDO layer$^{90}$.

Another approach to preventing moisture uptake in the damaged CDO is to convert surface silicon hydroxyl (Si-OH) functional groups to hydrophobic Si-O-Si(CH$_3$)$_3$ moieties by treating with hexamethyldisilazane (HDMS)$^{62}$. However, the resulting single carbon (-CH$_3$, Methyl) termination is too thin to prevent the further attack from oxygen and moisture. Hence, it is necessary to find a way to selectively derivatize hydrophilic
functional groups (Si-OH) to hydrophobic, without disturbing the backbone structure of the CDO film.

4.5. Results of developed pathway for selective derivatization of silanol

4.5.1. Developed scheme for surface derivatization

A gentle and spatially selective approach for the modification of surface exposed, hydrophilic silicon hydroxyl (Si-OH) functional groups to hydrophobic hydrogen-terminated (Si-H) is developed. The stability and surface hydrophobicity was further enhanced by long alkyl chain derivatization of the silicon hydride (Si-R, R= C_{18}H_{37}) as shown in Scheme 4.3. Compared to methyl groups, long chain alkyl groups form a thicker and more compact hydrophobic film that blocks moisture uptake effectively. The significant advantage of this approach is that the selective derivatization occurs only on the exposed or surface silicon hydroxyl (Si-OH) groups of CDO film without damaging the rest of the nano-porous cage structure. Additionally, silicon hydride formed during the intermediate stage may be further derivatized with different reactants to provide new surface functionalities. For example, it provides a convenient route for the insertion of additional functional groups such as terminal thiol (-SH) or amine (-NH\textsubscript{2}) alkene structure.

Scheme 4.1. Schematic of the new reaction pathway for hydrogen termination and octadecane (Si-R, R= C_{18}H_{37}) derivatization.
The viability of this approach has been established methodically by preliminary studies of derivatization of silanol groups in organic molecules, silica gel, SiOₓ wafer and CDO as shown in Figure 4.2 below.

![Scheme 4.2](image)

**Scheme 4.2.** Schematics of various compounds considered for testing surface derivatization reaction, a) tert-Butyldimethylsilanol b) silica gel, and c) SiOₓ wafer and d) nano porous carbon doped silicon oxide (CDO) wafer.

**Common synthetic procedure:** The first step in the reaction scheme employed 0.1 M triflic anhydride ((CF₃SO₂)₂O) in anhydrous toluene at 50-60 °C for 48 hours to esterify the Si-OH functional group to yield silyltrifluoromethane sulfonate (Si-O-Tf, where Tf indicates the SO₂CF₃ group). In the second step, 0.1 M DIBAL-H (di-isobutylaluminum hydride, (CH₃)₂CHCH₂₂AlH) reagent at 50-60 °C for 48 hours reduced the triflate intermediate (Si-O-(SO₂)CF₃) to silicon hydride. Because hydrogen-terminated silicon is unstable, Si-H was further treated with 1-octadecene (CH₃(CH₂)₁₅CH=CH₂) in the presence of UV light (254 nm) for 2 hours to produce Si-R (Si-C₁₈H₃₇). All reactions were carried out in dry N₂ within a glovebox. Between each set of reactions, silica gel, SiOₓ wafer, and CDO wafers were thoroughly washed with anhydrous toluene.

**4.5.2. Results on organic molecules**
Scheme 4.3. Schemes of a) Synthesis of TBDMS Triflate and tert-Butyldimethylsilane, b) Synthesis of tert-Butyltrimethylsilane c) Synthesis of Triphenylsilane.

a) tert-Butyldimethylsilanol

tert-Butyldimethylsilanol \(([\text{CH}_3]_3\text{CSi(\text{CH}_3)_2\text{OH}}]\) was used as a control molecule for the reaction scheme shown in Figure 4.3a. Reaction kinetics were studied by $^1$H-NMR spectroscopy and reaction products were confirmed by $^{19}$F-NMR, FTIR and Gas Chromatography-Mass Spectrometry (GC-MS). Transformation of the molecule to triflate and hydride molecules can be readily tracked by NMR, IR, and GC-MS techniques enabling optimization of the reaction conditions and yield. The kinetics of the triflation reaction was monitored by $^1$H-NMR of the reaction mixture over 72 hours, within 48 hours the reaction goes to completion (see Figure 4.3). The product of the reaction was further confirmed by parent ion peak at 265.0 g.mol$^{-1}$ in the mass spectrum.
In step 2, the reduction of TBDMS-OTf to tert-Butyldimethylsilane was achieved by treating the TBDMS-OTf with DIBAL-H reagent; the corresponding $^1$H-NMR spectrum was shown in Figure 4.4. After 24 hours of reaction, a new multiplet assigned to Si-H at δ 3.64 (m, 1H) suggest the formation of hydrogen-terminated product, completes the reaction within 48 hours. In $^{19}$F-NMR spectra, shown in Figure 4.5, fluorine peak corresponding to triflate become negligible after 48 hours, confirm the reaction completion. The observed FTIR peak at 2105.7 cm$^{-1}$ due to the formation of Si-H bond and a GC-MS parent ion peak at 116.1 g/mol$^{-1}$ further confirmed the formation of the expected silane.

**Synthesis of tert-Butyldimethylsilyl trifluoromethanesulfonate (TBDMS-OTf):** 1mmol of tert-Butyldimethylsilanol and 2mmol of triflicanhydride reagents were dissolved in 1ml of anhydrous toluene solvent and mixed at 50 to 60 °C for up to 72 hours. Percentage yields were calculated from $^1$H-NMR of the crude reaction mixture, suggests a 30, 60, 98 and 100% product yield for 2, 24, 48 and 72 hours respectively. $^1$H-NMR (C$_6$D$_6$, 400 MHz) δ 0.60 (s, 9H) and δ -0.00 (s, 6H); $^{19}$F-NMR (CDCl$_3$, 400MHz) δ -76.84 (s, 3F); GCMS calculated for CF$_3$SO$_3$Si(CH$_3$)$_2$C(CH$_3$)$_3$ 264.34, found 265.0; IR (KBr) $\nu_{max}$ 3027, 2926, 2857, 1228, 1193, 1039 cm$^{-1}$. 
Figure 4.3. $^1$H-NMR ($C_6D_6$, 400 MHz) reaction progress study of TBDMS Triflate synthesis at 2, 24, 48 and 72 hours respectively. Increase in peak intensities at $\delta$ 0.60 (s, 9H) and $\delta$ -0.00 (s, 6H) corresponding to TBDMS Triflate can be observed.

**Synthesis of tert-Butyldimethylsilane:** 1mmol of tert-Butyldimethylsilyl trifluoromethane sulfonate and 2 mmol of DIBAL-H reagents were dissolved in 1ml of anhydrous toluene solvent and mixed at 50 to 60 °C for up to 72 hours. Percent yields were calculated from $^1$H-NMR of the crude reaction mixture, suggests 0, 61, 89 and 91% product conversion after 2, 24, 48 and 72 hours respectively. $^1$H-NMR ($CDCl_3$, 400 MHz) $\delta$ 3.64 (m, 1H), $\delta$ 0.88 (s, 9H), $\delta$ 0.02 (d, $J$ = 3.1 Hz, 6H); GCMS calculated for HSi(CH$_3$)$_2$C(CH$_3$)$_3$ 116.28, found 116.10; IR (KBr) $\nu_{\max}$ 3027, 2926, 2857, 2105 cm$^{-1}$. 

