Investigation of Dye-Sensitized Solar Cell
Performance with Single Chirality
Semiconducting Carbon Nanotubes

by

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A Thesis Submitted in Partial Fulfilment of the Requirements for the
Degree of Master of Science in the Materials Science Program in the
Faculty of Science

University of Ontario Institute of Technology

March 2015

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Abstract

This study investigates the performance of dye-sensitized solar cells (DSSCs) assembled with single-walled carbon nanotubes (SWNTs) of a single chirality.

Single-chirality CNTs allow for the adjustment of the Schottky barrier height at the TiO₂/CNT/FTO interface and increases electron diffusion in the photoanode. Furthermore, chirality separated CNTs are expected to absorb less light than mixed chirality CNTs, resulting in better light absorption in the dye and promotion of electron injection into the TiO₂.

Areas of investigation in this study include: DSSC performance comparison between CNT/TiO₂ composites and CNT films, as well as comparison between different chirality-specific and mixed chirality photoanodes. Carbon-based counter-electrode material was also investigated using mixed and chirality-specific films.

Due to their absorption spectra, chiralities (6,5) and (7,3) were chosen and separated by size-exclusion gel chromatography. Characterization of the CNTs and composites included UV-vis-NIR spectroscopy, photoluminescence, XPS and XRD analysis. DSSC characterization includes I-V curves (performed in dark and light) and quantum efficiency. DSSCs demonstrated an improved performance with photoanodes composed of chirality-specific CNT films.
Acknowledgements

First and foremost I would like to offer my sincerest gratitude to my supervisor, Dr. Franco Gaspari, whose encouragement, expertise, and patience has added considerably to my graduate experience. Your guidance and support during my research has been highly valued. Simone Quaranta spent countless hours teaching me how to prepare DSSCs, and listened to me talk (and panic) about my research. He gave me a lot of support and friendship inside and outside of the lab and for that I am truly grateful.

There are many people at UOIT who helped make this project a success. I would like to extend my gratitude to Dr. Liliana Trevani who was always available to provide assistance and answer my many questions. I am sincerely thankful for the advice she gave, as well as generously allowing me to use her equipment and lab space. I would like to thank Dr. Brad Easton for our discussions and for providing resources to help me succeed. I would also like to thank Genevieve Barnes for her generosity. There were many times I needed labware and she was always very kind in lending me equipment. For the SEM and help with Raman spectroscopy, I’d like to thank Ranga Santhanam. I’d also like to thank Dr. Dario Bonetta for allowing me to use his ultracentrifuge.

There are also many from outside of UOIT who were very accommodating and helped me with much needed characterization. I’d like to thank Davit Yeghikyan from UofT for aiding me in the use of their profilometer, as well as Dr. Alex Adronov and Nicole Rice at McMaster University for introducing me to their photoluminescence spectrometer. I would also like to thank the staff at Politecnico di Torino and the Université de Picardie Jules Verne in Amiens for their assistance in characterizing the composites. Additionally, I am extremely grateful to Dr.
Alessandro Latini from La Sapienza Università di Roma for the time and effort he put into the characterization of my DSSCs. He went above and beyond to aid in the completion of this work.

Finally I’d like to thank my parents for all of the support, patience, and encouragement they have given me, as I’ve pursued my academic goals. Without you, this would not have been possible.
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### Abbreviations and Physical Constants

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
<th>Value</th>
</tr>
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<tbody>
<tr>
<td>k(_B)</td>
<td>Boltzmann constant</td>
<td>1.381x10(^{-23}) m(^2) kg/s K</td>
</tr>
<tr>
<td>e</td>
<td>elementary charge</td>
<td>1.602x10(^{-19}) C</td>
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<tr>
<td>h</td>
<td>Planck’s constant</td>
<td>6.620x10(^{-34}) m(^2) kg/s</td>
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<tr>
<td>N</td>
<td>Avogadro’s number</td>
<td>6.02x10(^{23})</td>
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<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
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<tbody>
<tr>
<td>AC</td>
<td>Alternating current</td>
<td></td>
</tr>
<tr>
<td>(a_{cc})</td>
<td>carbon-carbon bond length</td>
<td></td>
</tr>
<tr>
<td>BET</td>
<td>Bruanauer-Emmett-Teller Analysis</td>
<td></td>
</tr>
<tr>
<td>c</td>
<td>Concentration of absorption</td>
<td></td>
</tr>
<tr>
<td>(C_h)</td>
<td>Chiral vector</td>
<td></td>
</tr>
<tr>
<td>CB</td>
<td>Conduction band</td>
<td></td>
</tr>
<tr>
<td>CNT</td>
<td>Carbon Nanotube</td>
<td></td>
</tr>
<tr>
<td>VB</td>
<td>Valence band</td>
<td></td>
</tr>
<tr>
<td>CVD</td>
<td>Chemical Vapour Deposition</td>
<td></td>
</tr>
<tr>
<td>d</td>
<td>Diameter of carbon nanotube</td>
<td></td>
</tr>
<tr>
<td>DC</td>
<td>Direct Current</td>
<td></td>
</tr>
<tr>
<td>DGU</td>
<td>Density Gradient Ultracentrifugation</td>
<td></td>
</tr>
<tr>
<td>DOC</td>
<td>Sodium deoxycholate</td>
<td></td>
</tr>
<tr>
<td>DOS</td>
<td>Density of states</td>
<td></td>
</tr>
<tr>
<td>DSSC</td>
<td>Dye-Sensitized Solar Cell</td>
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</tr>
<tr>
<td>(E_{binding})</td>
<td>Binding energy</td>
<td></td>
</tr>
<tr>
<td>(E_F)</td>
<td>Fermi level</td>
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<tr>
<td>(E_{FC})</td>
<td>quasi-Fermi level for electron distribution in the conduction band</td>
<td></td>
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<tr>
<td>(E_{FV})</td>
<td>quasi-Fermi level for the electron distribution in the valence band</td>
<td></td>
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</table>
$E_g$ \hspace{1cm} \text{Bandgap energy}

$E_{\text{kinetic}}$ \hspace{1cm} \text{Kinetic energy of the electron}

EDS \hspace{1cm} \text{Energy-dispersive X-ray spectroscopy}

FF \hspace{1cm} \text{Fill Factor}

FFF \hspace{1cm} \text{Field-flow fractionation}

FTIR \hspace{1cm} \text{Fourier transform infrared spectroscopy}

FTO \hspace{1cm} \text{Fluorine doped Tin Oxide}

GHG \hspace{1cm} \text{Greenhouse gases}

HEM \hspace{1cm} \text{High energy mode}

HiPCO \hspace{1cm} \text{High-Pressure Carbon Monoxide}

$I_o$ \hspace{1cm} \text{Intensity of the incident light}

$I_{\text{dark}}$ \hspace{1cm} \text{Dark current}

$I_{\text{MP}}$ \hspace{1cm} \text{Current at maximum power point}

$I_{\text{ph}}$ \hspace{1cm} \text{Photocurrent}

$I_{\text{sc}}$ \hspace{1cm} \text{Short-circuit current}

$I_{\text{shunt}}$ \hspace{1cm} \text{Shunt current}

IPCE \hspace{1cm} \text{Incident photon-to-current conversion efficiency}

IR \hspace{1cm} \text{Infrared}

I-V \hspace{1cm} \text{Current-voltage}

$L$ \hspace{1cm} \text{Optical path length}

$m$ \hspace{1cm} \text{Ideality factor}

MWNT/MWCNT \hspace{1cm} \text{Multi-walled Carbon Nanotubes}

$P_{\text{in}}$ \hspace{1cm} \text{Incident solar power}

$P_{\text{max}}$ \hspace{1cm} \text{Maximum power point}

PE-CVD \hspace{1cm} \text{Plasma-Enhanced Chemical Vapour Deposition}

PLE \hspace{1cm} \text{Photoluminescence excitation spectroscopy}

PV \hspace{1cm} \text{Photovoltaic}
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>QE</td>
<td>Quantum efficiency</td>
</tr>
<tr>
<td>R&lt;sub&gt;shunt&lt;/sub&gt;</td>
<td>Shunt Resistance</td>
</tr>
<tr>
<td>RBM</td>
<td>Radial Breathing Mode</td>
</tr>
<tr>
<td>SC</td>
<td>Sodium cholate</td>
</tr>
<tr>
<td>SDBS</td>
<td>Sodium dodecyl benzene sulphonate</td>
</tr>
<tr>
<td>SDS</td>
<td>Sodium dodecyl sulphate</td>
</tr>
<tr>
<td>SEM</td>
<td>Scanning electron microscopy</td>
</tr>
<tr>
<td>SWNT/SWCNT</td>
<td>Single-Walled Carbon Nanotubes</td>
</tr>
<tr>
<td>TEM</td>
<td>Transmission electron spectroscopy</td>
</tr>
<tr>
<td>TGA</td>
<td>Thermogravimetric analysis</td>
</tr>
<tr>
<td>UV-vis-NIR</td>
<td>Ultraviolet-visible-near infrared</td>
</tr>
<tr>
<td>V&lt;sub&gt;MP&lt;/sub&gt;</td>
<td>Voltage at maximum power point</td>
</tr>
<tr>
<td>Voc</td>
<td>Open-circuit voltage</td>
</tr>
<tr>
<td>XPS</td>
<td>X-Ray photoelectron spectroscopy</td>
</tr>
<tr>
<td>XRD</td>
<td>X-Ray Diffraction</td>
</tr>
<tr>
<td>α</td>
<td>Absorption coefficient</td>
</tr>
<tr>
<td>γ&lt;sub&gt;0&lt;/sub&gt;</td>
<td>Tight binding nearest neighbour</td>
</tr>
<tr>
<td>η</td>
<td>Charge Collection Efficiency</td>
</tr>
<tr>
<td>η&lt;sub&gt;inj&lt;/sub&gt;</td>
<td>Electron injection efficiency from dye to TiO&lt;sub&gt;2&lt;/sub&gt;</td>
</tr>
<tr>
<td>θ</td>
<td>Diffraction angle</td>
</tr>
<tr>
<td>ν</td>
<td>Frequency of incident radiation</td>
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<tr>
<td>(a&lt;sub&gt;1&lt;/sub&gt;, a&lt;sub&gt;2&lt;/sub&gt;)</td>
<td>Unit vectors of hexagonal lattice</td>
</tr>
<tr>
<td>(n, m)</td>
<td>Chiral indices</td>
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Chapter 1

Introduction

1.1 Motivation

Since the Industrial revolution of the 18th century, humankind has steadily increased its consumption of fossil fuels and, as a result, has increased the amount of greenhouse gases (GHGs), most notably carbon dioxide, released into our atmosphere. The outcome includes significant changes in climate patterns resulting in extreme weather events at enormous human and financial cost. Furthermore, the world’s population is projected to increase by 1 billion over the next 12 years and reach 9.6 billion by 2050.\cite{1} With emerging markets growing and more people entering the middle class, it is expected that by 2050 world-wide energy use will approximately double, further increasing GHG emissions.\cite{2}

While energy demand is ever-growing, conventional energy resources are dwindling. The issue is not the amount of oil itself; but the amount of high-quality, easily extractable oil. In 2011, 40 out of 54 oil producing nations had peaked in oil production.\cite{3} According to the former head of exploration and production at Aramco, the Saudi national oil company, the international community would need to find a new Saudi Arabia every couple of years to compensate for energy demand, estimated annual growth, and expected annual decline in production from the older fields. The stability of both the biosphere and the global economy will depend on the transition from fossil fuels to a cleaner and more sustainable energy source.\cite{3}
Solar radiation is the single most abundant carbon-neutral energy resource. The amount of energy reaching the surface of the Earth in one hour is equal to approximately $4.3 \times 10^{20}$ J while the amount of primary energy consumed by humans on the planet in the year 2013 was $5.33 \times 10^{20}$ J.[4,5] Clearly solar energy can be part of a viable solution to easing our dependence on fossil fuels.

![Figure 1-1: Record Efficiencies of Different Types of Photovoltaic Cells. From NREL (http://www.nrel.gov/ncpv/images/efficiency_chart.jpg)](http://www.nrel.gov/ncpv/images/efficiency_chart.jpg)

Photovoltaic technology is the fastest growing energy technology with an annual growth rate of approximately 40% in the last two decades.[6] This rate of growth is due to a combination of decreasing prices and the development of devices with improved efficiency (see Figure 1-1). While the price of conventional solar cells, made from crystalline silicon, has dropped, the main disadvantage to them remains that a high purity of silicon is necessary for these cells to perform
acceptably. Unfortunately, many of the emerging photovoltaic technologies, while having lower production costs and efficient manufacturing methods, require the use of toxic materials, such as lead or cadmium (in CdTe, or Perovskite cells), and scarce resources such as indium (such as in CIGS cells). In order for photovoltaics to have a significant and sustainable impact on our future energy resources, the industry needs an efficient and cost-effective cell, easily manufactured, and using non-toxic materials. The dye-sensitized solar cell (DSSC), which is the focus of this work, fulfills these requirements and could therefore have a future as a contributor to the solar cell industry.

1.2 Basics of Photovoltaics

In simple terms photovoltaic (PV) devices are devices that convert the energy from the sun (photons) into electricity. The Photovoltaic Effect is a phenomenon that describes the generation of an electron-hole pair (collectively known as photocarriers or excitons) by photons that are absorbed into a semiconductor. The key element to a solar cell is the absorber, which has an intrinsic band gap to produce the photocarriers.

In the conventional silicon solar cell, two layers are made with one doped with phosphorous and the other doped with boron. Doping the silicon with phosphorous allows “extra” electrons in the structure that are then free to move around (called an n-type layer). On the other hand, doping the other silicon layer with boron allows spaces for electrons to fill, thereby creating holes which also move relatively freely (called a p-type layer). When the n-type and p-type layers are in contact, an electric field forms creating a boundary, or a depletion layer, at the junction (see Figure 1-2 a). The electric field acts like a diode, driving the electrons
to flow from the p-layer to the n-layer. When the device is exposed to sunlight, and therefore an applied bias (see Figure 1-2 b) an electron-hole pair is generated only if the energy of the photon

![Energy band diagram of a p-n junction at thermal equilibrium (V = 0) and under illumination (V > 0)](image)

**Figure 1-2:** a) Energy band diagram of a p-n junction at thermal equilibrium (V = 0) b) Energy band diagram of a p-n junction under illumination (V > 0)

is either equal to or greater than the energy of the bandgap (E_G). Absorption occurs and the electron is excited to the conduction band. Due to the applied bias, there is an increase in the energy of the electrons in the n-type layer, while the Fermi energy of the holes remains unchanged. The result is a split in the Fermi energy otherwise known as the quasi Fermi energy of the electrons (E_{Fn}) and holes (E_{Fp}). Separated by the differences in electrochemical potential in the n-doped and p-doped regions, the hole then travels to the junction, while the electron moves through the n-layer and diffuses to the p-type layer before traveling through the external circuit. Excitons that are not removed at the contacts end up recombining causing either the emission of
a photon or heat. The better the junction between the p-n interface can separate the charge, the higher the voltage produced by the cell. The difference in potential between the two contacts is known as the open-circuit voltage and is given by: [9]

\[ eV_{OC} = E_{Fn} - E_{Fp} \]  

where \( V_{OC} \) is the open-circuit voltage, \( e \) is the elementary charge, \( E_{Fn} \) is the quasi-Fermi level for the electron distribution, and \( E_{Fp} \) is the quasi-Fermi level for the hole distribution.

In summary photovoltaic devices rely on two important steps to ultimately convert sunlight into electrical energy: the successful absorption of radiation resulting in carrier excitation and the continued separation of the carriers as they travel through the cell.

1.3 The Dye-Sensitized Solar Cell

Dye-sensitized solar cells (or DSSCs) are different from classical p-n junction devices in the way they absorb radiation and separate carriers. Unlike conventional cells, which rely on the p-n junction, DSSCs rely on the dye to absorb the photons and generate carriers. The kinetics of the processes involved are responsible for charge separation. The following section discusses the structure of the DSSC and its mechanics in more detail.

1.3.1 DSSC Structure and Mechanics

There are three primary components that make up a DSSC: The working electrode (photoanode), the electrolyte, and the counter electrode (cathode). The working electrode is
conventionally made up of a thin film network of mesoporous titanium dioxide or TiO$_2$ (approx. 5-15 µm thick) covered with a dye monolayer and supported on conducting glass (in our case FTO glass). After exposure to sunlight, the dye (which is usually a metal-organic Ru-complex) becomes excited by the absorption of a photon. When excited, the dye injects an electron through its ligands, to the conduction band of the TiO$_2$ where it then diffuses through the nanostructure TiO$_2$ medium to the conducting glass. Figure 1-3 shows the absorption spectrum of the N719 dye which has two main absorption peaks located at 540 nm and 380 nm (with 380 nm being outside of the visible spectrum). The excited state has a lifetime of nanoseconds before the dye injects the electron into the conduction band of the TiO$_2$. For efficient charge injection, the energy level of the dye would ideally be between 0.2-0.3 V above the TiO$_2$ conduction band edge. Since the TiO$_2$ particles are too small for a macroscopic electric field to build, the main transport mechanism through the anode is by diffusion.

**Figure 1-3:** UV-vis absorption spectrum of N719 in acetonitrile and tert-butanol. Inset is the molecular structure of the N719 dye. (Reproduced with permission from ref [19])
Figure 1-4 demonstrates the key processes of the DSSC. The main role of the TiO$_2$ is to separate and allow photogenerated carriers to travel apart from one another. The electron-hole pair will do one of two things: either they will travel to a reaction site, or recombine. The successful transport of the electron through the nanometer-sized particle network is a compelling hurdle in attaining high efficiency since the DSSC’s performance significantly depends not only on minimizing carrier recombination; but also promoting the forward travel of the electrons.

**Figure 1-4: Schematic of Dye-Sensitized Structure and Mechanics (Reproduced with permission from ref [15])**
Once through the anode the electron flows through the outer circuit to perform electrical work, and returns to the cell via the counter-electrode. At the counter-electrode, the electron is transferred to a hole conducting medium. An iodide/iodine redox couple is used where the reduction is catalyzed by a thin layer of platinum. The reduced species in the electrolyte then diffuse to the oxidized dye and regenerates it in a two-step electron transfer:\textsuperscript{[14,16]}

\begin{align}
2I^- + D^+ &\rightarrow I_2^- + D \\
2I_2^- &\rightarrow I^- + I_3^- \quad (3)
\end{align}

One of the driving forces of the DSSC is the potential difference between the quasi-Fermi level and the redox potential of the electrolyte. In order for the electrons travelling through the circuit to be on a favorable path, and not recombine, the forward reaction must be faster than the backward reaction. Therefore, the regeneration of the dye is faster than the recombination of electrons injected into the conduction band of the TiO\textsubscript{2}.\textsuperscript{[15-18]}

There are many processes in the DSSC that can cause a limitation in the device efficiency; however as mentioned previously, a large part of that limitation is found by the slow electron transport and fast electron recombination in the TiO\textsubscript{2} photoanode. The next section is devoted to a closer look at the photoanode and its role in the DSSC.