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Figure 4.4. $^1$H-NMR (CDCl$_3$, 400 MHz) reaction progress study of tert-Butyldimethylsilane synthesis at 2, 24, 48 and 72 hours respectively. Peaks in the regions of $\delta$ 0.00 and $\delta$ 3.64 ppm were magnified and shown on the right and left-hand side of the original spectra. The appearance of multiplet at 3.64 (m, 1H) after 24 hours of reaction suggests the formation of the hydrogen-terminated product. The decrease in the peak intensities of reactant and increase in peak intensities of the product can be observed here.
Figure 4.5. $^{19}$F-NMR (CDCl$_3$, 400 MHz) reaction progress study of tert-Butyldimethylsilane synthesis at 2, 24 and 48 hours respectively. Fluorine peak intensity at $\delta$ -76.84ppm corresponding to the reactant TBDMS Triflate, decreases from 2 hours to 24 hours and becomes negligible after 48 hours of reaction can be observed.

b) Synthesis of tert-Butyltrimethylsilane.

It was possible to methyl terminate the TBDMS-OTf by treating it with Trimethylaluminum at 25 °C in 24 hours. In $^1$H-NMR spectra shown in Figure 4.6, the presence of 1:1 ratio singlet peaks corresponding to methyl groups of tert-Butyl and silicon suggest the methyl termination.

Synthesis of tert-Butyltrimethylsilane: 1 mmol of tert-Butyldimethyltrifluoro methane sulfonate and 2 mmol of trimethylaluminum reagents were dissolved in anhydrous toluene solvent and mixed at 25 °C for 24 hours. $^1$H-NMR (C$_6$D$_6$, 400 MHz) $\delta$ 0.93 (s, 9H), $\delta$ 0.01 (s, 9H); GCMS calculated for (CH$_3$)$_3$SiC(CH$_3$)$_3$ 130.30, found 130.1.
Figure 4.6. $^1$H-NMR (DMSO-$d_6$) reaction progress study of tert-Butyltrimethylsilane synthesis at 2, 24 hours of reaction. A gradual increase in peak intensities at $\delta$ 0.93 (s, 9H), $\delta$ 0.01 (s, 9H) with 1:1 ratio corresponding to methyl groups of the product i.e., tert-butyl and 3 methyl groups on silicon can be observed.

Thus, DIBAL-H and Trimethylaluminum reagents successfully converted the triflate ester of silanol to hydride and methyl groups. In addition to tert-Butyldimethysilanol, the hydrogen termination on Ethoxytriphenylsilanol was confirmed.

c) Ethoxytriphenylsilane

In addition to tert-Butyldimethysilanol, the reaction pathway on Ethoxytriphenylsilane was tested as shown in Scheme 4.3c. $^1$H-NMR data of the expected silane suggest the hydrogen termination of silicon by DIBAL-H reduction (see Figure 4.7).
Synthesis of Triphenylsilane: 1 mmol of ethoxytriphenylsilane and 2 mmol of DIBAL-H reagent were dissolved in anhydrous toluene solvent and mixed at 50-60 °C for 48 hours. Percentage yield calculated from $^1$H-NMR of the crude reaction mixture was 48%. $^1$H-NMR (DMSO-d$_6$, 400 MHz) $\delta$ 7.58 (m, 15H), $\delta$ 5.7 (s, 1H); IR (KBr) $\nu_{\text{max}}$ 3067, 2119, 1427, 1113, 802, 730, 697 cm$^{-1}$.

Figure 4.7. $^1$H-NMR (DMSO–D$_6$) of triphenylsilane. Peaks at $\delta$ 5.39 (s, 1H), $\delta$ 7.5 (m, 15H) were assigned to hydrogen-terminated silicon (Si-H) and protons of phenyl groups respectively. Peaks at $\delta$ 2.3 (S, 1H) and $\delta$ 3.52 (s, 9H) corresponding to toluene solvent and DIBAL-H reagent.

4.5.3. Results on Silica gel reaction

The reaction scheme discussed above can be readily adapted for modification of silica gel used in classic chromatography, pharmaceutical, cosmetic and in many other applications involving silanol group. In recent years, silica aerogel has been widely used in a variety of industries due to its exceptional properties such as low thermal conductivity,
high porosity, high optical transmission, high specific surface area, low dielectric constant, low refractive index and low sound velocity; however its moisture sensitivity has limited its practical use\textsuperscript{93, 94}.

In the literature, various derivatization strategies have been studied to alter the hydrophobicity of silica gel; these include methyltrimethoxysilane (MTMS), hexamethyldisilazane (HMDZ), dimethylchlorosilane (DMCS), trimethylethoxysilane (TMES), and hexadecyltrimethoxysilane (HDTMS)\textsuperscript{95}. These prevailing methods render the surface hydrophobic by replacing hydrogen from hydroxyl (SiO-H) functional group\textsuperscript{91, 96-104} which leads to the formation of siloxy groups (SiO-R). But the method developed here leads to the selective replacement of hydroxyl (-OH) functional groups with a nonpolar Si-C group bearing long alkyl chains (Si-R, R= C\textsubscript{18}H\textsubscript{37} or other long chain hydrocarbon) which resulted in less polar and a highly hydrophobic surface with lower dielectric constant.

Having established the feasibility of the reaction scheme on molecular species, its potential for derivatizing silica gel was explored. Silica gel, structurally similar to CDO, with its highly porous structure and a large number of air exposed hydroxyl functional groups provided a useful control to test the reaction scheme on a heterogeneous substrate\textsuperscript{105}. 

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Figure 4.8. Experimental setup of silica gel reaction. After triflation reaction silica gel turned to black and after DIBAL-H reaction turned to golden brown color.

The experiments used chromatography-grade silica gel, with particles ranging from 75 to 250 µm, $1.15 \times 10^{-6}$ m³/g specific volume, and surface area of 300 m²/g, with a pore size of approximately 15 nm. Before derivatization, the silica gel was dried by heating at 400°C for 4 hours and allowed to cool under modest vacuum (100 mTorr). Silica gel was then immersed in the reaction solution as shown in Figure 4.8. Reaction progress on the silica gel was studied by FTIR spectroscopy as shown in Figure 4.9. A broad peak at 3439 cm⁻¹ was ascribed to the superposition of stretching vibrational modes of physically adsorbed water molecules and silicon hydroxyl (Si-OH) functional group. Peaks at 1626 cm⁻¹, 1480 cm⁻¹, were assigned to the bending vibration of the water molecule and silanol respectively. 1102 cm⁻¹ and 969 cm⁻¹ peaks reflected the Si-O-Si stretch vibration modes of silica gel⁹¹, ⁹⁶, ⁹⁸, ¹⁰¹-¹⁰³. Triflation reaction with triflic anhydride resulted in new peaks (≡Si-O-Tf) at 1033 cm⁻¹, and 1257 cm⁻¹ corresponding to S=O asymmetric, symmetric stretches and an 1177 cm⁻¹ peak ascribed to C-F stretching corresponding to formation of (≡Si-O-(SO₂)CF₃) groups respectively. It should be noted that the trifilation reaction did
not result in complete removal of the broad OH peak since the silica gel has OH groups not accessible to the chemical reagent.

![FTIR spectra of silica gel](image)

**Figure 4.9.** FTIR spectra of silica gel in various stages of the reaction scheme. Pure Si gel (black), after treatment with triflic anhydride (blue), DIBAL-H reduced (pink) & after 1-octadecene treatment (red).

Further attempts to estimate the percent conversion of the surface silicon hydroxyl to Silyl trifluoromethanesulfonate ester were conducted by $^{19}$F NMR spectroscopy, based on the reaction between Si-OH on the surface of silica particles and triflic anhydride (See equation-1).
\[
\equiv \text{Si-OH} + (\text{CF}_3\text{SO}_2\text{O})_2 \longrightarrow \equiv \text{Si-O-SO}_2\text{CF}_3 + \text{CF}_3\text{SO}_2\text{OH} \quad (1)
\]

The chemical shifts of methyl fluorine in triflic anhydride (reactant) and triflic acid (product) occur at -72.6 ppm and -77.3 ppm, respectively. Their integrated peak intensities allowed determination of the extent of the triflation reaction, assuming that the only source of triflic acid was from silanol esterification. However, if moisture is present, it can also hydrolyze the triflic anhydride to give two moles of triflic acid as shown in the following equation.

\[
(\text{CF}_3\text{SO}_2\text{O})_2 + \text{H}_2\text{O} \longrightarrow 2\text{CF}_3\text{SO}_2\text{OH} \quad (2)
\]

Therefore, a control experiment was carried out in a glovebox to examine the effect of moisture on measurement accuracy. By stirring the triflic anhydride in anhydrous toluene solvent for 24 and 48 hours at 60 °C, \(^{19}\text{F}\)-NMR data showed that the peak integration ratio of triflic acid to triflic anhydride was 1:148 (or 1:296 wt. mole ratio) and 1:80 (or 1:160 wt. mole ratio), respectively. This indicated that moisture present in glovebox during triflation reaction was negligible. After 24 and 48 hours of triflation reaction at 60 °C with 0.2 g of silica gel, this ratio increased to 1:9 and 1:2, respectively (see Figure 4.10). Based on this, estimated conversion percentage in esterification reaction was 44% after 24 hours and then reached 100% after 48 hours (by assuming a surface silanol concentration of 14 mmol/5g silica gel with 4.6 groups SiOH per nm\(^2\))^106.
Figure 4.10. $^{19}$F NMR study of silica gel reaction with triflic anhydride. In a control experiment, a strong peak at -72.6 ppm corresponding to fluorine of triflic anhydride can be observed. In a reaction with silica gel peak intensity at -77.3 ppm corresponding to triflic acid increases and reaches to maximum after 48 hours of reaction.

After reduction with DIBAL-H reagent, new peaks in FT-IR appeared at 2259 cm$^{-1}$ and 888 cm$^{-1}$, which correspond to Si-H stretching, and bending vibrations$^{96}$; see the pink curve in Figure 4.9. Note that Si-H frequency depends on the polarity of the other three bonds with silicon, varying from 2109 to 2250 in going from C-Si-H to O-Si-H$^{107}$.

Silicon hydride groups (Si-H) were further modified to alkylated silica (Si-R) by treating with 1-octadecene in the presence of UV light (254nm) for 2 hours$^{108}$. Samples were washed with anhydrous chloroform (CHCl$_3$) solvent to remove excess adsorbed 1-
octadecene and then air dried\textsuperscript{108, 109}. New peaks were observed at 2856 cm\textsuperscript{-1} and 2927 cm\textsuperscript{-1} corresponding to -CH\textsubscript{2} symmetric and asymmetric stretches, and shoulder peak at 2974 cm\textsuperscript{-1} corresponding to -CH\textsubscript{3}, confirming the alkyl chain derivatization of silica gel surface\textsuperscript{91, 98, 99, 101-103}.

In FTIR of alkyl derivatized silica gel spectrum, the absence of peaks corresponding to alkene C-H stretch (C=C-H) at 3083 cm\textsuperscript{-1} and C=C stretch at 1644 cm\textsuperscript{-1} indicated the absence of 1-Octadecene molecules physically adsorbed on the silica gel surfaces. To further gauge the effect of adsorbed alkenes, as opposed to chemically bonded alkyl groups, a control experiment was undertaken. The experiment involved, immersing the dried silica gel in a 1-octadecene solvent and exposing it to UV light under similar conditions, and finally washed with chloroform solvent after the completion of the reaction. The FTIR spectrum (see Figure 4.11) of silica gel after these steps showed no significant peaks corresponding to alkyl chains, indicating negligible adsorption of 1-octadecene on the silica surface.
Figure 4.11. FTIR of silica gel control experiment. Comparison of pure silica gel before reaction and after treatment with 1-Octadecene in the presence of UV light. No peaks corresponding to alkyl chains can be observed in 2800 to 3000 cm\(^{-1}\) region.

After studying the reaction on organic molecules and heterogeneous silica gel, the method was tested on a classic SiO\(_x\) wafer. Reaction progress was studied by FTIR and water contact angle (CA) measurement.

4.5.4. Results on SiO\(_x\) wafer reaction

Derivatization studies of SiO\(_x\) coated wafer provided a suitable comparison with silica gel (\textit{vide supra}) and low k CDO (\textit{vide infra}). These studies utilized silicon wafers decorated
with 100 nm thick silicon dioxide treated in identical reaction conditions. A clean SiO$_x$ wafer before the reaction showed the peaks (the black trace in Figure 4.12) corresponding to the stretching (3376 cm$^{-1}$) and bending vibrations (1296 cm$^{-1}$) of the silanol (Si-OH); the symmetric (890 cm$^{-1}$) and asymmetric stretching (1105 cm$^{-1}$) of cage Si-O-Si; and the bending vibration (610 cm$^{-1}$) of Si-O-Si$^2$. Lower peak intensities associated with O-H stretch vibrations (at 3500 cm$^{-1}$) when compared to Si-O-Si peaks (at 1100 cm$^{-1}$), indicated a lower net content of surface Si-OH groups on SiO$_x$ wafer films than silica gel (cf. black curves in Figures 4.9 and 4.12).

**Figure 4.12.** FT-IR spectra of SiO$_x$ wafer at various stages of the reaction: clean SiO$_x$ film (black); after treatment with triflic anhydride (blue); DIBAL-H reduced (pink); and after treatment with 1-octadecene (red).
The Si-OH absorption peak intensity significantly reduced upon triflation consistent with the surface accessibility of most Si-OH groups in the nonporous SiOx film (cf. black and blue traces). After the triflation reaction, new peaks appeared at 1039, 1228 cm\(^{-1}\) and 1193 cm\(^{-1}\) corresponding to symmetric and asymmetric stretches of S=O bond, and C-F stretch, respectively, as in the case of silica gel. However, their relative peak intensities were proportionately smaller. After DIBAL-H reduction, the Si-H peak at 2259 cm\(^{-1}\) was not observed presumably because of the lower total content of SiOH groups on the SiOx film surface compared to silica gel (cf. pink curves in Figures 4.9 and 4.12). Nevertheless, silica gel Si-H peaks disappeared upon exposure to air within half an hour. Similarly, limited stability of the Si-H groups in the air has been previously noted when the remaining three group bound to silicon are highly polar, such as Si-O.

Hydrosilylation of the hydrogen-terminated Silicon surface with long-chain terminal alkene was conducted without exposing the sample to atmospheric moisture. After the photochemical reaction, the presence of peaks at 2853 and 2924 cm\(^{-1}\), corresponding to CH\(_2\) symmetric and asymmetric stretches, and a shoulder peak at 2976 cm\(^{-1}\) (red curve and red inset in Figure 3) confirmed the hydrosilylation. The ratio of C-H (2800-2950 cm\(^{-1}\)) to Si-O-Si (1100 cm\(^{-1}\)) peak intensities was higher than silica gel (cf. Figures 4.9 and 4.12) and could arise from a greater percentage of inaccessible surface SiOH groups in silica gel. The absence of peaks at 3083 and 1004 cm\(^{-1}\) corresponding to alkene C=C-H stretching and bending vibrations, and the C=C stretch at 1644 cm\(^{-1}\) of 1-octadecene, suggested that 1-octadecene was chemically bonded to the silicon surface rather than
physically adsorbed. Contact angles before and after derivatization of SiO₃ surfaces were 11±2° and 101±4° respectively.

### 4.5.5. Results on CDO wafer reaction

![Reaction Pathway Diagram]

**Figure 4.13.** Common reaction pathway for hydrogen termination, Octadecane derivatization and experimental setup of carbon doped silicon oxide (CDO) wafer.