1.3.2 The Photoanode

The separation and migration of charge carriers is a key process in the photocatalytic reaction of a DSSC. It is in the photoanode, often called the “working electrode”, where this
process takes place. The main component of the anode is a network of nanostructured TiO$_2$, which serves as a means to separate the carriers by providing a series of pathways for the electron to take when travelling to the conductive glass. The TiO$_2$ layer is typically made up of spherical anatase particles with diameters in the range of 20-30 nm and a band gap of ~3.2 eV.$^{[19]}$

TiO$_2$ is found in nature in four polymorphs (Figure 1-5), three of which are TiO$_2$ crystalline phases: anatase (tetragonal), rutile (tetragonal), and brookite (orthorhombic). The three structures differ by the contrast in each octahedral and by the pattern of the octahedral chains. The rutile structure is assembled by octahedrals that are connected by the edges. In anatase, the octahedrals touch only by their vertices, while in the brookite phase, the vertices and edges are connected.$^{[20]}$ Due to surface free energy and surface stress, anatase is the most thermodynamically stable at small particle sizes (typically of less than 11 nm), brookite being the most stable at sizes between 11 and 35 nm, and rutile is the most stable at larger particle sizes.

In DSSC application, a larger maximum voltage can be obtained from the anatase TiO$_2$ material because its conduction band is 0.2 V more negative than that of the rutile TiO$_2$. Therefore in the photoanode the anatase crystal polymorph is preferred over the rutile. This is due to the surface chemistry of the anatase structure; it also allows better electron mobility and has higher conduction-band energy.$^{[19-21]}$
The size and density of the TiO$_2$ nanoparticles can affect the ease of electron journey. The larger the nanoparticle size, the less defects the nanoparticle has and thus fewer electrons are lost to the electrolyte. On the other hand, the larger the nanoparticle is, the less dye it can be coated with (which is disadvantageous since the dye is the source of electrons for the TiO$_2$). The density of the nanoparticles in the film has a similar effect, where a higher density of nanoparticles yields less surface area to cover with dye; but will provide more paths for the electrons to take to the conductive glass. However, as photogenerated carrier recombination needs to be prevented to enhance efficiency, focusing solely on optimizing the morphology of the TiO$_2$ network is not sufficient.

One possible solution to decrease the amount of carrier recombination is to incorporate carbon nanotubes (CNTs) into the working electrode. Due to their exceptional ability to accept electrons, CNTs can facilitate the transport of electrons, and increase the photoconversion efficiency of the DSSC. When put into contact with the TiO$_2$ network (Figure 1-6), CNTs (which can be insulating, semiconducting, or metallic) could, depending on the electronic structure of a
given CNT, form a Schottky barrier at the TiO₂/CNT interface. TiO₂ is an n-type semiconductor; however, when in contact with a carbon nanotube, electrons from the TiO₂ will move towards the CNT surface, and if the CNT has a lower Fermi level, the CNT may act as an electron sink. This mechanism leaves the TiO₂ behaving similar to a p-type semiconductor as the holes in its valence band travels to the surface.¹²¹ ²⁵ As stated previously, CNTs come in varying electronic types. If a single bandgap of semiconducting CNTs were to be excluded from the mixture, this would allow the opportunity to “tune” the Schottky barrier height at the CNT/TiO₂ junction. Tuning the height of the Schottky barrier would not only encourage the electrons to travel in the forward direction (and therefore increase current collection efficiency in the anode) but it might also result in an increased recombination time between carriers.

**Figure 1-6:** CNT-TiO₂ morphologies: (a) A random mixture of TiO₂ nanoparticles and CNTs, (b) TiO₂ coated CNTs, and (c) CNTs wrapped around TiO₂ nanoparticles. (d) and (e) illustrates how single-walled CNTs may have better inter-phase contact with TiO₂ particles than tubes with multi-walls. (Reproduced with permission from ref [21])

¹ [Ref Number]
² [Ref Number]
1.4 Carbon Nanotubes (CNTs)

Carbon nanotubes (CNTs) are from the fullerene family, which is an allotrope of carbon. Similar to graphite, the chemical bonding of carbon nanotubes is entirely made up of $sp^2$ bonds; however, the rehybridization of bonds is due to its curvature causing three $\sigma$ bonds to shift out of plane. On the other hand, the $\pi$ orbital is delocalized outside the tube making up for the shift. It is due to this rehybridization and the strength of $sp^2$ bonds that CNTs have remarkable electronic, optical, and mechanical properties.\textsuperscript{[25]} The formation of a single wall carbon nanotube can be visualized as a single graphene sheet rolled into a cylinder. Similarly, a multiwall nanotube has one graphene sheet rolled over another. While this description is used to explain the basic structure of CNTs, this is not the way a nanotube actually forms. CNTs actually “grow” (Figure 1-7) from the substrate either from the tip (tip growth), or from the base (root growth).\textsuperscript{[26, 27]} CNT synthesis and deposition is described in section 1.4.1.

\textbf{Figure 1-7: CNT Growth Mechanisms: Tip Growth (top) and Root Growth (bottom)}

\textit{(Reproduced from ref [26])}
1.4.1 Summary of Common CNT Growth Methods

The past two decades saw significant research into efficient and high-yield carbon nanotube growth methods.\textsuperscript{[27]} Conventionally carbon nanotubes can be grown from a carbon source which is either in a pure solid form, or from an organic precursor. The carbon source is broken down in an inert environment

\textit{Physical Processes}

The processes discussed in this section outline the main physical methods of converting carbon precursors into nanotubes. Due to their immense popularity, they are the most widely used processes for nanotube production.

\textit{(i) Arc-Discharge Synthesis}

When CNTs were discovered in 1991,\textsuperscript{[28]} they were observed in the carbon soot on the electrodes from an arc discharge synthesis intended to produce fullerenes. While this technique is considered one of the easiest and most common methods to produce CNTs, it also produces a mixture of soot and catalytic metals which require purification to separate the nanotubes from the crude product. The arc discharge is generated by running a DC current of \textasciitilde100-200 amps at a low voltage of 20 to 30 V between two high-purity graphite rods electrodes, placed end to end with a space of approximately 1 mm between them. This setup (illustrated in Figure 1-8) is kept in an enclosure with an inert atmosphere typically composed of helium or argon at low pressure. The carbon in the negative electrode sublimes due to the high discharge temperatures. The
electric discharge brings the temperature up to \( \sim 6000^\circ C \). During sublimation, the pressure becomes high ejecting carbon atoms from the solid to create plasma between the electrodes. When the atoms travel toward the colder zones, the material accumulates as a small rod-shaped deposit on the counter-electrode. During this process, the electrode is consumed and thus a constant gap between the anode and cathode is kept constant by adjusting the position of the anode. \(^{[29-31]}\)

![Figure 1-8: Schematic of Arc-Discharge Setup (Reproduced with permission from ref [29])](image)

(ii) **Laser Ablation**

In 1996 Smalley et al.\(^{[32]}\) developed a laser ablation technique for the “mass production” of SWNTs. The laser ablation technique is similar to the arc discharge method as it also involves sublimating carbon in a reduced atmosphere resulting in fullerenes or CNTs being produced. The CNTs are prepared by vaporizing graphite rods with a 50:50 catalyst mixture of cobalt and nickel in an inert atmosphere and low pressure. The carbon target is ablated with a focused laser beam of either one of two types: a pulsed laser, or a continuous wave laser. The laser sublimes the
carbon target in a high-temperature chamber resulting in nanotubes developing on the cooler surface as the carbon condenses. This procedure is typically followed by a heat treatment in a vacuum at ~ 1000°C to remove carbon impurities.\[33\]

In the pulsed laser configuration (Figure 1-9), a ND-YAG laser pulse vaporizes a solid target of graphite or graphite/catalyst into a background gas that is flowing through a quartz tube placed inside a tube furnace. The surface of the target rises to about 6000K while at the end of the pulse the temperature drops to 4000K at which point the composite solid material is converted into a vapour that accumulates when placed in an external furnace that is heated to around 800°C. At this temperature CNTs can grow. On the other hand, in the continuous laser configuration, a continuous CO₂ laser is focused on a graphite target heating it to 3000-3500K. In this configuration the target is placed vertically where the inert gas flows from the bottom of the chamber to the top.\[32-34\]

![Figure 1-9: Schematic of Laser Ablation Setup. (Reproduced from ref [33])](image)

Improvements were made to the basic technique by Thess et al.\[35\] and Rao et al.\[36\] using a double beam laser producing a higher purity of approximately 90% and a better structure than tubes grown via arc-discharge.
Chemical Processes

Chemical techniques for growing carbon nanotubes have many advantages in comparison to the physical methods discussed previously. The first advantage is that the reaction process and often the reactor design, is simple and easy to control. The raw materials are abundant and readily available while most processes themselves are fairly cost-effective. These processes are capable of growing the CNTs directly onto substrates, possibly eliminating future steps. Vapour deposition techniques make depositing vertically aligned CNTs possible and high-pressure techniques allow for continuous operation to easily scale up the process. However, these processes come with large amounts of metallic and amorphous impurities which require further purification.

(i) Chemical Vapour Deposition

Chemical vapour deposition (otherwise known as thermal chemical vapour deposition) is a technique for growing nanotubes that involves preparing a substrate with a metal catalyst, generally iron, nickel, or cobalt (Figure 1-10). This method first emerged in 1996 [37] as a solution for large-scale production of CNTs. The CVD technique is called a “gas-phase” or “vapour-phase” technique because the carbon source is a hydrocarbon gas that continuously flows through the system. As it flows, it enters a hot zone that is heated by a furnace or IR lamp, where it attaches itself to a substrate and grows to form nanotubes from either the root or the tip. CNTs are formed as a result of decomposition of hydrocarbon gas which deposits itself onto a substrate and grows from a metal catalyst. CVD is conventionally performed at temperatures
ranging from ~ 700°C to 1200°C. However it has been reported that the minimum temperature in which CNTs can grow (using this method) is around 450°C. \[^{38-40}\]

\[\text{Figure 1-10: Schematic of a Basic CVD Setup (Reproduced from ref [26])}\]

An important aspect to CVD is the type of catalyst used and the substrate prepared. The substrate used defines the nature and type of carbon nanotubes grown. Conventional materials used as a substrate can be silicon, quartz, or even alumina. Similarly, the catalyst used has an effect on the nanotubes’ parameters. Catalysts are typically metal or organometallic nanoparticles containing iron, cobalt, or nickel. The particle size of the catalyst is proven to define the diameter of the tube and therefore the deposition technique of the catalyst should be chosen carefully. \[^{41, 42}\]

\(\text{\textit{(ii) Plasma-Enhanced Chemical Vapour Deposition (PE-CVD)}}\)

The plasma-enhances CVD process (PE-CVD) uses a glow discharge through a high-frequency voltage applied to two electrodes to create plasma from reactive gases (Figure 1-11).
A substrate is placed on the grounded electrode and the reaction gas is supplied from the opposite plate. Silicon or silicon dioxide substrates are typically used with a deposited catalyst material such as iron or nickel. After the catalyst particles are deposited, the CNTs grow from the catalyst by the glow discharge created from a high-frequency power source. The source of carbon is typically derived from a carbon-containing gas such as methane or carbon monoxide. [42, 43]

Figure 1-11: Schematic of a RF-DC Plasma CVD System. (Reproduced with permission from ref [44])

Thermal CVD and PECVD are different techniques in that PECVD is a film formation technique that causes excited species to react with one another in the gas phase. This makes deposition at lower temperatures possible. Thermal CVD requires temperature of 700°C-1200°C which brings two disadvantages. First the high temperature limits the use of certain kinds of substrates. Second, the nanotubes produced are not just randomly oriented and are not perfectly straight. [38] The PECVD technique overcomes these limitations by replacing some of the heat energy with energy from the plasma; this allows gas dissociation and tube formation to occur at
temperatures ranging from $500^\circ C$ to $700^\circ C$. Well aligned CNTs are grown through the adjustment of the geometry of the reactor. Using the electric field in the plasma, vertically or horizontally aligned CNTs are possible by using the electric field to guide the growth. For example, vertically aligned CNTs were accomplished with the use of a radio-frequency (RF) plasma and a direct-current (DC) plasma generated between a RF electrode and a DC cathode (illustrated in Figure 1-11). Without the presence of the plasma, the resulting carbon nanotubes tend to grow in a random formation.

(iii) High-Pressure Carbon Monoxide (HiPCO)

As its name suggests, this method is dependent on the catalytic decomposition of carbon monoxide (CO) under high-pressure (Figure 1-12). The technique was developed at Rice University in 1999 for the large-scale production of small-diameter carbon nanotubes. The catalysts are formed in the gas phase from an organometallic precursor that was introduced into the reactor. The organometallic species decomposes at high temperature creating metallic clusters on supported catalysts. At high temperature and pressure, these clusters act as nucleation sites for SWNTs to grow on. The group at Rice University first used CO with Fe(CO)$_5$ to yield good quality SWNTs at a rate of approximately 450 mg/hr. Bronikowski et al. investigated the effects of various growth parameters with SWNT yield. They varied the CO pressure, catalyst concentration, and temperature to conclude that the main limit of production is due to the gradual accumulation of layers of iron atoms onto the growing nanotubes.
Other variations of the HiPCO process include using a mixture of benzene and ferrocene to react in an environment of flowing hydrogen gas. Another group deposited high quality SWNTs by the pyrolysis of acetylene at 750-1000°C in a float-iron catalyst system. In both methods catalyst nanoparticles are formed through the thermal decomposition of organometallic compounds with the only difference being the orientation of the CVD system.

1.4.2 Structure and Electronic Properties of SWCNTs

As mentioned previously, carbon nanotubes are conceptualized as a single layer of sp² bonded carbon atoms in a honeycomb lattice that have been rolled up into a cylinder and with ends capped as hemispherical fullerenes of the same diameter. The concept of how the CNT is rolled and its resulting parameters are illustrated in Figure 1-13. Due to van der Waals forces, CNTs will often bond to each other to form bundles. The diameter of an SWNT usually ranges from 0.5 nm to 2 nm; however there are synthesis methods that have prepared tubes from 10 nm to 1 cm.
While the length is an important parameter for many electronic applications, the direction in which the graphene sheet is conceptually rolled has a direct effect on the electronic properties of the tube. This direction is described by its chiral vector \( \mathbf{C} \). In terms of developing photovoltaic devices, one important aspect is the ability to tune the band gap over a large range of wavelengths. Nanotubes are conventionally described by a pair of integers \((n, m)\) called the chiral indices (or chirality) which describes the location of each atom in the graphene sheet. The unit cell vectors of the hexagonal lattice are denoted as \( a_1 \) and \( a_2 \). These two parameters define the chiral vector of a given SWNT which in turn can provide the diameter and chiral angle \( \theta \) of a tube exhibiting properties ranging from metallic, to semi-metallic, and semi-conducting. The chiral vector is therefore described as:

\[
\mathbf{C} = na_1 + ma_2 \tag{4}
\]

It follows that the nanotube diameter is thus given by:

\[
d_t = \sqrt{3a_{cc}}(m^2 + mn + n^2)\frac{1}{2\pi} = \mathbf{C}/\pi \tag{5}
\]

where \( \mathbf{C} \) is the magnitude of the chiral vector and \( a_{cc} \) is the carbon-carbon bond length (1.42 Å). \[52\] The angle between the chiral vector and the indices \((n, 0)\) is described as the chiral angle given by:

\[
\theta = \tan^{-1}\left(\frac{m\sqrt{3}}{m + 2n}\right) \tag{6}
\]

In both semiconducting and metallic SWNTs, the dominant optical transitions are defined by their diameter and chiral vector and the bandgap in semiconducting SWNTs vary inversely with diameter.
Specifically, all CNTs with chiral indices such that \( n - m = 3j \) (where \( j = 0 \)) exhibit metallic properties; those with \( n - m = 3j \) (where \( j \neq 0 \)) are considered small gap semiconductors or semi-metallic CNTs. Finally when \( n - m = 3j \pm 1 \), they are large-gap semiconducting CNTs with a band gap around 1 eV for 0.7 nm of diameter. In terms of rolling up a graphene sheet, if the tube is rolled along the chiral vector which is dictated by chiral indices such that \( n = m \), and a chiral angle where \( \theta = 0^\circ \), it is labelled as an “armchair” carbon nanotube (since the carbon atoms in the graphene lattice form a pattern looking like an armchair). If, however, the tube has the chiral indices \((n,0)\) and the chiral angle is equal to \(30^\circ\) it is called a “zigzag” nanotube (since in this case the end of the tube makes a zigzag pattern). Finally, if the graphene sheet is rolled with a chiral angle between \(0^\circ\) and \(30^\circ\), it is simply called a chiral nanotube and has indices where \( m \neq \)
0 or n and n is any positive integer. Optical properties of SWNTs are typically identified using absorption and luminescence spectroscopy which are discussed in the following chapter. [53-55]

In order for the SWNTs to optimally perform in a dye-sensitized solar cell, nanotubes of a particular chirality, and therefore electronic type, need to be separated from the mixture. In the ideal situation, a separation process would result in a population of SWNT species with the same electronic type, diameter, and length.

1.4.3 Review of Chirality Separation Methods

In order for a CNT separation strategy to be effective, the method must be reproducible and iterative (such that further iterations improve the purity of the sample), scalable, non-destructive (such that desired properties are not degraded during the process), compatible for a large range of types of CNTs, and affordable. [57] The following section is a review of post-growth techniques in literature which have been shown to successfully separate CNTs in varying degrees.

Selective Chemistry and Functionalization

Both theoretical and empirical data have shown that SWNTs will interact with some compounds depending on their electronic structure and diameter. Interactions between CNTs and various compounds often result in a functional group attached, or an encapsulation, around the tubes (Figure 1-14) which leads to separation using an assortment of techniques such as chromatography, centrifugation, and electrophoretic separation. [58] Chemistry on CNTs can be
classified into two types: exohedral functionalization and endohedral functionalization. [59] This section will focus more on exohedral functionalization since it is the more selective of the two methods.