FT-IR, X-ray photoelectron spectroscopy (XPS) and water contact angles (CA) techniques were employed to monitor the progress of the reaction, and the appearance of new functional groups on CDO wafer surfaces. When compared to the SiO₃ wafer the water contact angle on CDO after RCA clean, was significantly higher (36±3° vs. 11±2°, see Table 4.2, below) consistent with a more hydrophobic surface due to its higher carbon content and a lower density of SiOH group on the top surface (cf. black curves in Figures 4.12 and 4.14).

Figure 4.14 shows the transmission FT-IR spectra of a CDO wafer at different stages of derivatization. Major peaks observed correlated well with the SiO₃ wafer IR peaks illustrated in Figure 4.12. However, significantly lower peak intensity in the OH vibrational frequency range (≈ 3300 cm⁻¹) compared to silica gel or SiO₃ wafer (Figures 2, 3), implied a lower net content of OH on the CDO surface. Accordingly, after triflation, intensities of
peaks associated with S=O and C-F stretches, (shown Figure 4.14, the blue curve (inset)), were smaller than the corresponding peaks observed in the triflated SiOₓ films (blue curve, Figure 4.12).

**Figure 4.14.** FT-IR spectra of CDO wafer in various stages of reaction; Clean CDO (black), after reaction with triflic anhydride (blue), after reaction with DIBAL-H (pink), after glacial acetic acid washing (green) and after octadecene derivatization (red). Stars indicated in the DIBAL-H treated spectrum correspond to alkyl groups arising from the surface-bound, excess DIBAL-H reaction.
products. Insets expand selected spectral regions of triflic anhydride Tf$_2$O and octadecene-derivatized silica in the IR spectra.

Reduction of triflate ester on the CDO surface using DIBAL-H proceeded as in the case of SiO$_x$.$^{110}$ After reduction with DIBAL-H several new peaks, not observed in silica gel or SiO$_x$ surfaces, appeared in the range from 2800-3000 cm$^{-1}$, attributed to alkyl C-H vibrations. These peaks could originate from traces of the degraded DIBAL-H reagent (e.g., diisobutyl groups), physically adsorbed or trapped in pores of CDO surface (pink trace, Figure 4.14). Also, supporting the above hypothesis was an observed increase in the peak intensity near the OH region (3300 cm$^{-1}$) due to the formation of Al-OH groups, a byproduct of the reduction reaction. Since an aluminum oxide or its hydroxide dissolves readily in an acidic medium, we treated the wafer with 25% glacial acetic acid in anhydrous toluene for five minutes to remove trapped/degraded DIBAL-H reagent from the CDO surface. The FT-IR spectra collected after the acid treatment displayed a reduction in OH peak intensity and disappearance of alkyl shoulder-peaks (the green trace, Figure 4.14).

A control experiment to further test the hypothesis of DIBAL-H binding on the CDO surface entailed treating clean CDO, SiO$_x$ wafers, and silica gel sample with a DIBAL-H reagent (see Figure 4.15) in anhydrous toluene. After 24 hours, samples were washed with dry toluene and dried. Their FT-IR spectra revealed the alkyl peaks only for the CDO confirming preferential DIBAL-H trapping/adsorption on the surface/pores of a slightly hydrophobic CDO surface (vide supra) as opposed to hydrophilic surfaces of silica gel or SiO$_x$ wafer.
Figure 4.15. Control experiment showing that only the porous structure of CDO traps DIBAL-H from anhydrous toluene solution. A. Silica Gel, B. SiOₓ wafer C. CDO wafer. Only C indicates the presence of C-H stretch vibrations in the 250-2950 cm⁻¹ region.

Like the SiOₓ wafer, Si-H stretching peaks were not detected presumably either because of a low surface concentration or rapid degradation in the air during FT-IR data acquisition. Furthermore, observed weaker IR peak intensities (blue inset in Figure 4.14) for the triflate groups on CDO and SiOₓ wafer surface compared to silica gel (blue curves in Figures 4.9, 4.12 and 4.14) supported the above interpretation. The FT-IR spectrum after hydrosilylation was similar to the SiOₓ wafer results (cf. red curves in Figures 4.12 and 4.14). A control experiment involving hydrosilylation of underivatized CDO surface revealed a lack of adsorption of octadecene or its photochemical reaction product (Figure 4.16). The observation of DIBAL-H adsorption/trapping (Figure 4.15) but not of
octadecene on a clean CDO surface (Figure 4.16) implies that a slightly hydrophilic CDO pore is unable to retain nonpolar octadecene, pointing to the amphipathic nature of the pore wall. This weaker pore-octadecene interaction would result in ready removal of octadecene, but not DIBAL-H, during the CDO wafer rinsing/drying stage, before collecting respective IR spectra. In principle, following DIBAL-H reduction of pore surface triflate groups, octadecene can enter CDO pores and derivatize the hydrophobic silicon hydride pore surfaces (red curve Figure 4.14). Alternatively, the steric effects such as nanopore size may preclude entry of the larger octadecane while allowing smaller DIBAL-H molecules. Future studies using smaller size alkenes are planned, however, with the data at hand, we are unable to verify whether the internal pore surface is derivatized or not.
Figure 4.16. FTIR of CDO wafer control experiment. Spectral comparison of clean CDO, i.e., before reaction and CDO after treatment with 1-octadecene in the presence of UV light under identical conditions, were shown. No peaks corresponding to alkyl chains can be observed in 2800 to 3000 cm\(^{-1}\) region.

A complementary technique of X-ray photoelectron spectroscopy (XPS) provided further support for derivatization on CDO wafers. Survey spectra obtained from a CDO surface before (Figure 4.17a) and after (Figure 4.17c) reaction revealed Si, C, and O elements. After derivatization, the intensity of the carbon (1s) peak increased, consistent with the derivatization of surface groups with alkyl chains. In deconvoluted high-resolution Si (2p) spectra of clean CDO wafers before the reaction (Figure 4.17b), three strong peaks at 101.6, 102.3 and 103.1 eV corresponding to Si-C, -Si-O-C- and -O-Si-O-, respectively were observed. The XPS spectrum after the reaction (Figure 4.17d) had the same three
strong peaks at similar positions but with different intensities. The area of the \(-\text{O-Si-O-}\) peak decreased from 26 % to 10 % of the Si (2p) peak area, and the relative area of \(-\text{Si-C}\) peak increased from 11 % to 22 %.

Figure 4.17. XPS spectra of CDO wafer. Survey spectrum, a, and Si (2p) high-resolution spectrum, b, collected before the reaction and c, d are the corresponding spectra collected after the reaction.

This finding validates the conversion of \(-\text{O-Si-OH}\) to \(-\text{O-Si-R}\) (R= octadecyl). No significant change was observed, however, in the peak area for the \(-\text{Si-O-C-}\) group, indicating no structural modification in the cage structure of the CDO film. More surface chemical studies are needed to confirm this indication.
Derivatization of the CDO surface with octadecene should change its hydrophobicity. The contact angle of water droplets deposited on the surface confirmed that shift (Figure 4.18). Clean CDO wafers (Figure 4.18a) had a low contact angle 36±3° caused by the presence of the hydrophilic hydroxyl groups (Si-OH). Derivatization with octadecene increased the contact angle to about 81±2° (Figure 4.18b, Table 4.2). The water contact angles on derivatized surfaces were stable for more than a year (See Figure 4.18) as were the corresponding IR spectra (Figure 4.19).

![Figure 4.18. Water contact angles measured on un-derivatized (clean CDO) and octadecene derivatized CDO wafers at various time intervals.](image)

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Contact angle (deg.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Before reaction (Clean CDO)</td>
<td>36 ± 3°</td>
</tr>
<tr>
<td>After reaction</td>
<td>81 ± 2°</td>
</tr>
<tr>
<td>9 months later</td>
<td>82 ± 3°</td>
</tr>
<tr>
<td>14 months later</td>
<td>82 ± 2°</td>
</tr>
</tbody>
</table>

*Table 4.2. Contact angle results of a water droplet on clean CDO and octadecyl derivatized CDO at various time intervals.*
Figure 4.19. FTIR spectra comparison of octadecyl derivatized CDO wafer before and after one-year duration.

4.5.6. Comparison of Octadecyl chains derivatized via DIHAL-H chemistry and Octadecyl trichlorosilane (OTS) routes

Octadecyltrichlorosilane (OTS), a commonly used silanization agent, readily forms a self-assembled monolayer on SiO$_x$ film transforming it to from hydrophilic to a hydrophobic surface by inserting an alkoxy ether (Si-O-R) group bearing a C18 alkyl chain. Hence, the results of octadecene-based vs. the OTS-based derivatization on CDO and SiO$_x$ surfaces were compared. Qualitatively, FT-IR spectra, in the relevant region shown in Figure 4.20, indicated that the thickness of the hydrocarbon layer is higher for octadecene
derivatization than the OTS SAMs (blue vs. red traces), while the silanol peak intensities were greater in the latter. An increase in OH peak intensity could come from the free Si-OH groups on the hydrolyzed OTS molecule as schematically illustrated in Figure 4.20 (inset). These free O-H groups provide interfacial hydrophilic sites for water adsorption underneath the alkyl chains of the OTS film. Thus, an insertion of a Si-OR group which has a higher dipole moment than Si-R group and new free –OH groups would be expected to increase the \( k \) values of CDO/Oxide films derivatized using OTS.

![Figure 4.20. Comparison CDO and SiO\(_x\) wafer derivatized by DIBAL-H assisted chemistry (blue) vs. OTS chemistry (red). Black curves depict IR spectra of the respective wafers before derivatization.](image)

4.5.7. Characterization of film dielectric constants \((k)\)

Effects of Maxwell-Wagner polarization (MWP) hampered the low frequency (0.1-1 MHz) electrical measurements of \( k \) values due to adsorption of moisture in the nanopores of CDO. Partial dissociation of trapped/adsorbed water molecules imparts to the film a finite electrical conductivity creating electrically leaky (resistive) as opposed to insulating films\(^{111}\). Such samples\(^{112}\) have a higher dielectric contribution arising from distortion and
dipolar polarization which raises their effective $k^{113}$. To disentangle MWP and other polarization effects require measurements of the film porosity and frequency dependent dielectric constants as a function of water saturation$^{114}$ (i.e., humidity). Since these data were unavailable, a qualitative consideration of capacitance (proportional to $k$ (1MHz)) measurements (Figure 4.21) revealed the following ordering in the magnitude of $k^{OTS} > k^{CDO} > k^{DIBAL-H}$, consistent with the expected interfacial polarity of these samples (vide supra). One way around this problem is to reduce surface SiOH group concentration by high-temperature baking in an inert environment.

![C-V of three samples](image)

**Figure 4.21.** Capacitance vs. voltage curves measured using Hg probe method. Hg contact area was $3.1 \times 10^{-3}$ cm$^2$. Samples 1, 2 and 3 are underivatized CDO, OTS and Octadecene derivatized CDO wafers respectively.

Another approach to access $k$ is through measurements of film refractive index; noting that high frequency (optical) $k$ values simply depend on electronic polarization and scale as the square of refractive index$^{115}$. Note, $k$ values at optical frequencies do not contain dielectric contributions from MWP and dipole polarizations$^{113}$ (see below). Analysis of reflectance spectra (Filmetrics) collected from these wafers yielded best-fit values of refractive index and film thickness; tabulated in Table 4.3. Best-fit refractive
index values for CDO films were lower than SiO\textsubscript{x} films commensurate with the porous structure of CDO that can trap either water or air, both of which have lower refractive indices than pure SiO\textsubscript{2}. Furthermore, the effective refractive index values of OTS derivatized films were consistently lower than the octadecene derivatized surface implying perhaps their larger adsorbed moisture content. In either SiO\textsubscript{x} or CDO films, the refractive index of DIBAL-H-treated films was greater than the OTS-treated films corresponding to its thicker hydrocarbon film of the greater refractive index (1.4439) than refractive indices of either water (1.33) or air (1.0).

To estimate film porosity of CDO crudely, the refractive index of these composite films ($\eta_{exp}$) was decomposed as a sum of volume fraction ($\phi$) weighted refractive indices of the constituents\textsuperscript{116}:

$$\eta_{exp} = \sum \eta_i \phi_i$$

where index $i$ denotes a specific component and $\phi_i$ its volume fraction. Assuming that the CDO matrix is made up of SiO\textsubscript{2} ($\eta_{SiO2} = 1.473$) and pores are filled completely with either air ($\eta_{SiO2} = 1$), or water ($\eta_{H2O} = 1.333$) gave approximate the effective porosity of 3 or 11 percent for CDO respectively. Within limits of optical measurements of the film thickness using filmetrics, the hydrocarbon layer thickness in the octadecene derivatization scheme was comparable ($\approx 5\text{nm}$) for both CDO and SiO\textsubscript{x} (Table 4.3). These values were significantly higher than expected for a densely packed upright monolayer of C18 chain (2 nm). Ellipsometric measurements, however, consistently gave a higher thickness of hydrocarbon layer in octadecene
derivatized samples supporting the observed trend of higher IR peak intensities for the C-H stretch vibrations (Figure 4.20). These higher IR intensities are consistent with partial cross-linking/polymerization of octadecene during the hydrosilylation step\textsuperscript{117}.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Refractive index (n)</th>
<th>$\mu = \frac{n^2}{n_2^2}$</th>
<th>Overlayer Thickness (nm)</th>
<th>Hydrocarbon layer thickness (nm)</th>
<th>CA (deg.)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Filmmetrics</td>
<td>Ellipsometry</td>
<td></td>
</tr>
<tr>
<td>CDO clean</td>
<td>1.457</td>
<td>2.12</td>
<td>101.2 ± 0.6</td>
<td>0</td>
<td>36 ± 3°</td>
</tr>
<tr>
<td>1-Octadecene deriv. CDO through DIBAL-H</td>
<td>1.447</td>
<td>2.09</td>
<td>103.1 ± 2.4</td>
<td>2 ± 3</td>
<td>87 ± 3°</td>
</tr>
<tr>
<td>OTS deriv. CDO</td>
<td>1.412</td>
<td>1.99</td>
<td>106.4 ± 0.3</td>
<td>3 ± 1</td>
<td>91 ± 4°</td>
</tr>
<tr>
<td>SiO\textsubscript{2} clean</td>
<td>1.473</td>
<td>2.17</td>
<td>107 ± 2</td>
<td>4 ± 2</td>
<td>11 ± 2°</td>
</tr>
<tr>
<td>1-Octadecene deriv. SiO\textsubscript{2} through DIBAL-H</td>
<td>1.448</td>
<td>2.09</td>
<td>111.2 ± 0.9</td>
<td>4 ± 3</td>
<td>101 ± 4°</td>
</tr>
<tr>
<td>OTS deriv. SiO\textsubscript{2}</td>
<td>1.435</td>
<td>2.06</td>
<td>118.9 ± 0.6</td>
<td>2 ± 3</td>
<td>105 ± 3°</td>
</tr>
</tbody>
</table>

**Table 4.3.** Contact angle, film thicknesses of CDO and SiO\textsubscript{2} wafers.

The contact angle measurements revealed similar trends. The contact angles for both derivatization schemes depended on the nature film, i.e., CDO vs. SiO\textsubscript{2} and not on the chemical mode of derivatization, i.e., silanization vs. hydrosilylation. The CDO surface which had higher contact angle before derivatization (36° (CDO) vs. 11° (SiO\textsubscript{2})) but exhibited lower contact angle after derivatization (87° vs. 91° for CDO derivatized with octadecene and OTS respectively). Applying the Cassie equation for a measured contact angle ($\theta_m$) on a surface made up of two types of
regions (labeled with superscripts 1 and 2) with differing contact angles, $\theta_1$ and $\theta_2$, gives\textsuperscript{118}:

$$\cos \theta_m = f (\cos \theta_1 - \cos \theta_2) + \cos \theta_2$$

where $f$ represents a fraction of the surface of type 1. Crudely, identifying region 1 as SiO$_x$ ($\theta_1 = 11^\circ$) and region 2 as pores (i.e., air with $\theta_2 = 180^\circ$) on the CDO surface, the estimated upper limit for $f = 0.91$ implying a 9 percent surface porosity. Should the lower OH density on the CDO surface be accounted for by setting ($\theta_1 > 11^\circ$) the measured porosity would decrease. For example, if $\theta_1 = 30^\circ$ for the nonporous CDO surface, then the calculated pore fraction would be 3 percent, similar to the estimates from the refractive index measurements (\textit{Vide Supra}). Since we did not have a nonporous CDO sample, we did not have a value for $\theta_1$.