Endohedral functionalization (Figure 1-14 E) is a type of modification of CNTs where nanoparticles are inserted into the CNT interior. [60] This method is done in one of two ways: either putting the CNT sample inside a suspension containing nanoparticles so that the sample can be penetrated from inside, [61] or, the CNT sample is filled with a material that reacts and produces nanoparticles which are trapped inside the tube. [62] This method of functionalization is particularly interesting for the development of nanowires and storage applications. For example the creation of nanowires has been done by encapsulating noble metals and metallofullerenes in CNTs. [63]

![Figure 1-14: Methods to chemically functionalize CNTs: A) Covalent functionalization at defect sites or open ends. B) Covalent sidewall functionalization. C) Noncovalent functionalization with surfactants. D) Noncovalent functionalization by polymer wrapping. E) Endohedral functionalization (Reproduced with permission from ref [62])](image-url)
Exohedral functionalization is the modification of the external walls of the CNT (Figure 1-14 A-D). This method is subcategorized into three methods:

1. Noncovalent functionalization: Typically this kind of functionalization is defined by either encapsulation of the CNTs from a surfactant (Figure 1-14 C), or wrapping the CNT with a polymer (Figure 1-14 D). Due to the CNT’s strong π-π interactions, they have poor solubility in water. To make them soluble in water and organic solvents, a variety of polymers and amphiphilic or π-surfactants can be used to encapsulate or be wrapped around the CNT. Not only does this process increase solubility but also aids in individualization after ultrasonication. Conventional surfactants such as sodium dodecyl sulphate (SDS), [65] sodium dodecyl benzene sulphonate (SDBS), [66] and sodium cholate [67] have been studied and have become the standard surfactants in CNT chemistry. These surfactants successfully attach themselves to the CNT surface and not only interact through π-π interactions but their unpolar tail may wrap around the nanotube as well. This concept of wrapping is applied to many synthetic and bio-polymers. A notable example was published by Zheng et al., [68] who reported a method allowing purification of 12 major semiconducting CNT chiralities by wrapping DNA of different sequences. The sequences recognize and enable chromatographic purification of a particular chirality from the mixture. This is accomplished through exhibiting a periodic purine-pyrimidines pattern which undergoes hydrogen bonding to form a two-dimensional sheet. The sheet folds over the tube into a three-dimensional barrel providing the structural basis of CNT selection. To filter the different CNTs out into electronically specific batches, an ultracentrifugation [69] or a chromatographic [70] method is used and is described later in this chapter.
2. Covalent functionalization at defect sites or open ends (Figure 1-14 A): this method uses the defects or open ends of grown CNTs to anchor functional groups. The process of growing carbon nanotubes often results in material with impurities, such as amorphous carbon, fullerenes, and catalyst residuals. \[^{[71]}\] In the case of HiPCO carbon nanotubes, where the residual catalyst particles are encased in a carbon shell, the removal of these impurities without damaging the CNTs is troublesome. \[^{[72]}\] A common method employed is to oxidize the impurities which have a higher reactivity than the 1D sp\(^2\) lattice of the carbon nanotubes. Since the end caps of the CNT are not perfect sp\(^2\) cylinders, the result is a partial oxidation of the material. \[^{[73]}\] Aside from the ends, CNTs have various defects due to topological rehybridization. The process of oxidation usually results in the opening of the sp\(^2\) carbon lattice at the defect sites yielding the formation of oxygen containing functional groups – commonly carboxylic acid. \[^{[74]}\]

Various means of purification and functionalization have been used, including refluxing in nitric and sulfuric acid, \[^{[75]}\] hydrogen peroxide, \[^{[76]}\] and potassium permanganate. \[^{[77]}\] Sonication is often used to expedite the reaction generating a larger amount of functional groups on the defect sites. \[^{[78]}\] These functional groups often act as an anchor for secondary functionalization, which in turn, aid in solubilizing CNTs in polar solvents. \[^{[79, 80]}\] Aside from physical degradation of the carbon nanotubes, the process regularly introduces negative effects on the electronic properties of the tubes as well. \[^{[81]}\]

3. Covalent side-wall functionalization (Figure 1-14 B): Many selective covalent side-wall functionalization techniques have been developed that use diazonium salts to functionalize metallic CNTs. \[^{[82]}\] The use of diazonium as a basis for functionalization
was first performed electrochemically using buckypaper as a working electrode. The reaction was done in situ decomposing the diazonium salts to radicals by the electron uptake of the CNTs. [83] Later, a modification of this reaction was done with water as a solvent. [84, 85] The availability of electrons near the Fermi level in metallic SWNTs encourages bonding interactions leaving the semi-conducting CNTs insoluble. [86, 87] Similarly, nitronium ions, [88] hydrogen peroxide, [89] and carbenes [90] have also been found to be methods of enriching the metallic population of SWNT species; however azomethine ylides (through cycloaddition) showed preference towards semi-conducting tubes. [91] This was achieved by pre-organizing the starting N-oxides on the nanotube surface prior to generating the reactive ylide. Following functionalization, the metallic SWNTs were separated from the functionalized semiconducting SWNTs through inducing the semiconducting tubes to solubilize in the presence of lignoceric acid. [91]

**Ultracentrifugation**

Centrifugation is a widely used process to separate mixtures by using their inertia. The denser components will travel away from the center of rotation resulting in the most dense material building up at the end of the centrifuge tube. When the effective gravitational force on the tube is increased, more of the dense components begin to precipitate and collect at the bottom of the tube. The less dense material is then collected in the supernatant. A very popular method to sort various lengths and diameters of carbon nanotubes is to use ultracentrifugation. This technique conventionally works by dispersing the CNTs into a constant density medium and ultra-centrifuging the CNTs which are then separated according to their sedimentation
coefficient. The problem of relying solely on their sedimentation coefficient is that the sorting results end up convoluted by several physical parameters (length, diameter, etc.). A paper published by O’Connell \textit{et al.} in 2002 \cite{92} reported the separation of individual SWNTs from bundles, ropes, and catalyst impurities by first suspending the CNTs in heavy water and sodium dodecyl sulphate (SDS). They ultra-centrifuged the solution at 122,000g for 4 hours resulting in the individual SWNTs in the supernatant and the rest of the components collected at the bottom.

A density gradient was introduced into the procedure using iodixanol (a non-ionic density gradient medium) which allowed for SWNTs to be sorted by their buoyant density. \cite{93, 94} In density gradient ultracentrifugation (DGU), a density gradient is created in a centrifuge tube before loading the surfactant/SWNT solution. This gradient is created by either layering the medium in discrete steps, or by using a linear gradient generator. During ultracentrifugation, the SWNTs migrate to the region where their density is the same as the density of the gradient (this is called the isopycnic point). This causes layers of homogeneous CNTs to form in the centrifuge tube according to their density which, if the surfactant surrounding each CNT is uniform, will depend on the diameter of the CNT. In Figure 1-15, bands of semiconducting chiralities are visibly distinguishable. Electronic-type sorting was achieved by DGU when using co-surfactant mixtures of sodium cholate (SC) and SDS. \cite{95, 96} This is due to unequal binding of the two surfactants as a function of SWNT polarization, resulting in subtle differences in density depending on the electronic type, bandgap, and diameter of the SWNT. When this technique is done in several iterations, purities approaching 97\% can be achieved.
Sucrose was experimented with as a density gradient medium in work published by Yanagi et al. in 2008. Sucrose is a common medium used in the separation of macromolecules and is more cost effective than iodixanol. It was reported that by lowering the temperature during the ultra-centrifugation process and tuning the concentrations of the co-surfactants, both metallic and semiconducting SWNTs were retrieved in purities of 69% and 95% respectively. Zhao et al. achieved a 97% purity of (6,5) SWNTs by further tuning the encapsulation of the co-surfactants. In this work sodium deoxycholate (DOC) was used to restrict the SDS wrapping around the individual tubes resulting in a change of density dictated by the chirality.

\textbf{Figure 1-15:} SWNT sorting using density gradient ultracentrifugation and its corresponding optical absorbance spectra for each fraction. (Reproduced with permission from ref [95])
Electrophoresis

CNTs have dimensions that are similar to many biomolecules and therefore it follows that techniques used to separate and sort biomolecules can be used to separate and sort carbon nanotubes as well. Electrophoretic separation is one such technique that has been shown to sort SWNTs by their physical and electronic properties. Conventionally a direct current is applied through a gel and the SWNTs are sorted by their relative mobility. The tubes with the smallest molecular weight migrate the fastest allowing for assortment by length. It would seem that this does not lend any electronic separation at all except that during the tip sonication process of debundling the tubes also cuts the tubes. How much the tubes are cut is dependent on the diameter of the tube which indirectly creates a trend between tube length and diameter. [99, 100]

Another approach was developed in 2003 [101] which used an alternating current dielectrophoretic method to separate metallic SWNTs from the semiconducting ones. When a CNT suspension is placed on an array of microelectrodes and under alternating current, an electric field gradient is created prompting the movement of metallic and semiconducting tubes in opposite directions. The metallic tubes were attracted to the array leaving the semiconducting tubes in the solvent. Shin et al. [102] further developed the concept by finding a scalable method to separate SWNTs using a H-shaped microfluidic channel (shown in Figure 1-16). There are two inlets and two outlets to allow two laminar streams, one with the suspended CNTs and the other only a surfactant solution, to flow through the channel. The technique works by using a dielectrophoretic force from an embedded electrode to extract the metallic CNTs from the flow of suspended CNTs to the surfactant solution. This method only allows for the separation of metallic tubes from the semiconducting ones and not the differences in bandgap of the
semiconducting tubes themselves. A more recent approach uses a diaelectrophoretic field-flow fractionation (FFF) technique to sort mixtures of SWNTs by diameter, thereby opening the possibility to use electrophoretic techniques to isolate semiconducting species by their bandgaps. [103]

**Figure 1-16: Dielectrophoresis of carbon nanotubes.** Semiconducting CNTs (white) remain in solution while the metallic CNTs (black) deposit themselves on the electrode array. (Reproduced with permission from ref [101])

**Chromatography**

Using biochemical methods to separate macromolecules, chromatography effectively separates not only semi-conducting nanotubes from metallic, but also different chiralities of semi-conducting species. Ion-exchange chromatography [70] of DNA-wrapped SWNTs (Figure 1-17) has shown a lot of promise in separating individual semiconducting chiralities. In this study, HiPCO SWNTs are individually wrapped with single-stranded DNA in an aqueous solution. After chromatography, CNTs separated by chirality was confirmed via absorption, fluorescence
and Raman spectroscopy. Each strand of DNA contains a different sequence and with each sequence there is affinity for a different chirality of SWNT. Therefore, the quality of SWNT sorting depends on DNA with a sequence of (GT)$_n$ where $n$ is 10 to 45. While the results of this method report a very high purity of single-chirality tubes, removing the DNA strands post-separation remains a challenge. Furthermore the length of the tubes introduces a convolution of results during the ion-exchange chromatography stage of separation.

**Figure 1-17:** A 2D DNA sheet composed of two hydrogen-bonded anti-parallel ATTTATTTATTTT strands wrapped around a (8,4) tube. (Reproduced with permission from ref [68])

Gel filtration, or size-exclusion gel chromatography, is a simple technique that has opened the possibility of scaling up the separation of CNTs. \(^{[104]}\) Separation into semiconducting and metallic fractions was initially done in work by Moshammer et al. \(^{[105]}\) which demonstrated the ability of an amide-functionalized hydrogel to separate a mixture of SWNTs in a solution of sodium dodecyl sulphate (SDS). Liu et al. \(^{[106]}\) made further progress by using several syringes in a vertical column and a single surfactant. After multiple iterations, the group succeeded in
separating 13 different semiconducting chiralities with ranging purities from a mixture of HiPCO tubes. Jain et al., \cite{107} slightly modified that process to separate a relatively pure (6,5) chirality from a mixture of HiPCO tubes in a single iteration.

The gel filtration method separates macromolecules (such as proteins and polymers) by their size in columns of gel beads with small pores. Small molecules spend more time within the pores of the medium and therefore move more slowly through the column than larger molecules. The small particles enter many pores in the gel, equilibrating between the gel and the moving buffer. Meanwhile, medium-sized particles enter only some of the pores in the gel equilibrating between the gel and the moving buffer. Large particles enter very few pores and as a result travel quickly through the column. Prior to loading the column, the CNTs were first suspended in a 2 wt% SDS solution and homogenized in a tip sonicator for 20 hours. In the first columns, tubes with the largest affinity for the gel were removed from the mixed-chirality solution while the latter columns removed tubes with less affinity (Figure 1-18). After the tubes ran through the column, the unbound tubes were washed with a 2 wt% SDS solution and then the tubes were eluted from each syringe independently using a 5 wt% SDS solution. Since CNTs that were trapped in the gel were easily eluted with a higher concentration of surfactant, this suggests that it is the amount of SDS coating the tubes that alters their interaction with the gel. Therefore it is believed that the chirality-dependent separation of the nanotubes originates from the different amounts of SDS micelles on different CNTs of different chirality. It has been reported that the morphology and coverage of SDS on nanotubes is dependent on tube structure. \cite{106-108}
The Kataura group [106] (Liu et al.) explored this concept further by analyzing the relationship between bond curvature and SDS coverage. They found that there is an increasing relationship between the smallest bond curvature radius belonging to a specific chirality and the separation order. Comparing this with the relationship between separation order and diameter, this would indicate that the tubes which interact with the gel the strongest are the tubes with the smallest diameter. The differences in the surface π-electron states of the varying semiconducting chiralities are responsible for the different curvatures. Metallic nanotubes are fully covered in SDS and therefore have a much weaker interaction with the gel.
1.5 Thesis Objective

As discussed previously, DSSCs containing carbon nanotubes form Schottky junctions with TiO$_2$ permitting electron transport to occur and limiting electron trapping inside the nanostructured film. There have been many studies done where single-walled and multi-walled CNTs have been coated by the TiO$_2$ in the anode film. One drawback to this, however, is that rarely are the tubes completely coated leaving carbon exposed to the electrolyte catalyzing the triiodide conversion into iodide. For this reason, creating a film of CNTs and then printing a film of TiO$_2$ on top of the CNT layer may decrease the amount of unwanted reduction in the electrolyte. Another drawback to a TiO$_2$/CNT composite electrode is the inevitable competition between the CNTs and the dye for light absorption. The main objectives can be summarized as follows:

- Implement a chirality separation process of CNTs to obtain single chiralities in a fast and efficient process;
- Minimize the competition for light between the film and the dye by using a specific chirality, and thus a specific bandgap. It is expected that a film where all of the tubes have the same band-gap will facilitate electron transport through the anode and improve the DSSC efficiency; and
- Decrease unwanted reduction in the electrolyte, thus improving DSSC efficiency, by separating the CNT films and the TiO$_2$ layer.

In this study chiralities (6,4), (6,5), and (7,3) were chosen and separated via size exclusion gel chromatography. These particular chiralities were selected as their respective CNT
species have band-gaps in the range of 2.1-2.5 eV and thus have absorption bands that only partially overlap with the N719 dye absorption spectra.
Chapter 2

CHARACTERIZATION METHODS

2.1 Optical Spectroscopy Techniques

Spectroscopy is the study of the interaction between matter and electromagnetic radiation. When radiation interacts with matter, a number of processes can occur, including reflection, absorption, scattering, and fluorescence. Optical absorption spectroscopy measures the absorption of electromagnetic radiation in the subject material.

When light travels through an absorbing medium, its intensity decreases according to the Beer-Lambert law:\[ I(L) = I_o e^{-\alpha L c} \] (7)

where \( I_o \) is the intensity of the incident light, \( \alpha \) is the absorption coefficient, \( L \) is the optical path length, and \( c \) is the concentration of absorption in the material. Measuring the transmitted intensity relative to the incident intensity, the absorption for a particular wavelength can be found by:

\[ A = -\ln\left(\frac{I}{I_o}\right) \] (8)
When discussing the optical properties of carbon nanotubes or titanium dioxide, one is specifically referring to its absorption, luminescence, and Raman scattering mechanisms. In this study, the optical properties of the CNTs are important as the electronic structure of a given nanotube depends on its chirality. Therefore, the optical properties of a tube of a particular electronic structure can provide a wealth of information such as the diameter, chirality, and electronic nature of the sample.\textsuperscript{[110]} The following sections introduce the basic concepts of the optical spectroscopic techniques used.

2.1.1 UV-vis-NIR Spectroscopy

Ultra-violet-visible-near infrared spectroscopy (UV-vis-NIR) is a technique where the incident light is scanned in the ultraviolet, visible, and near infrared regions of the spectrum. It is often used for the characterization of SWNTs, since it measures light absorbed in the region dominated by transitions between the valence and conduction bands.

As previously discussed in Chapter 1, a carbon nanotube can be described as either semiconducting, semi-metallic, or metallic. The energy gap ($E_g$) of the semiconducting CNT is related to the chirality, which is proportional to the inverse of the diameter of the tube.\textsuperscript{[110]} The electronic properties of a quasi-one-dimensional SWNT is described by a continuous electronic density of states (DOS) which divides into a series of peaks, known as van Hove singularities. The carbon atoms have $p$ electrons which yield a large $\pi$-electron system. The optical absorption spectrum in the UV-vis-NIR range shows the characteristic absorption behaviour of SWNTs, due to the transitions between the first ($E_{11}$) and second ($E_{22}$) pairs of singularities in the density of states. Each $(n,m)$ indices exhibits a different set of van Hove singularities in its valence and
conduction bands. Therefore, the optical absorption spectra can be used to identify the diameters and chiralities in a SWNT sample. [111-114]

**Figure 2-1:** Density of electronic states for a single nanotube. Solid arrows represent optical absorption and emission transitions while dashed arrows represent nonradiative relaxation. (Reproduced with permission from ref [55]).

A typical absorption spectrum of a mixed SWNT sample consists of broad absorption peaks: $E_{11}^{M}$, $E_{11}^{S}$, and $E_{22}^{S}$ which represent the first peak in the DOS of the metallic SWNTs, and the first and second transitions in the semiconducting SWNTs respectively. Based on the tight-binding model, the energy of each transition is described by:

$$E_{11}^{S} = \frac{2a_{cc}y_{o}}{d} \quad \text{and} \quad E_{22}^{S} = \frac{4a_{cc}y_{o}}{d} \quad \text{for semiconducting nanotubes}$$  (9)
where $a_{cc}$ is the distance between carbon atoms and $\gamma_0$ is the tight binding nearest neighbour hopping constant. \cite{115}

Absorbance peaks between 800-1600 nm are assigned as the first inter-band transition $v_1 \rightarrow c_1$ in semiconducting SWNTs ($E_{11}^S$). Second inter-band transitions $v_2 \rightarrow c_2$ in semiconducting SWNTs ($E_{22}^S$) are peaks between 500-900 nm. Absorbance peaks $v_1 \rightarrow c_1$ created by metallic SWNTs ($E_{11}^M$) lay between 400 – 600 nm. \cite{116} For the reasons discussed previously, where the absorption peak is located for any transition is dependent on the particular diameter and chirality of the nanotube. Since the transition energy depends on the diameter and electronic properties of the tube, many parameters including the growth method, diameter dispersion, and purity of the tubes can affect the spectra of the tubes making analysis difficult.