For SiO$_x$ surface the observed contact angles after derivation were $101^\circ$ (octadecene) vs. $104^\circ$ (OTS) implying a more densely packed monolayer as was noted elsewhere\textsuperscript{119}. Again, applying the Cassie equation, but now ascribing region 1 to the densely packed hydrocarbon monolayer ($\theta_1 = 107^\circ$) and region 2 to underivatized SiO$_x$ ($\theta_2 = 11^\circ$, from Table 1), estimated $f$ values were 0.97 for OTS and 0.92 for Octadecene derivatized SiO$_x$ surfaces. By contrast, the CDO surface had a thicker but more loosely packed hydrocarbon layer due to the lower density of surface OH groups, inferred from its higher initial contact angle. The corresponding $f$ values on the CDO surfaces were, as expected, smaller; 0.75 (OTS) and 0.69 (Octadecene) respectively. The results also illustrated that the hydrocarbon layer thickness did not affect the contact angle, but the areal fraction of derivatized (i.e., $f$) surface did.
The reaction of hydrofluoric acid (HF) can also enhance the hydrophobicity of the silicon surface by creating a hydrogen-terminated surface that may be hydrosilylated, but it would produce undesirable etching of the CDO film. Recently, our group has used sacrificial silanes to prevent the etching of SiOx by HF. The chemistry, however, is carried out in an aqueous medium of low pH, which would likely reoxidize Si-H groups² and hence is undesirable for this protecting the inner layer dielectrics from moisture.

4.6. Thiol functionalization of CDO surface to enhance NWs interaction to substrate

Thiol functional groups(-SH) shows strong interaction towards noble and coinage metals¹²⁰. Thiol interaction and formation of self-assembled monolayers (SAM) of alkane thiols on copper has been extensively studied¹²⁰-¹²³. Thiol functionalization of CDO surface was explored to enhance the chemical interaction of as-deposited Cu/Ni NWs to CDO substrate.

![Scheme 4.4. Reaction scheme describing thiol functionalization of CDO surface.](image)

Thiol functionalization of CDO was carried out with the DIBAL-H approach, See Scheme 4.4. The reaction was carried out under identical conditions, except 1-Octadecene is replaced with a 2-propene thiol in the photochemical reaction. In the FTIR characterization, 2888 and 2941 cm⁻¹ peaks are attributed to propyl chain (3 carbon chain)
and 2564 cm\(^{-1}\) is a characteristic peak corresponding to thiol (-S-H) stretch suggests the thiol termination, See Figure 4.22.

![FTIR characterization of as received CDO (black curve) and thiol (-SH) functionalized CDO (red curve).](image)

**Figure 4.22.** FTIR characterization of as received CDO (black curve) and thiol (-SH) functionalized CDO (red curve).

**Adhesion strength of Cu/Ni NWs to thiol (-SH) functionalized CDO surface:** The main purpose of thiol functionalization is to strongly hold the Cu/Ni NWs after deposition onto CDO surface. The adhesion strength of as received CDO (non-functionalized) vs. thiol functionalized CDO was qualitatively assessed by ultra-sonic treatments, see set up in Figure 4.23a. The procedure involves, drop casting of equal amounts of Cu/Ni NWs
solution (250 µL, 5 mg/mL NWs concentration) on as received CDO and thiol-functionalized CDO (2x2 cm coupons). After drying, wafers substrates were immersed in ethanol solvent in a beaker and placed in ultrasonicator (160 watts); Figure 4.23a. In case of non-functionalized CDO, traces of NWs were washed off within 4 seconds of ultrasonic treatment, see Figure 4.23b. Whereas for thiol functionalized CDO, holds the NWs strongly till 30 seconds of sonication treatment, see Figure 4.23c. And, after 150 seconds of ultrasonic treatments traces of NWs were washed off, this suggests the enhanced interaction of NWs to thiol functionalized CDO surface.

![Figure 4.23](image)

**Figure 4.23.** Digital photograph showing ultrasonication setup (a), for qualitatively testing the adhesion strength of Cu/Ni NWs to as received CDO (b), and thiol-functionalized CDO surface (c).
4.7. Summary of octadecyl and thiol functionalization

The current plasma treatments to prevent uptake of atmospheric moisture by low dielectric CDO cause thinning of the films. The classical HMDS derivatization to make surfaces hydrophobic does not provide sufficient protection. The studies presented here show a selective derivatization of surface hydrophilic Si-OH groups to form a hydrophobic alkyl-modified silicon (Si-R, R= \(-\text{C}_{18}\text{H}_{37}\)) without disturbing the rest of the cage structure of the CDO film. This selective hydrophobic modification should help maintain the ultra-low k of the insulating film. The method does require an additional processing step to remove pore-trapped DIBAL-H reagent that can be removed readily with mild acetic acid. The versatile method allows derivatization of Si-OH functional groups on the surface of silica gel or CDO wafers with a variety of other organic molecules useful in designing other biochips, sensors\textsuperscript{124-126}. Preliminary studies indicate that the degree of hydrophobicity created on surface compares favorably with the classic silanization approach, bypassing interfacial insertion of the polar Si-O-R group. It provides acceptable thermal processing stability for the interconnect processing. The same DIBAL-H reduction approach was used for the thiol (-SH) functionalization. And, the qualitative analysis by ultra-sonic treatments suggests the strong adhesion of Cu/Ni NWs to the thiol-modified CDO surface.
CHAPTER FIVE
NANOWIRES SELF-ASSEMBLY AND ELECTRICAL CHARACTERIZATION

Over the last two decades, substantial efforts have been devoted to the assembly of 1D nanostructures into nanoscale circuits and interconnects. In this chapter, the self-assembly of Cu/Ni NWs into an interconnect structure on a CDO wafer using the combination of photolithography and magnetic field assisted NWs alignment techniques is presented. CDO is an important nanoporous low-

The Strong interaction between the substrate and deposited NWs is essential for the successful assembly of NWs. Here the chemical interaction of thiol (-SH) functional groups to copper and nickel was exploited to strongly adhere the NWs after deposition onto the CDO substrate. CDO surface was functionalized with dangling thiol (-SH) functional groups using the protocol described in chapter 4, section 4.6. Thiol functionalization and its characterization were discussed in detail in chapter 4.