2.1.2 Photoluminescence Excitation (PLE) Spectroscopy

If a sample contains more than a single chirality, the peaks in the UV-vis-NIR spectra can be very broad. This becomes accentuated if two species in the same sample have absorption wavelengths that are close together. To identify semiconducting species in the sample, a photoluminescence map can be constructed to accompany the absorption spectra. The figure in the previous section (Figure 2.1) can also be applied to illustrate the processes of absorption and emission of photoluminescence in a DOS diagram for a semiconducting SWNT. To excite the nanotube for PL characterization, transitions involving the second van Hove singularity are
generally used. Photoluminescence excitation spectroscopy (PLE) is used to determine all of the semiconducting chiralities in the sample. This technique records the photoluminescence emission spectra while the excitation wavelength changes. From this one can create a 3D map of the PL intensity vs. the emission and excitation wavelength (Figure 2.2). Each spot in the map represents a single (n, m) species. Since metallic tubes do not luminesce, having a large amount of metallic species in the sample will dilute the semiconducting species causing them to be less intense in the map. Therefore, to produce the desired results, it is important to limit the amount of metallic tubes in the sample.

![Figure 2-2: PLE Map of mixed SWNTs grown via HiPCO method and dispersed in SDS (Reproduced with permission from ref [55]).](image)

"Figure 2-2: PLE Map of mixed SWNTs grown via HiPCO method and dispersed in SDS (Reproduced with permission from ref [55])."
2.1.3 Infrared (IR) Spectroscopy

Infrared spectroscopy is a fast, non-destructive, and fairly cost-effective technique to characterize materials. The infrared portion of the electromagnetic spectrum ranges from 4000 to 400 cm\(^{-1}\). When a molecule is exposed to infrared radiation, it will absorb the radiant energy if it matches the vibrational energy of that molecule. IR radiation causes the excitation in the vibrations of covalent bonds including the stretching and bending modes. Generally speaking the IR spectrum is split into two regions: the group frequency region which lies from 3600 cm\(^{-1}\) to approximately 1250 cm\(^{-1}\) and the fingerprint region which is located from 1200 cm\(^{-1}\) to 600 cm\(^{-1}\). The fingerprint region is useful as small differences in the structure of a molecule results in large changes in the peaks in this region of the absorption spectrum.\(^{[117-119]}\)

IR spectroscopy is a technique mainly used to clarify the structure of a material and determine its attached functional groups. Since individual interatomic bonds may vibrate in several different motions, each bond may absorb at more than a single frequency. While stretching modes typically result in stronger absorption peaks than bending modes, the absorption due to bending vibrations are useful in differentiating between similar bonds.\(^{[119, 120]}\)

Since carbon nanotubes absorb quite strongly in the IR range, not a lot of information is gathered in comparison to other optical spectroscopy techniques. Therefore in CNT characterization, IR spectroscopy is typically used to analyze impurities resulting from the growth process and to identify functional groups covalently attached to the nanotube surface.\(^{[121]}\) The application of IR spectroscopy in the study of CNT surface chemistry provides a means to observe the interactions occurring at the surface during absorption and to determine the structure of the species absorbing the radiation. In an early transmission IR study, two absorption peaks
were recorded: a weak peak at 868 cm\textsuperscript{-1} and a broad asymmetric peak at 1575 cm\textsuperscript{-1}. \cite{122} In similar studies using Fourier-transform IR spectroscopy and different carbon nanotube symmetries, absorption peaks were located at approximately 850 cm\textsuperscript{-1} and 1590 cm\textsuperscript{-1}. It was found that these phonon modes were present regardless of the diameter of the tube studied. \cite{123,124} Work involving the functionalization of CNTs found that the peaks of carboxylic groups absorb at an IR frequency of 1719 cm\textsuperscript{-1}, and create broad bands ranging between 2800 cm\textsuperscript{-1} and 3500 cm\textsuperscript{-1}. \cite{125-127}

### 2.1.4 Raman Spectroscopy

Raman scattering is inelastic scattering of a photon (usually from a laser in the visible, near infrared, or near ultraviolet spectrum) off of a molecule. Most types of scattering are elastic or Rayleigh scattering; however, a small fraction of incident photons are inelastically scattered by interacting with optical phonons. The scattering process can be broken down into three main steps: 1. Absorption of the incident photon with an energy $\hbar \nu$ and consequently exciting an electron; 2. The generation and absorption of a phonon ($\hbar \omega$); and 3. Relaxation of the electron from its excited state and the emission of a photon ($\hbar \nu \mp \hbar \omega$).
Conventionally there are two types of Raman scattering: Stokes scattering and anti-Stokes scattering. As the photon travels through the material there is either a loss or gain in the photon energy. Anti-Stokes Raman scattering occurs when an electron in a molecule is initially in a higher electronic state and, following interaction with the photon, it relaxes to its ground state, resulting in the scattered photon having more energy than the incident photon. Stokes Raman scattering is the opposite case, where the absorption of a photon excites the electron and then it relaxes to still a higher electronic state from where it started. The shift in wavelength of the material provides substantial information on the sample being analyzed. In Raman spectroscopy only the Stokes part of the spectrum is used because it has greater intensity.
Figure 2-4: Raman spectrum of a mixed multi-wall and single-wall carbon nanotubes sample

The Raman spectrum of CNTs can be categorized into three distinguishable features: the radial breathing mode (RBM), the D mode, and the high energy mode (HEM). Each of these features demonstrates different properties of SWCNTs. From these categories two dominant features appear in every Raman spectra of CNT samples: the radial breathing mode, which is usually found in the range of 100-300 cm\(^{-1}\), and the double peak structure found around 1500-1600 cm\(^{-1}\). These bands signify the difference between SWCNTs and other forms of carbon since these components of the Raman spectra are not observed in any other sp\(^2\) bonded carbon material. The RBM band is a low-frequency band that arises from the scattering in the radial breathing mode. This is due to the symmetric in-phase displacements of all of the carbon atoms around the tube. This mode provides some indication of the diameters in the sample, and also identifies whether the tubes are multi or single-walled. In MWNTs, the RBM band can be seen if
the inner tube diameters are very small (less than 5nm usually). The frequency of the RBM ($\omega_{RBM}$) band depends on the diameter of the tube and is expressed by:

$$\omega_{RBM} = \frac{A}{d} + B$$  \hspace{1cm} (11)

where $d$ is the diameter of the tube, $A$ is a proportionality constant, and $B$ is a damping variable caused by the environment around the tube. If the tube is isolated, B is expected to be zero. \cite{130-132}

![Diagram showing atomic vibrations responsible for RBM and G bands](image)

**Figure 2-5:** *Illustration of the atomic vibrations responsible for the RBM and G bands. (Reproduced with permission from ref [128])*

The double peak structure, referred to earlier, is associated with the G-band, tangential vibrational modes of the CNTs, and is associated with the Raman spectra of graphite. The G band in carbon nanotubes is due to the curvature of the graphene sheet and is split into two peaks: the $G^-$ peak and the $G^+$ peak. The $G^-$ peak has a lower frequency and is more visible in semiconducting tubes, as in metallic tube samples it also has a lower intensity and is much broader, making it less visible. The $G^+$ peak, however, is at a slightly higher frequency and has
no significant difference in frequency or width between metallic and semiconducting tubes. In samples of MWNTs where a large portion are metallic, the G-band is visible as a single peak.

The D-band is found at frequencies between 1250 and 1450 cm\(^{-1}\), while its overtone appears at approximately double the frequency ranging between 2500 and 2900 cm\(^{-1}\). The D-band is commonly called the defect band because the intensity of the D band is related to the number of defects in the tube. The approximate intensity and width of the D band can be used to determine the fraction of graphitic material present in the sample.\(^{[131},^{133]}\) A general idea of the quality of nanotubes grown can be obtained by taking the ratio of the G and D bands. Furthermore the intensity of the D band can be used to oversee damage being inflicted on the sample from various processes used in its purification, such as tip sonication or centrifugation.

### 2.2 Surface Characterization Techniques

In some subfields of materials science, understanding the physical and chemical structure of the surface of a material is more important than the bulk of the material. The general methods used to analyze the surface are physical in that the images of the surface are derived from microscopy techniques. These techniques can determine the surface roughness, area, general alignment and porosity of the material. While this information is useful to surface characterization, they provide little information regarding the chemical nature of the material. Spectroscopic methods provide qualitative and quantitative chemical information about a solid from a few angstroms to a few nanometers deep. It is ideal to use both microscopic and spectroscopic techniques in conjunction, as together they provide not only information of the physical surface, but also identification of the chemical species and their concentrations.
2.2.1 Scanning Electron Microscopy (SEM)

A scanning electron microscope is the classical method to see objects at the nanoscale and obtain surface information at a higher resolution than when using regular microscopy. The SEM works by generating a fine beam of electrons from a tungsten filament source or for higher resolution, a field-emission electron gun. The electrons are accelerated to energies between 1-30 keV. The electrons flow in a vertical path through a magnetic condenser and objective lens system reducing the spot size to between 2-10 nm before impinging on the sample surface. Several types of electrons are then produced: backscattered, secondary, and Auger. X-ray fluorescence photons and other photons of various energies are also generated from the process. In SEM, the backscattered and secondary electrons are used to produce the image. The angle at which the electrons strike the surface determines the intensity of the emission of secondary and backscattered electrons. The electrons are ejected from the sample to detectors which convert the electron current into a signal that is sent to a screen producing the final image. The magnification produced by the SEM is a ratio between the field scanning of the sample and the dimensions of the final image. While transmission electron spectroscopy (TEM) creates images with a resolution an order of magnitude higher then SEM techniques, SEM images rely on the surface mechanics rather than transmission and can therefore produce images of bulk samples with a greater depth of view. This results in the production of images that are a good representation of the overall 3D structure of the sample.
A common sample preparation technique consists of coating the sample with a conductive material (such as gold, or platinum) approximately 10 nm thick. This is required as SEM cannot image samples that are not conductive and would otherwise charge the sample. SEM is typically used to analyze the general alignment of CNTs and view them as a bulk sample.  

2.1.5 X-Ray Photoelectron Spectroscopy

X-Ray photoelectron spectroscopy (XPS) is a surface technique used to accurately detect and quantify the elemental composition of a sample. Additionally, it can determine the chemical and electronic state of the elements making up the material’s composition. An XPS spectrum is obtained by exposing a material to a projected beam of aluminum or magnesium x-rays causing

Figure 2-6: Schematic of a Scanning Electron Microscope (Reproduced from ref [136])
electrons to escape from the top 1-10 nm of the sample. Simultaneously the kinetic energy and number of electrons that escape the surface of the material are measured under a very high vacuum. The electrons emitted have kinetic energies proportional to the energy of the incident radiation. Since the energy of the incident ray wavelength is known, the kinetic energy and thus the binding energy of the ejected electron can be found by:

$$E_{kinetic} = h\nu - E_{binding} - \phi$$  \hspace{1cm} (12)

where $E_{kinetic}$ is the kinetic energy of the electron, $h$ is Planck’s constant, $\nu$ is the frequency of the incident radiation, $E_{binding}$ is the binding energy of an electron, and $\phi$ is the work function which is dependent on the spectrometer used. The XP spectrometer only detects electrons that have escaped into the vacuum of the instrument having originated from approximately 10-12 nm into the surface of the material. Electrons that have escaped from deeper inside the sample are either trapped or recaptured in various excited states within the specimen examined. Each element produces a signature of XPS peaks located at characteristic binding energy values specific to that element. High resolution scans of the peaks can provide information regarding the hybridization, bonding, and functional groups in the sample. XPS is a non-destructive technique to investigate the surface chemistry of inorganic compounds, polymers, metal alloys, or semiconductors.  

In CNT/TiO$_2$ composites, XPS is typically used to obtain the structure of the nanocomposite and the chemical compositions on the surface of the nanotubes. Major peaks due to the carbon 1s and oxygen 1s photoelectrons are observed in the scan. Oxygen/carbon
atomic ratios can be calculated to indicate the extent of surface oxidation. The higher the ratio, the more oxygen atoms are bonded to the nanotubes. [142]

In the process of synthesizing TiO$_2$ composites, other titanium oxide impurities can be created which can enhance or hinder the performance of the composite in the photoanode of a DSSC. While TiOx can provide extra electrons to the composite, TiO acts as an electron trap consequently lowering the conduction of the material. [143,144] The XPS technique can then determine the percentage of TiO on the surface of the composite.

2.1.6 Energy-Dispersive X-Ray Spectroscopy

Energy-dispersive X-ray spectroscopy (EDS) is both a qualitative and quantitative analytical technique used for the elemental analysis of a sample. A beam of electrons is focussed on the sample in either a scanning or transmission electron microscope. The electrons from the primary beam interact with the atoms in the sample, which results in two types of X-rays: Continuous (or background radiation) and Characteristic. Characteristic X-rays allow the identification of individual elements in the sample as well as their concentration, based on the intensity of the rays. An energy dispersive detector receives the rays and displays the resulting signal as a spectrum with respect to the X-ray energy. [145,146]
Figure 2-7: Example of an EDS Spectrum of 5.58 at% Au sample. Analysis indicates no detectable impurities since all labeled signals are all due to the presence of gold. (Reproduced with permission from ref [147])

2.3 Other Materials Characterization Techniques

The sections following introduce the basic concepts of other useful characterization techniques that fall under neither the optical nor surface microscopy/spectroscopy categories.

2.3.1 Thermogravimetric Analysis (TGA)

Thermogravimetric analysis (TGA) is an analytical technique in which the mass of a substance is measured as a function of temperature or time. The sample is placed in a controlled environment and the mass is measured as the temperature is increased or decreased at a specified rate. The TGA instrument consists of a sample pan that is supported by a precision balance in a furnace. A purge gas, which can be either inert or reactive, is used to control the environment
and flows through to an exhaust. TGA can determine the liquid content and the presence of volatile species in a sample, the decomposition temperatures of each component, and the rate of thermal degradation.

Combustion of carbon nanotubes typically occurs between 400°C and 640°C. SWNTs with a smaller diameter will combust at lower temperatures due to structural strain. Defects in the nanotube walls will also lower the thermal stability of the sample. Functionalization of CNTs will also alter their thermal stability. Oxidation of CNTs leads to decomposition and reduced thermal stability. [148] In general, measuring the thermal stability of a CNT sample is a method to determine the quality of the nanotube sample, where higher combustion temperatures are associated with better quality CNTs. [149]

In the CNT/TiO₂ composite, performing TGA will determine the weight percent of the CNTs in the composite. Similarly, because the TiO₂ film in the photoanode is made of specific percentages of each of its components, TGA can determine the quantity of TiO₂ in the paste making up the film.

2.3.2 X-Ray Diffraction

Powder X-Ray Diffraction (XRD) is one of the main techniques used to characterize solid-state materials. X-ray diffraction patterns provide information with respect to the crystallinity of a material and the average crystallite size. The X-ray diffraction of a crystal is formulated by Bragg’s law:

\[ 2d \sin(\theta) = n \lambda \] (13)
where \( d \) is the perpendicular distance between pairs of adjacent planes in the crystal, \( \theta \) is the incident angle, \( n \) is the layer of planes, and \( \lambda \) is the wavelength of the X-rays. The distance between adjacent planes can be expressed by cell parameters and the Miller index:

\[
\frac{1}{d^2} = \frac{h^2}{a^2} + \frac{k^2}{b^2} + \frac{l^2}{c^2}
\]  

where \( a, b, \) and \( c \) are the sides of the unit cell; and \( h, k, \) and \( l \) are the Miller indices which describe the lattice planes in the crystal.

**Figure 2-8: Schematic of Bragg’s Law of Diffraction**

In this study XRD is used to characterize TiO\(_2\) and TiO\(_2\) composites. The anatase phase of TiO\(_2\) (101) and the rutile phase (110) are used to determine the average size of the nanocrystallites using Scherrer’s equation: \(^{[150-152]}\)

\[
D_{crys} = \frac{0.9\lambda}{\beta\cos\theta}
\]  

(15)
where $D_{\text{crys}}$ is the average nanocrystallite size (nm), $\lambda$ is the wavelength of the X-ray (1.542 Å), $\beta$ is the full width at half maximum intensity, and $\theta$ is half of the diffraction peak angle.

### 2.3.3 Brunauer-Emmett-Teller Analysis (BET)

Brunauer-Emmett-Teller (BET) theory aims to explain the physical adsorption of gas molecules on a solid surface and is a method for determining the specific surface area of a material. When a solid is exposed to a gas or vapour in a closed space at a definite pressure, the solid begins to adsorb the gas. The adsorption of the gas on the surface of the solid is used to determine the specific surface area of the material by calculating the amount of adsorbate gas corresponding to a monomolecular layer on the surface. Physical adsorption results from van der Waals forces between the adsorbed gas molecules and the adsorbant surface. The amount of absorbed gas is typically carried out at the temperature of liquid nitrogen and measured by a volumetric or continuous flow procedure. Using BET theory, the surface area is calculated from the amount of monolayer gas molecules required to cover the solid surface and the cross-sectional area of the gas molecule being adsorbed. The BET equation is written as:

\[
\frac{P}{v(P_0-P)} = \left( \frac{c-1}{cv_m} \right) \frac{P}{P_0} + \frac{1}{v_m} \tag{16}
\]

where $P$ and $P_0$ are equilibrium and saturation pressures of the adsorbed species, $v$ is the amount of absorbed gas, $v_m$ is the amount of gas adsorbed as the monolayer, and $c$ is the BET constant. The BET equation illustrates an adsorption isotherm as a straight line where $P/v(P_0-P)$ is plotted
on the y-axis and $P/P_0$ is on the x-axis. With the equation written in this way, it is the clear that the slope is $c^{-1}$ and $v_m^{-1}$ is the y-intercept. The slope and y-intercept are used to calculate the amount of monolayer gas, $v_m$, and the BET constant $c$. The BET constant, which is dependent on temperature, is calculated by:

$$c = \exp\left(\frac{E_1 - E_L}{RT}\right)$$

(17)

where $T$ is the temperature, $R$ is Boltzmann’s constant, $E_1$ is the heat of adsorption of the first layer, and $E_L$ is the heat for liquefaction. The total surface area and specific surface area can be calculated by:

$$S_{total} = \frac{v_m \cdot N \cdot s \cdot \Delta x}{M}$$

(18)

$$S_{sp} = \frac{S_{total}}{m}$$

(19)

where $S_{total}$ is the total surface area, $N$ is Avogadro’s number, $s$ is the adsorption cross section, $M$ is the molecular weight of the adsorbed species, $S_{sp}$ is the specific surface area, and $m$ is the mass of the sample analyzed.