5.1. Self-assembly of Cu/Ni NWs into interconnects

Schematic diagram shown in Figure 5.1 describes the process flow for the fabrication of Cu/Ni NWs based interconnects. The protocol involves first transferring the desired circuit pattern (300µm width and 4mm length channel pattern) onto the thiol-functionalized CDO substrate using a photolithography process, see Section 2.2.5 for more details about the photolithography process. The above-patterned substrate was placed in between the
electromagnet poles like the schematic shown in Figures 3.10 and 5.1. Cu/Ni NWs aqueous solution (100 µL of 0.5 mg/mL solution) was placed on to the substrate and allowed to dry in the presence of magnetic field (2500 G). In the presence of magnetic field, NWs oriented in the applied magnetic field direction and begin to form NW chains in the solution; see Figure 3.12. After allowing the solvent to dry under ambient conditions, well aligned NWs deposited on to the surface of photoresist as well as in the thiol-modified trenches on CDO. The NWs which were deposited in the trenches are strongly held to that surface through the covalent bond to the thiol functional groups on CDO surface. The NWs which were deposited on photoresist were loosely bound. These loosely bound NWs and photoresist were removed by washing with acetone solvent (the so-called lift-off process). Acetone solvent dissolved the photoresist and simultaneously washed off the loosely bound NWs from the CDO surface. Finally, the selectively deposited and well-aligned Cu/Ni NWs in the circuit pattern remained on the substrate like the optical micrographs shown in Figure 5.2.
5.2. Electrical characterization of Cu/Ni NWs based interconnect channels

5.2.1. Effect of nanowire type (pure Cu NWs vs. Cu/Ni NWs) and orientation (random vs. aligned) on channel resistance.

Electrical characterization studies of as-fabricated NWs channels were carried out by measuring the electrical resistance using a Keithley-617 programmable digital...
multimeter. Contact pads on both ends of the interconnect channel were fabricated by depositing a small amount (20µL) of silver ink, see inset image in Figure 5.2a. The contact resistance of the dry silver ink was observed as < 1Ω, negligible in comparison to the resistance of interconnect channels. For the comparison, along with the well-aligned Cu/Ni NWs deposited in the channel, several samples were fabricated with similar dimensions of interconnect channels wherein randomly arranged Cu/Ni and pure Cu NWs were deposited. All the above-fabricated channels used 0.5 mg/mL (dry mass of NWs) concentration of NWs in aqueous suspensions. The common channel lengths and widths were 4 mm and 300 µm respectively. Schematic diagrams, digital photographs and optical micrographs of the as mentioned NWs interconnect channels appear in Figures 5.4, 5.4, 5.5 & 5.6.

**Figure 5.3.** Schematic diagrams and digital photographs showing interconnect channels fabricated by using different types and arrangement styles of NWs. The random arrangement of Cu NWs channel in a), unidirectionally aligned Cu/Ni NWs channel in b), and randomly arranged Cu/Ni NWs channel is shown in c).
**Figure 5.4.** Optical micrographs of Cu NWs based interconnect channel at various magnification scales showing the random arrangement of Cu NWs.

**Figure 5.5.** Optical micrographs of Cu/Ni NWs based interconnect channel at various magnification scales showing directionally well-arranged NWs.
Figure 5.6. Optical micrographs of Cu/Ni NWs based interconnect channel at various magnification scales showing randomly arranged NWs.

Figure 5.7. Plot showing resistance values of well-aligned and randomly arranged Cu/Ni NWs, and randomly arranged pure Cu NWs based interconnect channels.

The average resistance values of interconnect channels which were made of well-aligned Cu/Ni NWs, and randomly arranged Cu/Ni NWs and pure Cu NWs were $290 \pm 20$ kΩ, $780 \pm 40$ kΩ, and $370 \pm 30$ kΩ respectively, See Figure 5.7. Average resistance values and uncertainties were determined by measuring the resistance values of 3 interconnect
channels fabricated for each type of NWs arrangement. Lower resistance for well-aligned Cu/Ni NWs is attributed to the more number of effective conductive channels due to end-to-end magnetic contacts as illustrated in Figure 5.8.

![Randomly arranged vs Well arranged](image)

**Figure 5.8.** Schematic depicting the randomly aligned Cu/Ni NWs channel fabricated in absence of magnetic field and orientationally controlled NWs channel fabricated in presence of applied magnetic field.

High resistance in case of randomly arranged Cu/Ni NWs suggests the percolating network of NWs leads to a higher number of inter NW hops and effectively a lower number of effective channels. Whereas for the pure Cu NWs, the resistance is close to the well-aligned Cu/Ni NWs. This could result from the high conductivity of copper in comparison to the nickel.

### 5.2.2. Effect of nanowires alignment order on electrical performance

The above studies reveal the importance of NWs alignment over randomly organized NWs on achieving a lower channel resistance. For randomly oriented nanowires, the probability of making interwire contact is uniform over the length of NW while for magnetized NWs should have a significantly higher probability near its ends. This anisotropic sticking probability distribution should be a function of magnetic strength.
Thus, the effect of applied magnetic field strength (0-2500 G) on the alignment order and sample resistance was investigated. Optical micrographs of channels mentioned above appear in Figure 5.9. As the magnetic field increased, the resistance of NWs channels decreased, See Figure 5.10. The observation differs in comparison to the other metallic or carbon nanotube-based results$^{131}$. This could be attributed to the ferromagnetic nature of nickel which enhances the end to end NW connections (Figure 5.8) increasing the number of effective channels formed.

![Figures 5.9](image)

**Figure 5.9.** Optical microscope images of interconnect channels fabricated at various applied magnetic field strength from 0 to 2500 Gauss.
5.2.3. Effect of NWs density on resistance of well-aligned Cu/Ni NWs channels

The resistance of the NWs based interconnect channel is determined by the number of effective conductive channels that successfully allows the electrons to pass through from one end to the other end of the channel. The number of effective conductive channels can be increased by increasing the number density of NWs, i.e., increasing the NW concentration higher than required for the percolation threshold. The NW concentration in depositing suspension was increased from 0.5, 1.0, and 2.0 mg/mL for the deposition on the patterned substrate. Optical micrographs of the corresponding interconnect channels are shown in Figure 5.11. Plot in Figure 5.12, showed a decrease in the resistance values from 290 ± 20 kΩ (0.5 mg/mL), 160 ± 20 kΩ (1.0 mg/mL) to 100 ± 20 kΩ (2.0 mg/mL) as the NWs density increased consistent with a simple percolation picture.

Figure 5.10. Interconnect channel resistance values of Cu/Ni NWs fabricated using various applied magnetic field strength from 0 to 2500 G.
Figure 5.1. Low and high magnification optical micrographs showing of interconnect channels fabricated by using 0.5 mg/mL, 1.0 mg/mL and 2 mg/mL respectively.

Figure 5.12. Resistance values of Cu/Ni NWs based interconnect channels as a function of NW concentrations in suspension.

5.2.4. Thermal stability of Cu NWs and Cu/Ni NWs based interconnect channels

The thermal stability of NWs is an important factor for the reliable performance of NWs based devices. Nickel coated NWs were found to be more oxidation resistant comparison to pure Cu NWs.$^{59}$ The thermal stability of as-fabricated interconnects channels were
examined by heating the substrate at 100 °C in the air while simultaneously measuring the resistance. For the thermal stability studies, interconnect channels were fabricated by using 0.5 mg/mL NWs solutions. Figure 5.13 shows the resistance value as a function of heating duration for randomly arranged pure Cu NWs and Cu/Ni NWs channels and well-aligned Cu/Ni NWs channels. The resistance of pure Cu NWs channel rapidly increased by over tenfold and crossed 20 MΩ in less than 10 minutes. The resistance increase resulted from the rapid oxidation of pure copper in the air. Whereas for randomly arranged and well-aligned Cu/Ni NWs, the resistance rapidly increased by barely 11% and remained at constant throughout the heating duration. And even after heating for 72 hours, no significant change in resistance was observed suggesting a reliable performance of Cu/Ni NWs at high temperature.