Since the surface area and porosity of the TiO$_2$ affects the amount of dye coating and electron transport as well as the introduction of shunt paths to the cathode, BET analysis was done to the hydrothermally synthesized TiO$_2$ to determine these parameters. This was required to decide if thermal treatments of the TiO$_2$ powder caused a large decrease to the particle size.
2.4 Photovoltaic Characterization

The quantum efficiency and current-voltage measurements are the basic characteristics of a solar cell. These techniques are performed under a solar simulator with a radiation intensity of 100 mW/cm\(^2\), or 1 sun, and are measured with respect to a standard reference irradiance distribution: the air mass 1.5 global, or AM 1.5G.

2.4.1 Quantum Efficiency (IPCE)

The current generated at short circuit by a solar cell is dependent on the energy of the incident light. The quantum efficiency (QE) describes the relationship between the incident light and the short-circuit current (I\(_{sc}\)). By definition the quantum efficiency is the probability that an incident photon, with an energy \(E\), will excite an electron to the external circuit. The short-circuit current is given by:

\[
I_{sc} = q \int b_s(E)QE(E) \, dE
\]

where \(b_s(E)\) is the photon flux density of the incident light and \(q\) is the electronic charge. The external quantum efficiency is also called the incident photon-to-current conversion efficiency, or IPCE. In simple terms, the IPCE is a dimensionless quantity which represents the ratio between the amount of electrons generated and the amount of photons incident on the solar cell. [157]
2.4.2 Current-Voltage Measurements (I-V Measurements)

The key performance parameters of a solar cell are determined from the current versus the applied voltage (I-V) characteristics under illumination. When the cell is illuminated, a photocurrent ($I_{ph}$) is generated. When there is no illumination and a forward voltage bias is applied, a dark current ($I_{dark}$), flows in the opposite direction. Charge recombination in the photoactive layer induces a shunting current ($I_{shunt}$) which gives rise to a shunt resistance ($R_{shunt}$). The series resistance of the cell is a summation of the contact resistance at the interfaces, bulk resistance, and the sheet resistance of the electrodes.

By analyzing the I-V curve under illumination (see Figure 2-9), the short-circuit current ($I_{sc}$) can be extracted when $V = 0$ and similarly the open-circuit voltage ($V_{oc}$) can be found when $I = 0$. The maximum power point can be found on the curve where $P_{max}=I_{MP}V_{MP}$. This point defines the condition in which the power output of the device is at its maximum. Conventionally, the solar to electric power conversion efficiency, which is the key parameter of a solar cell, is described in terms of $V_{oc}$ and $I_{sc}$ which requires the introduction of a dimensionless parameter called the fill factor (FF):

$$FF = \frac{V_{MP}I_{MP}}{V_{oc}I_{sc}}$$

(21)

A decrease in the shunt resistance reduces the open-circuit voltage as well as the fill factor. Alternatively an increase in series resistance results in a decrease in the fill factor and the short-circuit current. It is the goal for any solar cell to have the lowest possible series resistance.
to avoid loss of current. The values for both the series and shunt resistance can be found from the slopes of the IV curve at \( V_{oc} \) and \( I_{sc} \) respectively. \[157, 158\]

**Figure 2-9:** Example I-V Curve labelling the maximum power point (MPP) and its associated voltage and current \( (V_{mp}, I_{mp}) \), the open-circuit voltage \( (V_{oc}) \), and the short-circuit current \( (I_{sc}) \).

The last parameter measured from the IV curve under illumination is the energy conversion efficiency which is described by the ratio of the maximum extractable power to the incident solar power \( (P_{in}) \). It is quantitatively expressed as:

\[
\eta = \frac{P_{max}}{P_{in}} = \frac{V_{MP}I_{MP}}{P_{in}} = \frac{FFI_{SC}V_{OC}}{P_{in}} \tag{22}
\]

Two main parameters are derived from the curves generated under dark current. The first is the saturation dark current \( (I_o) \) which is a measure of the recombination in the cell. A cell with
a large amount of recombination will have a large $I_0$ value. For an ideal diode, the dark current density is expressed as:

$$I_{dark}(V) = I_0(e^{qV/k_BT} - 1)$$  \hspace{1cm} (23)$$

where $I_0$ is the saturation dark current, $q$ is the elementary charge constant, $V$ is the voltage, $T$ is the temperature, and $k_B$ is the Boltzmann constant. The saturation dark current is empirically found by taking the ln of the current. Plotting this versus the voltage gives us the linear relationship:

$$\ln(I) = \frac{qV}{k_BT} + \ln(I_0)$$  \hspace{1cm} (24)$$

where $\ln(I_0)$ is the y-intercept.

The last parameter taken from the dark I-V curve is the ideality factor ($m$) which measures how well the device behaves like a diode. A PV device has an ideality factor between 1 and 2; however values higher than 2 are possible due to the presence of non-linearity or a high (ohmic) contact resistance at the metal-semiconductor junction. The ideality factor is calculated from the dark saturation current by taking the diode equation including the ideality factor and again plotting the $\ln(I_{dark})$ versus the voltage:

$$I_{dark}(V) = I_0(e^{qV/mk_BT} - 1)$$  \hspace{1cm} (25)$$

60
\[ \ln(I) = \frac{qV}{mk_BT} + \ln(I_0) \] (26)

The ideality factor can be found by taking a linear fit, finding the slope, and rearranging the slope parameter for \( m \).
Chapter 3

EXPERIMENTAL

3.1 Carbon Nanotube Growth and Preparation

In this study carbon nanotubes produced using high-pressure carbon dioxide (HiPCO) were purchased from Nanointegris. Additionally, samples of carbon nanotubes were grown “in-house” using the chemical vapour deposition method described in the following section. Both of these samples of CNTs were phase separated to remove iron impurities and prepared into an aqueous suspension for gel filtration.

3.1.1 Chemical Vapour Deposition of Carbon Nanotubes

The CNTs were grown via chemical vapour deposition (CVD) using a Barnstead tube furnace, a 121.92 cm long quartz tube with a 5 cm diameter, and a heating coil. A flow meter used to control air flow out of a nitrogen gas cylinder, and a bubbler was arranged with respect to the CVD system, as shown in Figure 3-1. The right end of the heating coil (heated from the variac), was placed 10 cm from the right edge of the tube and over the mixture of camphor and ferrocene. This ensured that the mixture remained inside the tube; but away from the furnace. The left end of the tube was placed 50 cm away from the center of the furnace with the right end about 71cm away. The nitrogen gas cylinder was connected to the tube on the right side, via a
floating ball flow meter. The left side of the tube was connected to the exhaust, via a bubbler, resulting in a flow of air and carbon material from right to left.

**Figure 3-1:** Chemical Vapour Deposition Schematic (top) and Picture (bottom) for the Growth of Carbon Nanotubes
The “hot zone” was controlled by both the variac on the right end of the tube and a microcontroller in the furnace. The microcontroller was programmed to dwell at a set temperature while the variac was slowly increased during the first 10 to 12 minutes of dwell time. The temperature of the heating coil was measured with an IR probe that was connected to a digital temperature meter. The CNTs growth was observed primarily in the area covered by the furnace, with minimal growth occurring outside of the furnace.

Approximately 3 grams of camphor and 2% wt. of ferrocene were each loaded into opposite sides of a ceramic boat which was then inserted approximately 10 cm into the right side of the (clean) quartz tube, and subsequently covered with a heating coil. The furnace was positioned as explained previously with the flanges secured. The tube was then purged with nitrogen gas for 20 minutes before readjusting the gas flow to the furnace to establish a rate temperature rise of 10 °C per minute to a maximum value of 800 °C.

### 3.1.2 Preparation of Carbon Nanotubes Grown by Chemical Vapour Deposition (CVD) and High-Pressure Carbon Monoxide (HiPCO)

Raw CNTs were treated using the organic aqueous phase separation procedure outlined in Tvrdy et al.\textsuperscript{[159]} One gram of raw CNTs, prepared by either high pressure carbon monoxide (Nanointegris), or chemical vapour deposition (UOIT), was placed into a beaker with ~ 20 mL of deionized water. The mixture was then sonicated for 5 minutes in a bath sonicator, and then transferred into a separation funnel. A small amount of hexane was added and the mixture was shaken to produce a phase separation. The procedure was repeated with the addition of small amounts of hexane, followed by shaking the funnel to ensure there were no CNTs left in the
aqueous phase. Periodically the stopcock of the funnel was adjusted to release gas building up in the funnel. After several iterations, the aqueous phase turned a yellow colour due to the presence of impurities from the CNT growth process. The aqueous phase was then removed by draining the fluid into a beaker placed beneath. The organic phase (a mixture of hexanes and purified SWNTs), was allowed to flow into a round-bottom flask. The flask was then placed in a rotary evaporator, under normal vacuum, which in turn was placed inside an oven for approximately 6 hours.

3.1.3 Preparation of an Aqueous Suspension of Carbon Nanotubes

A CNT suspension was created at a rate of 1 mg/mL in an aqueous solution of 2% wt. sodium dodecyl sulphate (SDS, 99%, Sigma Aldrich). The mixture was first placed in an ultrasonic bath for 5 min to break up the large pieces of CNTs before being placed under a tip sonicator (Sonifire 450D, Branson) with a 0.5 inch flat tip at 20 W cm$^{-2}$ for 20 hours. Before sonication, the tip was set 1 cm away from the bottom of the beaker and placed in a temperature-controlled bath at approximately 10 °C.

Immediately following tip sonication the sample was ultracentrifuged at 197,000g for 15 minutes at 10 °C. The top 90% of the supernatant was removed from each ultracentrifuge tube and directly poured into the gel column described in the section below.
3.2 Separation by Chirality: Size-Exclusion Gel Chromatography

Filtration and chirality separation was done in two phases by using the gel chromatographic method outlined by Liu et al.\cite{106} The original method was modified to a single pass to increase the yield and concentration of SWNTs purified for PV application.

3.2.1 General Procedure

The general procedure, introduced and outlined by Liu et al.\cite{106}, is a two-pass process where chirality separation was performed using the “SS-MUGEC” method. In this method a number of 10 mL syringes were connected vertically in series and filled with 1.4 mL of allyl dextran-based size-exclusion gel beads (Sephacryl S-200 HR or S-300 HR, GE Healthcare). An illustration of the separation process is presented in Figure 3-2. After the columns were equilibrated with a 2 wt% SDS solution, 5-10 mL of SWNTs, held in an aqueous suspension and ultracentrifuged, were added to the top column. After the SWNT solution had gone through the column, the column was then taken apart and 6 mL of a 2 wt% SDS aqueous solution was added to each syringe to elute the unbound nanotubes. The semiconducting tubes that were trapped inside the gel were eluted with 4 mL of a 5 wt% SDS solution.

In order to separate the majority of the semiconducting tubes from the metallic ones, the column was equilibrated with 2 wt% SDS solution. After the column was equilibrated, the procedure was repeated by loading the unbound tubes into the column until there were no more semiconducting tubes trapped in the gel leaving only highly enriched metallic tubes.
In order to get an enriched single chirality, a second pass of the previously filtered tubes was required. Prior to secondary separation, each of the samples were diluted to a 2 wt% SDS solution. Again, a number of 10 mL syringes were arranged vertically in a series and equilibrated. The diluted semiconducting fractions were then added to the top column and allowed to flow through the syringes. The column was then disassembled and the unbound tubes eluted with a 2 wt% SDS solution. The semiconducting tubes that were absorbed into the gel were eluted by inserting 4 mL of 5 wt% SDS solution. The result was a solution with an enriched single chirality.

Figure 3-2: Schematic of Separation Process
3.2.2 Modified Procedure: Obtain a Single Chirality in a Single Pass

One disadvantage to requiring two passes through the column is a reduction in CNT concentration following each pass. As the separated CNTs are to be used in a DSSC, it was necessary to obtain as much of the enriched-chirality CNTs in one pass as possible. In our study, we modified the original procedure to maximize the collection of enriched single-chirality in a single pass.

Similar to the original procedure, a number of syringes were vertically arranged in series and plugged with glass wool. The number of syringes arranged depended on the chirality we were targeting to separate from the rest. 1-2 mL of gel was injected into each syringe and equilibrated with 10 mL of 2 wt% of SDS solution. Unbound tubes were eluted with 1.8 wt% SDS solution.

Figure 3-3: Photo of Modified Separation Process
This study examined chiralities (6,5) and (7,3). It was determined that in order to separate the desired chiralities in a single pass, different ratios of gel/SWNTs were required. By injecting 11 mL of SWNT/SDS solution into 1.4 mL of gel/column, into the top column of a 4 column stack, the separation of highly enriched (6,5) in the fourth column occurred. To separate enriched (7,3), 17 mL of SWNT/SDS solution was injected into the top column with 1.4 mL of gel in each syringe. The (7,3) chirality eluted in the first three columns; however the first column yielded the least impurities.

### 3.3 Hydrothermally Synthesized TiO₂ and TiO₂/SWCNT Composite

Two mL of titanium isopropoxide (>97%, Aldrich) was hydrolyzed by adding 6 µL of deionized water. When synthesizing only TiO₂, the solution was immediately transferred to a Teflon-lined acid digestion vessel (45 mL, Parr Instrument) along with 3 mL of 1M H₂SO₄ (95-98%, Sigma-Aldrich). The vessel was then covered and sealed before placing it into a muffle furnace for 24 hours at a temperature of 175 ℃. When preparing the TiO₂/SWNT composite, 3 mg of phase-separated CNTs were added to the solution, and then sonicated in a bath sonicator for 5 minutes. Following sonication the solution was transferred to the vessel with 3 mL of 1M H₂SO₄ and then placed in a muffle furnace at 175 ℃ for 24 hours. For both the TiO₂ and TiO₂ composite, the autoclave was removed from the muffle furnace and left to cool for 2 hours. The autoclave was then opened and the CNT/TiO₂ composite placed in a beaker of deionized water to precipitate. Deionized water was drained out of the beaker and re-filled with fresh deionized water three times before filtering through a Whatmann paper filter (Grade 1, 15 mm, Sigma-
Aldrich) and left to dry. Once dry, the powder was then ground with a mortar and pestle and stored in a dry place.\textsuperscript{[160]}

### 3.4 Dye-Sensitized Solar Cell Fabrication and Assembly

The DSSCs were fabricated using glass purchased with one side coated in fluorine doped tin oxide. The glasses were cleaned using the standard RCA cleaning procedure and those that were to be used as the photoanode were treated with TiCl\(_4\). The anode was prepared following three different procedures: the first involved screen printing the TiO\(_2\) paste onto one of the treated FTO glasses. The second method screen printed the paste made from the CNT/TiO\(_2\) composite onto the treated FTO glass. The third procedure required creating a CNT film (mixed or chirality-specific), then depositing it onto the FTO glass and screen printing the TiO\(_2\) paste on top of the film. The cathode was then prepared and both electrodes sandwiched together and sealed.

#### 3.4.1 Cleaning and Treatment of FTO Glasses

The RCA-1 and RCA-2 cleaning procedures\textsuperscript{[161]} were used to decontaminate the FTO glasses of organic residue and metal ions. These procedures were developed by Werner Kern at RCA laboratories in the late 1960’s and are now considered standard procedures for cleaning silicon wafers. Both processes consist of sequential oxidative desorption and complexing with H\(_2\)O\(_2\)-NH\(_3\)OH-H\(_2\)O in the RCA-1 procedure and with H\(_2\)O\(_2\)-HCl-H\(_2\)O in the RCA-2 procedure. The procedures leave the FTO glass with a thin oxide on the surface. Following RCA-1 and
RCA-2 cleaning, the FTO glasses were treated with 40 mM of TiCl$_4$. The procedures of both cleaning methods and TiCl$_4$ treatment are outlined below.

(i) RCA-1

This method was used to decontaminate the FTO glass from organic residues and films. Five parts water (H$_2$O) and 1 part ammonium hydroxide (NH$_4$OH) were added to a Pyrex beaker and then heated to 70 ± 5 °C. Once the solution reached the desired the temperature, the beaker was removed from the hot plate and 1 part 30% hydrogen peroxide (H$_2$O$_2$) was added. After 1-2 minutes the solution began to bubble vigorously indicating it was ready for use. The FTO glasses were then inserted into the bubbling solution and soaked for 15 minutes after which they were removed and placed into a fresh beaker filled with deionized water. The beaker was then placed under flowing deionized water for a couple of water changes to keep organic residue from re-depositing onto the surface of the FTO glass. The RCA-1 solution was then diluted with cold water and left to cool at the back of the fume hood before disposal.

(ii) RCA-2

This method was utilized to decontaminate metal ions from the surface of the FTO glass. In this procedure 6 parts of deionized water (H$_2$O) was mixed with 1 part hydrochloric acid (HCl) in a Pyrex beaker and heated to 70 ± 5 °C on a hot-plate. Once the desired temperature was reached, the beaker was removed from the hot plate and 1 part hydrogen peroxide (H$_2$O$_2$, 30%) was added. After 1-2 minutes the solution began to bubble indicating it was ready for use.
The FTO glasses were inserted into the beaker and left to soak for 10 minutes. When finished, the glasses were removed and rinsed with deionized water. The RCA-2 solution was left to cool and then poured into a waste container.

(iii) Titanium Tetrachloride (TiCl₄) Treatment

This treatment was utilized to improve the adhesion of the TiO₂ film to the FTO glass and to act as a blocking layer. The procedure required 4 mL of TiCl₄ to be added and stirred in a beaker with 100 mL of deionized water. The solution was then heated to 70 ± 5 °C (any more than this and the TiCl₄ hydrolyzes) and the glasses were left to soak at a constant temperature of 70 °C for 30 minutes. When finished the glasses were removed, washed with deionized water, then washed with ethanol, and then finally blow dried with compressed air. The thin film should be mildly visible when the glass is inspected under light.

3.4.2 Preparation of the Photoanode

(i) SWNT Thin-Film

The films prepared in this work were deposited using a vacuum filtration method: cellulose nitrate membranes with a pore size of 0.2 μm and diameter of 25 mm were used to filter the SWNTs from the surfactant solution. The membranes were clamped to a glass frit and (with the vacuum on) washed with 30 mL of deionized water. The SWNT solution obtained
from the chromatography process was diluted with deionized water to a 1 wt% SDS solution. This decreases the concentration of SDS and SWNTs to help create a more uniform film. The now diluted SWNT/SDS solution was filtrated through the vacuum filtration apparatus. A thin film of CNTs accumulated on the filter surface as the solution passed through the cellulose nitrate membrane. Once all of the solution passed through, the vacuum was left on, and the film was allowed to sit for 15 minutes to set the CNT film. Once the film was dry and set, 1 mL of isopropanol was passed through the filter. The isopropanol decreases surface tension and lowers the likelihood of cracks forming in the film. Approximately 30 mL of deionized water was then passed through the film to wash any remaining SDS in the membrane. After rinsing, the film was allowed to set again by sitting under vacuum for another 15 minutes.

The film was then transferred to the FTO glass by a method originally introduced by Wu et al. First the film was dipped in ethanol and then, applying gentle pressure, the film was pressed (film side down) against the conductive side of the glass. In the original method, outlined by Wu et al., the substrate and film were placed, film side up, into a stirred bath of liquid acetone. However, when this was attempted it was found that the adhesion between the cellulose nitrate filter and the CNT film was larger than that of the film and the substrate. As a result, the film failed to be kept intact as the filter dissolved and it became evident that a gentler method was needed to digest the filter. The subsequent method used was mentioned in Wu et al, where the substrate was placed into an acetone vapour bath instead.
Figure 3-4: Schematic (top) and Photo (bottom) of an Acetone Vapour Bath in a Double Boiler System
In this study, a modified version was performed by creating a double boiler to slowly increase the temperature of the acetone and maintain it at its boiling point. The FTO glass and filter was lowered into the acetone vapour (filter side up) until it was about 2 inches from the liquid. Aluminum foil was used to cover the beaker and to keep the vapour from evaporating. The acetone vapours were allowed to gradually dissolve the cellulose nitrate filter. The filter and substrate were allowed to sit in the vapour bath for approximately an hour and a half to ensure the entire filter was visibly dissolved. The filter was then placed into a stirred bath of liquid acetone for 15 minutes to remove any remaining cellulose nitrate residue. It is important to make sure the entire filter was visibly digested prior to putting the substrate into the liquid bath, as placing the substrate into the bath too soon will result in the thin film breaking apart. Following the liquid acetone bath, the glass was placed into a stirred bath of ethanol for 15 minutes, then removed to dry. After the film dried on the substrate, it was baked for 2 hours at 250 °C in a muffle furnace. The purpose of this last step was to reduce the sheet resistance of the film.

(ii) TiO$_2$ and TiO$_2$/SWNT Composite Paste

The first part of the investigation of DSSC performance compared the CNT/TiO$_2$ composite prepared by hydrothermal synthesis with the TiO$_2$ composite produced by the same method without CNTs. The hydrothermal procedure is outlined in section 2.3 of this document.

The original mixing procedure to produce TiO$_2$ paste for the photoanode was introduced by Ito et al. [164] and was modified in this work to increase the ratio of TiO$_2$ to decrease delamination during heat treatments. The increased ratio of TiO$_2$ resulted in a thicker paste
required only 3-4 layers of screen printing to achieve a film thickness of 15 µm. The paste was made using the procedure outlined in Figure 3-5.

The paste for the TiO\textsubscript{2} photoanode in the chirality specific experiments was made using Aeroxide TiO\textsubscript{2} P90, instead of the hydrothermally synthesized TiO\textsubscript{2}. The paste was produced using the same procedure as the composite paste.

![Figure 3-5: Modified procedure for anode paste preparation](image-url)
(iii) Final Preparation Steps

The majority of anodes were prepared by screen printing the TiO$_2$ paste over the thin film composed of filtered SWNTs onto the FTO glass.$^{[164]}$ For comparison, one anode was prepared with the TiO$_2$ paste screen-printed without a filtered SWNT film underneath, and another prepared with the TiO$_2$/SWNT paste also without the presence of a filtered SWNT thin film. The screen-printing apparatus is shown in Figure 3-6.

![Screen-printing mesh to produce TiO$_2$ films](image)

**Figure 3-6: Screen-printing mesh to produce TiO$_2$ films**

FTO glass was placed under a 1 cm$^2$ square mesh and the paste was spread across the mesh, using a fill blade, leaving a film on the FTO glass. After each layer was printed, the film was allowed to settle on top of a sponge soaked with ethanol until the film visibly relaxed (when the ripples of the film flattened). The first layer relaxed almost immediately and each subsequent
layer took a little longer following each printing. The film was then dried on a hotplate for 5 minutes at 125 °C before the next layer was applied. To get the desired 15 µm thickness, only 3-4 layers were printed before annealing the film. If the anode being prepared had a film of filtered SWNTs underneath a TiO$_2$ film, then the exposed parts of the SWNT film were scratched with glass taking care not to scratch off any of the FTO. Annealing the film was done in stages: 325 °C for 5 minutes in air, 350 °C for 5 minutes in air, 350 °C for 5 minutes under nitrogen, 450 °C for 15 minutes under nitrogen, and then finally 500 °C for 15 minutes also under nitrogen. The furnace temperature was ramped up under air at a rate of 5 °C/min to limit temperature overshooting and then under nitrogen after 350 °C to avoid combustion of the SWNT films. Following the first heat treatment, the glasses were again soaked in a TiCl$_4$ solution at 70 °C for 30 minutes and then rinsed with deionized water and ethanol. Once dry, the films underwent another heat treatment at 350 °C in air for 30 minutes, and then increased to 500 °C under nitrogen for another 30 minutes. When the films cooled to between 70 – 80 °C, they were immediately placed into a bath of 5 mM N-719 dye solution, sealed, and stored in a dark place for 24 hours. When the glasses were removed from the dye, the films were then rinsed with acetonitrile and ready for cell assembly.

### 3.4.3 Preparation of Cathode

Two holes, approximately 0.5 mm in diameter, were drilled into the FTO glass slides about 1.5 cm apart. The glasses were cleaned using the RCA-1 and RCA-2 procedures outlined in section 3.4.1. Using scotch tape, an area of 1 cm$^2$ was masked out on the glass (so that it lined
up with the film on the anode) and platinized with 5 layers of Platisol T paint (Solaronix). The platinum electrode was then treated in air at 450 °C for 15 minutes.

### 3.4.4 Final Assembly of DSSCs

The DSSC was assembled by sandwiching the photoanode and counter-electrode together using a 25 µm Meltonix gasket (Solaronix) to seal the cell together and to keep the two electrodes from touching. A T shape was cut in the center of the gasket with the stem 1 cm² (which lines up with the film) and the sides large enough to go over the holes. This is illustrated in Figure 3-7. When the electrolyte is injected into one of the holes of the counter-electrode, the electrolyte will travel across the film to the other hole.

**Figure 3-7: Looking down on the cell from the photoanode side**

When cutting the gasket to fit the cell, it was ensured that some slack was left on either side of the glasses to ensure that it sufficiently filled the space in between the electrodes. The
glasses were clamped together and placed in an oven at 120 °C for 20 minutes. When the cell is removed from the oven, the gasket should be completely transparent and without bubbles.

The electrolyte was injected into the cell via one of the drilled holes in the cathode. The electrolyte was pulled through the cell, across the film, to the other hole before injecting more electrolyte into the other hole. To seal the electrolyte in the cell, the holes of the cathode were covered with a 60 µm Meltonix gasket (Solaronix) and pieces of thin glass microscope slide were cut and soldered onto the glass. Using an ultrasonic soldering iron, the copper wire contacts were made using a Cerasolzer soldering alloy which specifically adheres to glass.

3.5 Characterization Methods

Experimental procedures for optical, surface characterization and other materials characterization experiments are described in the sections below. Description of how the DSSC cells were characterized using IPCE curves and current-voltage curves under both dark and illumination are also included.

3.5.1 Optical Spectroscopy Techniques

*UV-vis-NIR Spectroscopy*

CNT solutions were analyzed using a Perkin Elmer Lambda 750S UV-vis Spectrometer with a 60 mm Integrating Sphere with wavelengths ranging from 200 nm to 1350 nm. The UV-
Vis resolution of the instrument was from 0.17 to 5.00 nm and the resolution of the NIR ranged from 0.20 to 20.0 nm.

**Photoluminescence Excitation (PLE) Spectroscopy**

PLE spectra were obtained with a Jobin-Yvon SPEX Fluorolog 3.22, equipped with 450 W Xe lamp and an InGaAs detector using a 10 mm quartz cuvette. Slit widths for both excitation and emission were set to a 10 nm band-pass, and correction factor files were applied to account for instrument variations. Photoluminescence maps were obtained at 25°C, with 5 nm intervals in both excitation and emission.

**FTIR Spectroscopy**

IR spectroscopy was done using a Perkin Elmer Spectrum 100 with a Horizontal Attenuated Total Reflectance (HATR) accessory. The sample was spread over the ZnSe crystal and pressed with the force arm attached. Spectra were collected from 400 cm\(^{-1}\) to 4000 cm\(^{-1}\) in 10 accumulations using a resolution of 4 cm\(^{-1}\).

**Raman Spectroscopy**

Raman Spectra were acquired with a Renishaw Raman microscope using an argon laser with a 514 nm wavelength. Sample preparation required dispersing approximately 5 mg of
CNTs into 100 µL of isopropanol. The mixture was placed into a bath sonicator for about 5 minutes to make a dark solution. A drop of the mixture was then placed on a silicon wafer and left to dry for 30 minutes. The microscope was focussed and the spectra were collected at 20 accumulations, with an objective lens setting of 50x, and from 100 nm to 4000 nm.

3.5.2 Surface Characterization

Scanning Electron Microscopy (SEM)

A small amount of CNT sample was placed on a stage and placed in a JEOL 6400 SEM. Once loaded, three or four images were taken of each sample at varying magnifications and accelerating voltages.

Field-Emission Scanning Electron Microscopy (FE-SEM)

The TiO$_2$ and TiO$_2$/CNT composites were characterized using a ZEISS SUPRA 40 Field-Emission Scanning Electron Microscope equipped with a lanthanum boride electron gun. The equipment generates images with a nominal resolution of 1.5 nm at 10 KV and a WD of 2 mm. The composites were first metallized by sputtering chromium, and then placed in the chamber. Images were taken at varying magnifications. The FESEM is equipped with an energy-dispersive X-rays detector.
X-Ray Photoelectron Spectroscopy

XPS experiments were conducted using a Perkin-Elmer, Optica 3100 XL and carried out with a Leybold-Heraeus LHS10 spectrometer using Al Kα (1486.6 eV) radiation. Samples were deposited on a gold sheet and were analyzed in the chamber at a residual pressure of $10^{-9}$ Torr.

3.5.3 Other Materials Characterization Techniques

Thermal Gravimetric Analysis (TGA)

TGA analysis was accomplished using a Q Series SDT Q600 Model (TA Instruments) by oxidizing the samples in air, with a flow rate of 50 mL/min. Samples of approximately 3 mg were gradually heated from 30 °C to 1000 °C at a rate of 5 °C/min. The samples were placed in a small, circular alumina sample pan and loaded into the heating compartment of the instrument.

X-Ray Diffraction

A Panalytical X’PertPRO MPD diffractometer was used with a Cu Kα source ($\lambda = 0.154184$ nm) and an X’Celerator ultrafast RTMS detector. XRD configurations consisted of an angular resolution (in 2θ) of 0.001° and a 0.04 rad soller slit, and a 1° divergence slit. A 20 mm mask was used on the incident beam path, while a 6.6 mm anti-scatter slit and a 0.04 rad collimator were used on the diffracted beam path.
**Brunauer-Emmett-Teller Analysis (BET)**

BET analysis of the TiO$_2$ composites was performed to investigate the relative porosity and surface area of the composites before and after heat treatments. The analysis was completed using Quantachrom Instruments Nova1200e Surface Area and Pore Size Analyzer, and NovaWina software for data acquisition.

**3.5.4 Photovoltaic Analysis**

**Current-Voltage Measurements (I-V Measurements)**

Current-Voltage curves were generated by a 1286 electrochemical Interface from Solartron Analytical, U.K., using full combo Zplot/corrware software by Scribner Associated Inc., U.S.A. During irradiation measurements, the cell was placed underneath an Asahi Spectra HAL-320 Sun Simulator measuring 1 sun at AM 1.5 G. Prior to experiments, calibration was performed with an Asahi Spectra sun checker to verify the intensity of the irradiance to be within ± 1% of 1 sun.

**Quantum Efficiency (IPCE)**

IPCE was measured in DC mode by a custom made apparatus between wavelengths of 400-800 nm and a scan interval of 5 nm. The instrument was made with a 250 W halogen lamp
(Newport-Oriel), a Spectral Products CM110 1/8 m monochromator with variable resolution slits, a Newport broadband beam splitter, a calibrated poly-Si detector, and all of the necessary lenses to focus the beam of light. Data was collected with a National Instruments NI 9219 24-bit universal analog input, and the control software was written by LabView. All of the measurements were done in wavelengths ranging from 400 to 1100 nm with a scan interval of 5 nm.
Chapter 4

Carbon Nanotube Preparation and Separation Results

4.1 Carbon Nanotubes Grown by Chemical Vapour Deposition and High-Pressure Carbon Monoxide

4.1.1 Characterization of CNTs Grown by Chemical Vapour Deposition

Utilizing the procedure outlined in Chapter 3, carbon nanotubes prepared by chemical vapour deposition (CVD) consistently contained aligned forests of nanotubes mixed with CNTs in spaghetti formation and amorphous carbon. Previous work using this growth method has shown that samples prepared under a nitrogen or argon gas flow of around 100 sccm have a larger portion of CNTs growing in straight and uniform bundles. Samples grown under a high gas flow (> 300 sccm) have larger portions of CNTs in a spaghetti formation.

In this study, CNTs were prepared by the CVD method using a mixture of the carbon precursor and catalyst in the solid and vapour phases. Raw samples of the prepared tubes were characterized using SEM, Raman and UV-vis-NIR spectroscopy, and TGA. Similarly, carbon nanotubes prepared by the HiPCO method were also characterized using the same techniques.

SEM images, at varying magnifications, of CNTs grown with mixed precursors in the vapour and solid phases are shown in Figures 4-1 and 4-4. The images in Figure 4-1 indicate the height of the bundled CNTs to be around 40 µm.
Figure 4-1: SEM images of carbon nanotubes grown by chemical vapour deposition where the catalyst and carbon precursor were mixed in the vapour phase. Deposition was done at 800°C, 150 sccm, with a 3.3 wt.% catalyst.

The Raman spectrum in Figure 4-2 reveals the varying kinds of CNTs grown by this technique. RBM bands are observed peaking at 210 cm$^{-1}$, 274 cm$^{-1}$, and 380 cm$^{-1}$. The presence of RBM bands indicates that there are either very small diameter multi-walled tubes or a large
fraction of single-walled tubes. Even though RBM bands are typically associated with single-walled tubes, if there is a small diameter inner tube (less than 2 nm), an RBM can be established at the right resonance condition. RBM signals are not observed from very large diameter tubes because the signal is too weak and the average inner tube in multi-walled tubes will cause the signal to broaden. The two characteristic peaks, namely the D band and G band are located at 1343.5 cm\(^{-1}\) and 1580.3 cm\(^{-1}\) respectively. The D band, otherwise known as the defect band, is due to the scattering of defects that break the symmetry of the graphene sheet. The G band is an intrinsic peak of carbon nanotubes. In primarily semiconducting CNTs, the G peak is split into two components: The G\(^-\) band and the G\(^+\) band. The lack of G band splitting indicates that a large fraction of the nanotubes are metallic species, or multi-walled. The ratio of intensities between the G-band and D-band indicates the quality of the CNT sample. The Raman spectrum

![Raman spectrum](image)

**Figure 4-2:** Raman spectrum of CNTs grown using chemical vapour deposition with the carbon precursor and catalyst mixed in the vapour phase.
depicted in Figure 4-2 has a G to D band ratio of approximately 0.5, indicating a moderate quality sample.

The thermal degradation behaviour was studied using thermo-gravimetric analysis (TGA). The thermal stability of the carbon nanotubes displayed no significant weight loss up to ~500°C with the peak rate of CNT oxidation observed at 604°C, when approximately 71% of the original mass had been lost. At 1000°C, the residual weight of catalyst particles is 6.8%. While the presence of defects lowers the onset temperature of CNT oxidation, the sample contains a significant fraction of large diameter and multi-walled tubes which therefore raises the starting temperature of thermal degradation. The narrow temperature range in which material combusts indicates a smaller fraction of carbonaceous impurities in comparison to other CNT growth techniques.

Figure 4-3: TGA of carbon nanotubes grown by CVD.
Looking at the SEM images in Figure 4-4, CNTs grown with the precursors mixed in the solid phase did not grow as neatly aligned as the ones prepared with the precursors mixed in the vapour phase. Mixing the carbon precursor with the catalyst in the solid phase is not ideal because of the different boiling points of the two substances. Ferrocene vaporizes at 249°C, while the camphor boils at 204°C. This results in an insufficient amount of catalyst particles in the growth environment prior to the ferrocene completely entering the furnace followed by an overabundance of the same catalyst. Having a growth environment high in catalyst particles increases the amount of amorphous carbon around the CNTs which affects the growth alignment.

![SEM images of carbon nanotubes grown by chemical vapour deposition where the catalyst and carbon precursor were mixed in the solid phase. Deposition was prepared at 800°C, 150 sccm, and a catalyst of 3.3 wt. %.

Raman analysis for solid-phase precursor mix samples (see Figure 4-4) indicates RBM bands at approximately 210 cm\(^{-1}\), 276 cm\(^{-1}\), 390 cm\(^{-1}\), 481 cm\(^{-1}\), and 579 cm\(^{-1}\), with 210 cm\(^{-1}\) and 271 cm\(^{-1}\) being the most intense. Again, RBM bands in the spectrum indicate the presence of
SWNTs or very small diameter MWNTs that are under good resonance conditions. The G-band is located at 1572 cm$^{-1}$ and isn’t split into G$^-$ and G$^+$. This indicates large diameters in the mixture with a large fraction of metallic species. A high intensity D band is found at 1343 cm$^{-1}$ indicating that a large portion of the nanotube mixture is amorphous. The ratio of intensities between the D band and G band is approximately 0.71 indicating these tubes are of lesser quality than the ones where the precursors were mixed in the vapour phase.