Figure 5.13. Thermal stability of well-aligned and randomly arranged Cu/Ni NWs channels and randomly aligned Cu/Ni NWs channels at 100 °C in air. Inset plot in c) highlights the resistance curves of two type of Cu/Ni NWs arrangements.
5.3. Summary of Cu/Ni NWs self-assembly and electrical characterization

Despite numerous efforts, there were no successful techniques reported for the large-scale assembly of copper nanowires. Here, these studies show the potential of the magnetic field assisted technique to control the orientation and ability to self-assemble the NWs on the large-scale area. The result was achieved by coating the copper nanowires surface with a thin layer of ferromagnetic nickel. Electrical characterization results of self-assembled Cu/Ni NWs channels suggest the importance of orientational alignment for achieving lower resistance. Also, thermal stability studies reveal Cu/Ni NWs are highly resistant to oxidation under open-air conditions even at high temperature.
CHAPTER SIX

CONCLUSIONS AND FUTURE WORK

6.1. Conclusions

As the on-chip feature sizes are getting smaller, copper filling in the ever-decreasing channels is getting difficult and reducing the process yield. Therefore, new approaches are needed to supplement the standard Dual Damascene (DD) process. Fabrication of interconnects in a bottom-up approach, using 1D nanocrystals as building blocks offers a potential solution for future high performance integrated circuits. This work demonstrates a novel and simple solution processable technique to self-assemble the Cu/Ni NWs into interconnect channels on the low-k material; carbon doped silicon dioxide (CDO).

CDO is a highly nano-porous and hybrid low-k dielectric used in ultra large-scale integration (ULSI). For the self-assembly process, CDO surface was functionalized with thiol (-SH) functional groups to bind Cu/Ni NWs. Thiol functionalization was achieved via a novel approach developed to functionalize selectively the surface exposed hydrophilic Si-OH groups. Also, the same reaction was used to address the moisture uptake issue of CDO. In the chip integration process, an oxygen plasma irreversibly changes the silicon functional groups on the CDO surface from hydrophobic to hydrophilic, which causes an adverse effect on increasing the $k$ value by taking up moisture. This present work solves this issue by selectively functionalizing the surface exposed Si-OH functional groups to hydrophobic octadecyl terminated silicon and helps in maintain the low $k$ value.

In a bottom-up method, dimensionally controlled nanowires synthesis and precisely controlling the orientation and positioning of NWs are very critical. Here, in a wet synthesis
process, reagent concentrations and reaction temperatures were optimized to generate high aspect ratio and surface morphology controlled Cu NWs. This thesis work developed the magnetic field assisted technique to control the orientation of Cu/Ni NWs on a large surface area. Coating a thin layer of ferromagnetic nickel (Ni) onto the Cu NWs surface imparted magnetic field-mediated alignment of the composite NWs. Furthermore, the nickel coating helped in protecting the copper core from oxidation.

Finally, Cu/Ni NWs self-assembly into interconnect channels was demonstrated using the combination of photolithography and magnetic field assisted techniques. The electrical measurements revealed that well-aligned NWs exhibited less than two-fold lower resistance than the randomly organized Cu/Ni NWs channels. Also, as the degree of alignment of the NWs increased, a decrease in resistance was observed. This result is in stark contrast to the simulations results discussed for other types of NWs (Ag NWs, Au NWs and CNT). The enhanced performance of the magnetic field assisted alignment technique appeared to be related to the propensity of end-to-end contacts thereby reducing an overall number of nanowire contacts for electron transport between the two contacting electrodes. Furthermore, Cu/Ni NWs showed a higher thermal stability than the pure Cu NWs, due to their high oxidation resistance. This led to a more reliable electrical performance under ambient conditions.

6.2. Limitations

One of the important limitation of the as developed technique is, it can only apply to the ferro/para magnetic nanowires. For non-magnetic nanowires (pure metallic or carbon nanotubes etc.), they have to be coated with a magnetically active material to give a net
alignment when subjected to a modest magnetic field (2000 G). Here, the Cu/Ni NWs self-assembly process was demonstrated in 300 µm width channels, but the future technology requires fabrication of ultra-thin (< 5nm) interconnect channels⁹. For this, the self-assembly has to be performed with ultra-thin nanowires.

Another area needing improvement is a reduction in the observed high resistance values of NWs based interconnect channels. The significant contribution to the high resistance has been attributed to NW to NW junctional resistance¹³¹,¹³². This nanowire junction resistance can be minimized by fusing or welding the nanowire contacts either by thermal or laser ablation techniques¹³¹,¹³². Also, this technique only demonstrates the NWs self-assembly in a two-dimensional manner, but it needs to be adopted for the fabrication of copper vias (vertically aligned NWs which would connect to the two adjacent layers). Therefore, more efforts are needed to apply the as developed magnetic field assisted self-assembly technique to fulfill the future interconnect fabrication needs.

6.3. Other applications

Copper nanowires (Cu NWs) and nickel coated Cu NWs (Cu/Ni NWs) have been successfully demonstrated as an excellent material of choice for the preparation of highly flexible and transparent electrodes³³,³⁴,⁵⁹,⁶⁶. In case of Ag NWs, it was demonstrated that orientationally well-aligned NWs (deposited as a crossed arrays or mesh, like the image shown in Figure 3.11c), yields high transparency and lower sheet resistance values as opposed to randomly organized NWs arrays⁴⁵,⁴⁹. Cu/Ni NWs are the better choice than Ag NWs due to its cheap precursor cost, high oxidation resistance and orientability that can be well controlled in a large-scale area. Currently, studies of Cu/Ni NWs deposition as a mesh
on a polyethylenetetraphthalate (PET) substrate for the application of highly transparent and flexible devices are underway in our lab.

6.4. Future work

In this work, 300 µm width and 4 mm lengthy channels were used to demonstrate the self-assembly process. However, one may use the magnetic field assisted technique to assemble the single NWs based ultra-thin channels. In the literature, researchers have demonstrated the synthesis of three-atom wide copper nanowires\textsuperscript{133}. The developed magnetic field assisted self-assembly process is very promising, and perhaps it can be used to fabricate the three-atom wide interconnect channels which are highly desirable for future commercial chip fabrication.
REFERENCES


42. Lu, W.; Lieber, C. M., Nanoelectronics from the bottom up. *Nature materials* 2007, 6 (11), 841-850.


APPENDICES

Appendix-A. Measurement of nanowire length distribution

The average size of nanowires and its distribution were measured by uploading SEM or optical micrographs into the Image-J application. The analysis employed the distance measuring tool as shown in Figure A. The typical sample size consisted of well over 250 nanowires from each batch.

![Image-J](image.png)

**Figure A.** SEM image and Image-J application depicting the nanowire length measurement technique.
Appendix-B. Measurement of nanowires alignment angle

Figure B. Optical micrographs depicting the nanowire alignment angle measurement using Image-J application for NWs dried in the absence (a), and in the presence of applied magnetic field (b). The red arrow in image-a represents the hypothesized axis and in image-b represents the applied magnetic field direction.

The Cu/Ni nanowires alignment angle distribution was determined by using Image-J application. The procedure involved uploading the optical micrographs into the Image-J
application. The measurement angular distribution was performed with respect to a randomly chosen orientation (in the absence of magnetic field) or applied magnetic field direction. These studies utilized images such as shown in Figure B. Several (>250 NWs) isolated nanowires, with two visible ends, were analyzed for NWs alignment angle distribution and average alignment angles.

**Appendix-C. Issues of using ultra-long (>30 µm) nanowires**

As the length of the Cu/Ni nanowires increases, the flexibility also increases. Especially, the magnetic interactions enhance the bending nature of nanowires resulting in permanently bent nanowire arrangements, see Figure C. The interconnect fabrication requires the nanowires to be deposited as the straight conductive channels. Therefore, for the current study, nanowires with the length of less than 30 µm were used.

![Figure C](image_url)

**Figure C.** Optical micrograph highlighting the flexibility of ultra-long nanowire (shown in the yellow box).