![Raman Spectroscopy of carbon nanotubes grown by CVD. The precursors were mixed in the solid phase.](image)

**Figure 4-5**: Raman Spectroscopy of carbon nanotubes grown by CVD. The precursors were mixed in the solid phase.

The thermal degradation of the CNT samples prepared with a solid-phase precursor mix begins at 510°C; the rate of combustion peaks at 621°C, resulting in a 74% reduction in weight (see Figure 4-6). The narrower range of temperature indicates a smaller variety of CNTs, with less carbonaceous impurities. The high onset temperature confirms the Raman and SEM data,
revealing a high fraction of multi-walled CNTs and nanotubes with large diameters. The large fraction of multi-walled structures negates the thermal degradation effects of defects in the graphene structure. The residual weight was 10.5% of the original mass loaded into the TGA instrument indicating a high amount of iron catalyst left in the CNT mixture.

![TGA graph](image)

**Figure 4-6:** TGA of CNTs grown by CVD with precursors mixed in the solid phase.

Home grown CNTs were dispersed in a 2 wt% SDS solution and analyzed using UV-vis-NIR spectroscopy. Peaks in the absorption spectra (Figure 4-7) show the presence of semiconducting species at a range of diameters. Narrow peaks indicate the absence of a continuous range of semiconducting tubes in the mixture. The absorption of many of the intense peaks at higher wavelengths indicates a greater fraction of semiconducting tubes with larger diameters.
Figure 4-7: UV-vis-NIR spectroscopy of CNTs grown by CVD dispersed in a 2 wt% SDS aqueous solution.

4.1.2 Characterization of CNTs Grown by High Pressure Carbon Monoxide

SEM images of the tubes prepared via the HiPCO method (Figure 4-8) reveal tubes that are not aligned, but form a mass of tubes that have grown in the spaghetti formation. The tubes measured approximately 600 nm in length, and are therefore longer than those grown by CVD. HiPCO tubes have small diameters ranging from 0.8 to 1.2 nm.
Figure 4-8: SEM images of carbon nanotubes grown by high pressure carbon monoxide.
Figure 4-9 shows that the RBM bands for HiPCO carbon nanotubes are located at 200 cm\(^{-1}\), 242 cm\(^{-1}\), and 263 cm\(^{-1}\). The bands' close proximity confirms the small range in diameters obtained by this growth method. The low intensity of the D band (located at 1330 cm\(^{-1}\)) suggests the nanotubes grown by the HiPCO procedure contain few defects. The G-band is split into \(G^-\) (1529 cm\(^{-1}\)) and \(G^+\) (1587 cm\(^{-1}\)) illustrating a large fraction of small diameter, semiconducting, and single-walled carbon nanotubes. The ratio of intensities between the D and G bands is 0.09, indicating a high quality sample.
The thermal degradation of HiPCO prepared carbon nanotubes was also studied using TGA (Figure 4-10). The CNTs retained thermal stability until approximately 350°C, where the amorphous carbon begins to oxidize. At approximately 450°C, the smaller diameter SWNTs lose thermal integrity. At this temperature approximately 10% of the bulk weight is lost. The largest burning rate of the CNT mixture (the derivative weight peak), occurs at 550°C. The residual weight, composed of iron particles, makes up approximately 3.0%.

*Figure 4-10: Thermogravimetric analysis of HiPCO prepared carbon nanotubes.*
Although it is difficult to make out the individual peaks, the spectra in Figure 4-11 reveal a series of broad peaks in the lower wavelength range, which was not evident in the CVD absorption spectra. This can be explained by the narrower range of diameters in the HiPCO tubes with a smaller variety than for those prepared by CVD. The existence of so many broad peaks in the mid to lower wavelengths confirms Raman spectroscopy data that shows a prevalence of semiconducting carbon nanotube species.

4.2 Carbon Nanotube Purification and Separation

The purity of single-chirality carbon nanotubes is important for a variety of electronic applications. The work presented in this section discusses the results obtained by two procedures reporting single-chirality nanotubes of varying purity. The impurities in each separated aliquot are metallic nanotubes and other semiconducting chiralities. The estimated purities of each of the separated (n,m) species were found using the area underneath each of the peaks in the optical
absorption spectra. Only the separated chirality fractions with the highest purity are presented and used for DSSC application.

Figure 4-12: Photo of separated tubes via size-exclusion gel chromatography. Chiralities separated from vials left to right: (7,3)/(6,4); (6,5)/(6,4); (7,5)/(8,3)/(8,4); and (8,6)/(9,4)

4.2.1 Separation of Carbon Nanotubes Grown by CVD

To obtain some understanding of the distribution of the diameters for the CVD grown CNTs, these were first separated in a single column using 1 mL of Sephacryl gel (200HR). This was done to determine the presence of SWNTs small enough to become caught in the pores of the gel. UV-vis-NIR spectroscopy of the eluted tubes was then performed (Figure 4-13). A purple-blue colour indicated that this growth method was able to produce small diameter tubes; furthermore, separation of small diameter single chiralities from home grown tubes is possible, with some optimization, using gel chromatography. The prominent $E_{11}$ peaks in Figure 4-13 are located at wavelengths 873 nm, 975 nm, 1053 nm, and 1122 nm which correspond to chiralities (6,4), (6,5), (10,2), and (7,6) respectively. Dual or single chirality separation was achieved by
increasing the quantity of initial CNT solution loaded into the column thereby improving the selectivity of the gel.

\[ \text{Figure 4-13: UV-vis-NIR spectrum of 1 mL single column separation of carbon nanotubes grown via chemical vapour deposition.} \]

\[ 4.2.2 \quad \text{Separation of Carbon Nanotubes Grown by HiPCO} \]

Single chiralities were successfully separated following the two-pass procedure written in Chapter 3 and reported in a paper by Liu et al.\textsuperscript{[106]} Figure 4-14 shows the optical absorption spectra of the semiconducting SWNTs after the first separation (left) and second separation (right) resulting in a highly enriched solution containing the (7,3) chirality. The (7,3) chirality has an \( E_{11} \) absorption band located at 990 nm and an \( E_{22} \) absorption band at approximately 510 nm. For DSSC application, the combination of being a relatively simple chirality to isolate, and
the location of its absorption peaks being where they do not overlap the dye, makes the (7,3) species an ideal chirality to target in separation procedures.

![Optical absorption spectra of the two-step separation of the (7,3) chirality. Absorption spectrum of the 1st column is on the left and the absorption spectrum of its second separation is on the right.](image)

**Figure 4-14:** Optical absorption spectra of the two-step separation of the (7,3) chirality. Absorption spectrum of the 1st column is on the left and the absorption spectrum of its second separation is on the right.

As discussed in Chapters 1 and 3, the trapped semiconducting SWNTs are easily desorbed and eluted by loading a higher concentration of surfactant. This indicates that CNTs with greater coverage of SDS have reduced interaction with the gel. In a 2013 paper, the Strano group[159] suggested a model to explain the interactions between SDS and hydrogels that results in the chiral separation of SWNTs made from the HiPCO process. They demonstrate that the separation of single-chirality SWNTs is a kinetically driven forward adsorption process. [159] As SWNTs pass through the stationary bed of ally dextran gel, the chiralities with the largest affinity are adsorbed into the pores of the gel and separated from the bulk solution. After the solution passes through many gel beds, the chiralities with less affinity are adsorbed as well leaving multi-walled, large diameter, and metallic tubes behind. Since a larger micelle reduces
interaction, smaller diameter SWNTs are first captured by the gel. The results presented in this study agree with this trend as the smaller diameter (7,3) chirality is always observed in the 1st column’s absorption spectrum, and mixtures of other chiralities are observed in subsequent columns in order of their diameter.

One drawback to separating by gel chromatography is a limited yield of separated tubes obtained after every pass, as after each pass through the gel, approximately 1% of the tubes are irreversibly adsorbed. \[10^{6}\] This suggests that it would be ideal to limit the number of iterations. In this work we targeted and isolated single chiralities from the bulk SWNT solution by adding enough solution to overload the column in order to obtain a high enough purity in a single pass. This involves using a different loading procedure for each of the chiralities desired. One disadvantage to this method is that the procedure requires optimization for every batch of tubes prepared. See section 3.2.2 for the procedures used to separate enriched (7,3) and (6,5) solutions.

**Figure 4-15: UV-vis-NIR spectrum of single-pass separation of (7,3) and (6,4) chiralities.**
Depending on the amount of CNT solution used to overload the column stack, columns mixed with (6,4) typically eluted from the 2\textsuperscript{nd} column and were either mixed with (7,3) or (6,5). An attempt to isolate (6,4) from the bulk was unsuccessful. To demonstrate a trend in the performance of DSSCs with increasing chirality purity, the (6,4) mixed samples were used in the photoanode of the DSSCs for comparison. Figure 4-15 shows the absorption spectrum of a (7,3)/(6,4) mixture with purities of respectively. In Figure 4-16, the optical absorption spectrum is accompanied by a PL contour map to confirm the mixture of those two chiralities. The calculated purity of the (6,4) and (6,5) is 49.7\% and 45.3\% respectively.

\begin{align*}
(6,4) \, \varepsilon_{11} &= 873 \, \text{nm} \\
(6,5) \, \varepsilon_{11} &= 975 \, \text{nm}
\end{align*}

\textbf{Figure 4-16:} \textit{UV-vis-NIR Spectrum of single-pass separation of (6,5)&(6,4) chiralities. (Left) Photoluminescence map of (6,5)&(6,4) chiralities. (Right)}

Figure 4-17 shows the absorption spectra of the highly-enriched (6,5) solution dispersed in 5 wt.% SDS. A photoluminescence counter map was used to reaffirm the purity of the (6,5) chirality, as in Figure 4-16. Using the UV-vis-NIR spectrum from Figure 4-16, the purity of (6,5) was calculated to be 93\%. Based on the intensity of the metallic range of the UV spectra, both the (6,4)/(6,5) and (6,5) fractions had very little metallic species and few semiconducting
impurities. Based on the absorption spectrum of (7,3) in Figure 4-14, the intensity of the metallic range was quite small in comparison to the semiconducting peaks. In the case of the mixed (6,4)/(7,3) solution, however, there was a comparatively higher absorption intensity in the $M_{11}$ range, indicating a larger presence of metallic species in the sample.

![Absorption spectrum of (7,3) and (6,4)/(7,3) solution](image)

$E_{11} = 975$ nm

**Figure 4-17**: UV-vis-NIR Spectrum of single-pass separation of (6,5) chirality. (Left) Photoluminescence map of separated (6,5) chirality. (Right)

If a specific chirality is desired, the amount of CNT solution loaded can be optimized to increase the selectivity of the gel and target the chirality to elute out of a convenient column. However some chiralities will only separate at a low purity. As there is only a single surfactant used, low purity fractions can be collected for a second pass through the column, and if required, several rounds of separation can be performed until the desired purity is achieved. In 2013, the Kataura group, who published single-chirality results through gel chromatography, introduced temperature-controlled gel chromatography to isolate specific (n,m) species. Instead of relying solely on the interaction between the SDS, SWNTs, and the ally dextran-based gel to elute
desired chiralities, they incorporated temperature control to selectively modulate these interactions. The result is obtaining a larger range of chiralities with a single pass through the gel. The best optimization of this technique would occur if a specific chiral index was present in higher amounts in the starting CNT material (for example the enriched (6,5) nanotubes grown by the CoMoCAT method).
Chapter 5

Characterization of TiO\textsubscript{2} Composites and SWNT Thin Films

5.1 TiO\textsubscript{2} and CNT/TiO\textsubscript{2} Composite

Hydrothermal synthesis involves the growth of crystals under high water vapour pressure and temperature in order to promote nucleation. This results in TiO\textsubscript{2} with a high specific surface area. By adding carbon nanotubes, TiO\textsubscript{2} is chemically linked by functional groups to cover the CNTs. Synthesizing TiO\textsubscript{2} and SWNT composites by hydrothermal functionalization is a useful method as it does not require strong acid/base baths that result in the structural degradation of SWNTs. Milder techniques of functionalization are not favourable alternatives since they have a poor success rate and introduce impurities into the composite that are difficult to eliminate (for example, potassium in potassium persulfate). Similarly, templating agents, (i.e. surfactants) also introduce unwanted species. The high pressure and temperature required in the process creates an environment for intimate contact between the TiO\textsubscript{2} and SWNTs, resulting in improved mixing, less agglomeration, and better coating than the conventional methods of CNT/TiO\textsubscript{2} functionalization.

The surface morphologies of hydrothermally synthesized TiO\textsubscript{2} (see Figure 5-1) and TiO\textsubscript{2}/CNT composites (Figure 5-2) were characterized by FESEM. The resulting images display a surface of TiO\textsubscript{2} with coarse grains and surface areas of varied size. With the addition of carbon nanotubes, the images reveal that there is a moderate amount of agglomeration.
Figure 5-1: Field Emission Scanning Electron Microscopy (FESEM) images of varying magnifications of TiO$_2$ grown using the hydrothermal method.
Figure 5-2: Field Emission Scanning Electron Microscopy (FESEM) images with varying magnifications of a CNT/TiO$_2$ composite grown using the hydrothermal method.

The EDS (see Figure 5-3) of TiO$_2$, synthesized without the presence of carbon nanotubes, reveals elemental compositions of 28.3 wt% of oxygen and a 70.5 wt% of titanium. The remaining 1.2 wt% was the escape peak of silicon, originating from the EDS detector. The
element composition of the TiO$_2$/CNT composite were identified by the EDS as 11.3 wt% of carbon, 46.3 wt% of oxygen, and 42.3 wt% of titanium.

**Figure 5-3:** Energy Dispersive X-Ray Spectroscopy (EDS) of hydrothermally synthesized TiO$_2$ (top) and a CNT/TiO$_2$ composite (bottom).
The carbon on the surface of the TiO$_2$ powder (Figure 5-4) results from the residual species of the synthesis. Carbon bonds formed from the acetate and alkoxide by-products, as well as from the residual isopropoxide in the synthesis. Carbon was an inevitable impurity (otherwise known as adventitious carbon) during XPS as it occurs from exposure to the atmosphere. Adventitious carbon is mainly composed of single and double-bonded (short chain) hydrocarbon species that have peaks identified between 284.5 and 285 eV. In the TiO$_2$/CNT composite (Figure 5-5), the carbon bonds resulted from both the adventitious carbon and the carbon nanotubes incorporated into the composite. The carbonyl group in the XPS from the composite was at a higher percentage than in the XPS of the TiO$_2$ powder due to the functional groups attached to the CNTs.

<table>
<thead>
<tr>
<th>Peak (eV)</th>
<th>%</th>
<th>Bond</th>
</tr>
</thead>
<tbody>
<tr>
<td>C$_{1s}$</td>
<td>284.5</td>
<td>47.6</td>
</tr>
<tr>
<td>285.0</td>
<td>16.9</td>
<td>C-O/C=O</td>
</tr>
<tr>
<td>288.6</td>
<td>7.8</td>
<td>C=O⁻</td>
</tr>
</tbody>
</table>

| Peak (eV) | %   | Bond   |
| Ti$_{2p}$ | 459.7 | 10.7   |
| 462.9    | 8.4   | TiO    |
| 464.3    | 25.3  | TiO$_2$ |
| 466.5    | 1.2   | TiO$_x$ |
The CNT/TiO\textsubscript{2} composite (Figure 5-5) yielded TiO\textsubscript{2} peaks located at 458.4eV and 464.2eV, with no indication of non-stoichiometric phases. This is due to the CNT surface acting as a crystallization seed for TiO\textsubscript{2} (as supported by the XRD data in Figure 5-8). The TiO\textsubscript{2} synthesized by the hydrothermal method (Figure 5-5) indicated the presence of TiO\textsubscript{x} and a very small amount of TiO on the surface. This is both an advantage and a disadvantage, as TiO\textsubscript{x} introduces n-type defects (oxygen vacancies), whereas the TiO acts as an exciton quenching site.

The FTIR spectra in Figure 5-6 and Figure 5-7 revealed peaks located at 1140 and 1735 cm\textsuperscript{-1}, which are CNT and TiO\textsubscript{2}-CNT signatures.\textsuperscript{165} The small peaks occurring near 1735 cm\textsuperscript{-1}...
(in TiO$_2$) and 1745 cm$^{-1}$ (in TiO$_2$/CNT composite) indicated the presence of a –COOH group. Bonding between Ti-O is represented in the approximate region of 650 cm$^{-1}$. Carboxylic groups coordinated with Ti absorb at 1454 cm$^{-1}$ and 1530 cm$^{-1}$ respectively and belong to symmetric and asymmetric stretching modes.\textsuperscript{[166]} The small bands located between 3000 and 2950 cm$^{-1}$ are due to the stretching modes of C-H bonds from the residual isopropoxide and acetate and alkoxide by-products.\textsuperscript{[167]}

*Figure 5-6: FTIR of hydrothermally synthesized TiO$_2$ powder*
Figure 5-7: FTIR of hydrothermally synthesized CNT/TiO$_2$ Composite

Figure 5-8: Raman spectra of TiO$_2$ and TiO$_2$/CNT composite
The anatase form of TiO$_2$ was represented by three peaks in the Raman spectrum (Figure 5-8) at approximately 393, 518, and 642 cm$^{-1}$. The other peaks located at approximately 1300 and 1624 cm$^{-1}$ were the D and G bands of the HiPCO prepared CNTs. The intensity of the D band is higher than the Raman spectrum observed in section 4.1.2 of Chapter 4 due to the introduction of defects during the functionalization process. The intensity of G band splitting and RBM bands were reduced as a result of the weakened signal of the inner-diameter of the tubes and sparse distribution of CNTs. The results of the Raman data were confirmed by XRD. Figure 5-9 compares the XRD patterns of the hydrothermally synthesized TiO$_2$ and TiO$_2$/CNT composite. The peaks at 25.28° (101), 37.80° (100), 48.18° (200), and 54.09° (105) represent the anatase phase of TiO$_2$ and confirmed the absence of other TiO$_2$ phases. The XRD CNT peak was expected at 26.38° (002), however it was barely visible due to the small percentage of CNTs in the composite.

![XRD Patterns of TiO$_2$ and TiO$_2$/CNT Composite](image)

**Figure 5-9: XRD Patterns of TiO$_2$ and TiO$_2$/CNT Composite**
5.2 SWNT Thin Films

Due to the limited yield of the SWNT separation methods, an option that would allow less CNTs to be incorporated into the DSSC was the subject of some investigation. It was determined that not only would a film of SWNTs require less material for assembly, but would produce an interface closer to an ideal Schottky barrier in the photoanode of a DSSC. This section examines transmittance and sheet resistance results of the deposited SWNT films of both mixed and single-chirality.

The transmittance spectra, in Figure 5-10, demonstrated the amount of light that can travel through the film. The transmittance of the (7,3) chirality indicated no absorption peak in the range of maximum absorption of the dye. The other films yielded peaks in the region of the maximum light absorption in the dye. The (7,3) chirality is ideal for DSSC application with dyes that absorb in the IR region (i.e. black dye).
The resistance was lowest for the mixed chirality film due to the presence of metallic tubes. The mixed semiconducting chirality films ((6,5) & (6,4) and (7,3) & (6,4)) measured to have the highest resistance due to the film being composed of two semiconducting chiralities with different bandgaps and the absence of metallic species. The single-chirality films demonstrated a low resistance as the high purity film was composed of a single bandgap.
However, this resistance was not as high as the mixed chirality film, due to the purity of semiconducting species.

**Table 5-1: Sheet Resistance measurements of chirality-specific films.**

<table>
<thead>
<tr>
<th>Chirality</th>
<th>Resistance (Ω)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mixed Chirality</td>
<td>3 695</td>
</tr>
<tr>
<td>(6,4) &amp; (6,5)</td>
<td>51 464</td>
</tr>
<tr>
<td>(6,4) &amp; (7,3)</td>
<td>46 493</td>
</tr>
<tr>
<td>(6,5)</td>
<td>5 992</td>
</tr>
<tr>
<td>(7,3)</td>
<td>5 822</td>
</tr>
</tbody>
</table>
Chapter 6

Dye-Sensitized Solar Cell Performance

6.1 Performance of Cells with TiO$_2$, TiO$_2$/SWNT Composite, and SWNT Thin Film

The short circuit current ($I_{sc}$) is due to the generation and collection of charges. Its value is dependent on the amount of dye absorbed on the TiO$_2$ surface and the injection of electrons from the dye to the TiO$_2$. When the carbon nanotubes are put in contact with the TiO$_2$ surface, the Fermi level in each of the materials equalizes, as observed in standard semiconductor p-n junctions. The equalized Fermi level causes the conduction band of the TiO$_2$ to shift downwards with respect to the LUMO of the dye and the conduction band of the CNTs to shift down with respect to the conduction band in the TiO$_2$. The result is a greater drive for the electrons to travel from the dye to the conduction band of the TiO$_2$ and CNTs.

In Table 6-1 and in Figure 6-1 the solar-cell characteristics of the 3 configurations tested in this work are shown. In particular, a standard TiO$_2$ DSSC performance is compared with a TiO$_2$/CNT composite and a TiO$_2$ + CNT film configuration.
**Figure 6-1:** I-V Curves of DSSCs taken under an irradiance of 1 sun. The DSSC with the photoanode composed of the CNT/TiO₂ composite had the highest fill factor (FF) however the DSSC that performed with the highest efficiency (η) was assembled with separate CNT and TiO₂ films.

**Table 6-1:** DSSC performance parameters of devices made with photoanodes containing TiO₂/CNT composite, separate CNT and TiO₂ films, and TiO₂ alone.

<table>
<thead>
<tr>
<th></th>
<th>η</th>
<th>FF</th>
<th>I&lt;sub&gt;sc&lt;/sub&gt; (mA)</th>
<th>V&lt;sub&gt;oc&lt;/sub&gt; (V)</th>
<th>R&lt;sub&gt;s&lt;/sub&gt; (Ω)</th>
<th>R&lt;sub&gt;sh&lt;/sub&gt; (kΩ)</th>
<th>I&lt;sub&gt;0&lt;/sub&gt; (A)</th>
<th>m</th>
<th>IPCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>TiO₂</td>
<td>1.85%</td>
<td>58%</td>
<td>4.94</td>
<td>0.651</td>
<td>24.80</td>
<td>2.9</td>
<td>7.9 x 10⁻⁷</td>
<td>2.56</td>
<td>15%</td>
</tr>
<tr>
<td>TiO₂/CNT Composite</td>
<td>1.88%</td>
<td>63%</td>
<td>5.40</td>
<td>0.551</td>
<td>19.03</td>
<td>1.1</td>
<td>1.0 x 10⁻⁴</td>
<td>1.80</td>
<td>26%</td>
</tr>
<tr>
<td>TiO₂/CNT Film</td>
<td>2.52%</td>
<td>52%</td>
<td>7.60</td>
<td>0.642</td>
<td>24.62</td>
<td>0.7</td>
<td>1.1 x 10⁻⁵</td>
<td>3.40</td>
<td>47%</td>
</tr>
</tbody>
</table>
A definite improvement can be observed for the DSSC with CNT. Furthermore, it is evident that the short-circuit current improves more with the film than with the composite. The possible reasons for this behaviour are:

1. In the composite, only a portion of the CNTs form a Schottky junction with the TiO₂. As not all of the CNTs are in contact with the TiO₂, layering the TiO₂ and CNTs in the form of a film creates a truer Schottky barrier.

2. There are more CNTs in the composite than there are in the film. This causes more competition for light absorption between the CNTs and the dye thereby lowering the charge injection from the dye to the TiO₂. This difference in light absorption is also reflected in the difference in quantum efficiency (Figure 6-2), where it is doubled in the DSSC made with the film compared to the TiO₂/CNT composite.

The open circuit voltage ($V_{oc}$), found in Figure 6-1, corresponds to the amount of forward bias on the cell stemming from the Fermi level of the TiO₂ and the redox potential of the electrolyte. Adding the CNTs to the photoanode causes the Fermi level of the TiO₂ to fall towards the valence band. This formation of a Schottky junction results in a drop in the open-circuit voltage. Using the CNTs as a film, instead as part of a composite, reduces both recombination and saturation dark current, resulting in an increase in the open-circuit voltage.
Figure 6-2: Quantum Efficiency (IPCE) performance of the DSSCs with TiO$_2$, CNT/TiO$_2$ composite, and CNT Film in the photoanode. Maximum incident photon to charge carrier efficiency for each of the DSSCs occur at 525 nm where the anode with the CNT film performs the best at almost double the photoanode with the CNT composite. The DSSCs with CNTs in the photoanode performed better than the DSSC without.

Since TiO$_2$ is a mesoporous structure, there are grain boundaries which have resistive effects on the DSSC. Defects in the TiO$_2$ will also cause a series resistance in the device. The addition of CNTs, many of which are metallic, increases the 1D transport structure of the electrons. During the preparation of the electrodes, the FTO glasses are treated with TiCl$_4$ (see section 3.4.1 for details). The purpose of this is to improve the adherence between the semiconducting films and the FTO glass by creating a chemical linkage. The deposition of the CNT film on the FTO glass significantly decreases this linkage and consequently decreases the adherence between the TiO$_2$/CNT film and the electrode. The result of this is an increase in the series resistance of the DSSC.
Figure 6-3: I-V Curves measured in a dark environment. The composite shows the largest increase in dark current with an increase in voltage. The DSSCs assembled with CNT and TiO$_2$ layered films and only TiO$_2$ have relatively comparable dark current behaviour.

The shunt resistance ($R_{sh}$) results in significant power loss in the cell by providing alternate paths for the current to travel. In the DSSC, a shunt will allow electrons to travel from the anode film back to the electrolyte, instead of travelling to the external circuit. In DSSCs constructed with CNTs, this means that the tri-iodide in the electrolyte is reduced from the CNTs in the photoanode, instead of the cathode. Shunt pathways were introduced by the technique used to deposit the films since it did not ensure that the CNT film will be compact. This could lead to spaces in the film where CNTs are not adequately covering the FTO electrode. The cell containing only TiO$_2$ in the photoanode has the highest shunt resistance; however there are still shunt pathways introduced between the conductive FTO glass and the electrolyte.
The saturation dark current \( (I_o) \), which is derived from taking the linear fit of the \( \ln(\text{dark current}) \) shown in Figure 6-4, is a measure of the leakage current density from the anode to the electrolyte. It is directly related to thermal excitation and recombination with the electrolyte. The kinetics is described by:

\[
I_o = k[I_3^-][e^-]
\]  

(27)

where \( k \) is the kinetic constant which is based on the speed of reaction, \( [I_3^-] \) is the concentration of tri-iodide, and \( [e^-] \) is the concentration of electrons leaking to the electrolyte. The speed of the reaction is influenced by the number of catalyst sites causing the reduction of tri-iodide. The kinetic constant is high due to the grain boundaries and dangling bonds in the TiO\(_2\). However, this is compensated by the concentration of electrons being kept low by its high band gap of 3.2 eV which makes it difficult for the current to leak to the cathode. The DSSC with the composite had the highest saturation dark current due to a larger percentage of CNTs in the composite. Additionally many of the CNTs were exposed to the electrolyte. Ensuring that the CNTs be completely covered by TiO\(_2\) is still an area of ongoing research and a priority, as the partial covering causes the composite to not behave as a real Schottky junction and produce a lower forward bias. The CNT film has the same issues as the composite in that there are CNTs exposed to the electrolyte; however the amount is much less in comparison to the composite. The incorporation of the CNTs in the form of a film creates an electronic structure closer to that of a real Schottky junction, thereby improving the forward bias and decreasing the recombination current.
**Figure 6-4**: $\ln(I)$-$V$ Curves of DSSCs in a dark environment. Two junctions are evident in the CNT film curve due to the layering in the composition of the anode.

Another parameter that is directly influenced by the recombination in the cell is the ideality factor which is a measure of how well the device behaves like a diode. An ideal diode has an ideality factor of 1, however good solar cells typically have ideality factors between 1 and 2. DSSCs having ideality factors larger than 2 are also frequently observed. The ideality factor of the basic TiO$_2$-DSSC is attributed to a higher order kinetic recombination due to electron traps and defects in the film. The anode film composed of the CNT/TiO$_2$ composite is a first order kinetic mechanism which implies the electrons are travelling directly from the tubes to the electrolyte. However, the value is closer to 2 than to 1 because the Schottky barrier is not ideal resulting in tunnelling and image charge. The photoanode composed of the layered CNT and TiO$_2$ film has a high ideality factor which can be explained by the method used to deposit the CNT film on the electrode. The resulting CNT film was not as uniform as anticipated.
6.2 Performance of Cells with Chirality Separated SWNTs

This section discusses the results from an internal comparison between different SWNT films. The TiO$_2$ was switched from the hydrothermally synthesized composite to the better quality commercial P90. Table 6-2 summarizes the DSSC parameters of cells made with chirality-specific CNT films. Overall, the DSSCs did not perform as well as expected. The main reason for this is the exposure of the CNTs to the electrolyte. The films deposited in this section were much larger than the dimensions of the TiO$_2$ film screen-printed on top, and while there was an attempt to scratch the exposed film off of the conductive glass, care had to be taken to not ruin the electrode or FTO coating. Therefore it was not possible to scratch off all of the exposed film. In the future, depositing CNTs using a spray pyrolysis technique to coat the films, make them more compact, and have them fit the dimensions of the layered TiO$_2$, will significantly decrease the catalysis of the electrolyte from the anode.

While this study does not reveal the best efficiencies, it does report a trend of increasing efficiency with increased chiral selectivity and an improvement over the mixed chirality film. Furthermore, the results found in Figure 6-5 and Table 6-2 indicate which chiralities improve the performance of the anode and which do not.
Figure 6-5: I-V Curves of DSSCs taken under an irradiance of 1 sun. DSSCs assembled with a single-chirality anode had the highest fill factor (FF) and efficiency (η).

Table 6-2: DSSC performance parameters of devices composed of chirality specific SWNT films in the photoanode.

<table>
<thead>
<tr>
<th></th>
<th>η</th>
<th>FF</th>
<th>I_{sc} (mA)</th>
<th>V_{oc} (V)</th>
<th>R_s (Ω)</th>
<th>R_{sh} (Ω)</th>
<th>I_o (A)</th>
<th>m</th>
<th>IPCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mixed</td>
<td>0.54%</td>
<td>32%</td>
<td>5.51</td>
<td>0.307</td>
<td>22.4</td>
<td>31.4</td>
<td>2.2 x 10^{-4}</td>
<td>3.93</td>
<td>7%</td>
</tr>
<tr>
<td>(7,3) &amp; (6,4)</td>
<td>0.28%</td>
<td>30%</td>
<td>3.99</td>
<td>0.235</td>
<td>28.6</td>
<td>53.4</td>
<td>7.3 x 10^{-5}</td>
<td>2.80</td>
<td>6%</td>
</tr>
<tr>
<td>(6,5) &amp; (6,4)</td>
<td>0.24%</td>
<td>33%</td>
<td>2.79</td>
<td>0.263</td>
<td>48.5</td>
<td>352.1</td>
<td>2.0 x 10^{-5}</td>
<td>2.92</td>
<td>8%</td>
</tr>
<tr>
<td>(6,5)</td>
<td>0.74%</td>
<td>47%</td>
<td>4.62</td>
<td>0.340</td>
<td>20.1</td>
<td>110.4</td>
<td>4.5 x 10^{-5}</td>
<td>2.47</td>
<td>12%</td>
</tr>
<tr>
<td>(7,3)</td>
<td>0.98%</td>
<td>48%</td>
<td>5.03</td>
<td>0.374</td>
<td>19.8</td>
<td>122.4</td>
<td>2.4 x 10^{-5}</td>
<td>2.12</td>
<td>15%</td>
</tr>
</tbody>
</table>
The open-circuit voltage shown in Figure 6-5 indicates the Fermi level position of the TiO$_2$ and the redox potential of the electrolyte. If the TiO$_2$ is brought into contact with CNTs of different bandgaps, the Fermi level of the semiconductor will shift either up or down, influencing the open circuit voltage. The mixed film has the largest variation in bandgaps, as well as metallic species. The two-chirality mixed films vary in their purity: the (7,3) & (6,4) film has a greater amount of metallic tubes than the (6,5) & (6,4) film. The single-chirality films: (6,5) and (7,3) have the highest purity in semiconducting tubes, as well as a high single-chirality purity. As reported in Table 6-3, the barrier height decreases with the amount of metallic tubes. Decreasing the barrier height increases the saturation dark current (Figure 6-6 and Figure 6-7) since more electrons can be thermally generated. The mixed film has the shortest height and the purer single-chiralities have the largest. When the TiO$_2$ and single-chirality film come into contact, the Fermi levels equilibrate, causing the Fermi level in the TiO$_2$ to shift upwards. This causes the difference between the conduction band of the TiO$_2$ and the redox potential of the electrolyte to become higher – thus increasing the open-circuit voltage of the DSSC.

**Table 6-3:** Surface area of TiO$_2$ films and estimated Schottky barrier heights at the FTO/TiO$_2$ interfaces influenced by varying chirality films.

<table>
<thead>
<tr>
<th></th>
<th>Surface Area (cm$^2$)</th>
<th>Schottky Barrier Height (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mixed</td>
<td>0.105</td>
<td>0.36</td>
</tr>
<tr>
<td>(7,3) &amp; (6,4)</td>
<td>0.031</td>
<td>0.39</td>
</tr>
<tr>
<td>(6,5) &amp; (6,4)</td>
<td>0.100</td>
<td>0.45</td>
</tr>
<tr>
<td>(6,5)</td>
<td>0.062</td>
<td>0.65</td>
</tr>
<tr>
<td>(7,3)</td>
<td>0.086</td>
<td>0.68</td>
</tr>
</tbody>
</table>
When a metallic and semi-conducting material comes in contact, they typically form a conventional Schottky barrier, where the Fermi level of the semiconducting material shifts to become a constant through the system. When two semiconducting materials come into contact with one another, the Fermi levels of each of the materials shift to equilibrate with one another creating a heterojunction at the interface. There are three types of junctions that could form at a semi-conductor/semi-conductor interface:

1. Straddling gap (Type I): Where the bandgap of one semiconductor is completely contained in the bandgap of the other. This is the most common heterojunction.

2. Staggered gap (Type II): The bandgaps overlap; however either the conduction band of the smaller gapped material might lie above that of the larger gapped material or its valence band might lie below.

3. Broken gap (Type III): If the staggering of the gaps becomes too extreme, they do not overlap at all.

Since the electron affinity of the TiO$_2$ film, or the work function of the individual single and mixed chirality films have not been found, it cannot be known exactly what kind of heterojunction is at the CNT/TiO$_2$ interface in each DSSC. Furthermore, there are inevitable impurities in each of the films including vacancies and grain boundaries in the TiO$_2$, and even small amounts of sodium from the surfactant used to separate the SWNTs. These impurities might cause a “pinning” of the Fermi level in the semiconductor and not a shift as predicted; they also contribute to an unknown type of barrier at the junction between the electrode and the TiO$_2$. 
A linear fit of the I-V curve measured in a dark environment of each DSSC (Figure 6-6) was used to estimate the barrier height and tabulated in Table 6-3.

![Figure 6-6: I-V Curves measured in a dark environment. The (6,5) & (6,4) chirality film produced the smallest increase in dark current with an increase in voltage due to having the highest shunt resistance](image)

The (6,4) chirality incorporated into the dual-mixed chirality films has a comparatively larger band gap than either the (7,3) or (6,5) chiralities. The effect of the larger band gap shifts the Fermi level up, which should increase in theory the open-circuit voltage. However, the barrier height remains low because the (6,4) chirality has its Fermi level close to the conduction band of the TiO₂. The result is less charge collection and more recombination, and therefore a decreased short-circuit current and a lower open-circuit voltage. This concept of the (6,4) chirality has the same effect on the films’ quantum efficiency (in Figure 6-8). The Fermi level is
too high, so the injection from the LUMO of the dye is again not favoured. Therefore it can be concluded that the (6,4) chirality is not a useful material to have in the photoanode.

![Figure 6-7](image)

**Figure 6-7: ln(I)-V Curves of DSSCs in a dark environment. Two junctions are evident in the curves due to the layering composition of the anode.**

The quantum efficiency is low for the mixed chirality anode, for if the Fermi level is high (and the barrier height low), there is a smaller difference between the LUMO of the dye and the conduction band in the TiO₂. The result is the injection of electrons from the LUMO of the dye to conduction and of the TiO₂ to be less favoured, which consequently decreases the quantum efficiency. Similarly, competition between the dye and the carbon nanotubes also decreases the quantum efficiency of the DSSC. The metallic species present in both the mixed film and the (7,3) & (6,4) film demonstrates this effect.
The metallic tubes in the mixed and (7,3) & (6,4) films are responsible for lowering the series resistance in the photoanode. Having a mixture of CNTs with different bandgaps introduces carrier traps in the material which increases the material's resistance. This is clearly observed in the (6,5) & (6,4) film, where the amount of metallic species is low but the semiconducting species with varying bandgaps is high.

In the single chirality films, the open-circuit voltage is expected to be low because the bandgap in these materials is small and the Fermi level is therefore close to the redox potential of the electrolyte. However, the barrier height in these anodes is higher, resulting in improved collection efficiency, higher quantum efficiency, and a smaller saturation dark current thereby.
raising the open-circuit voltage. Since the barrier is higher, the Fermi level of the single chirality films has an energetic distance from the LUMO of the dye (See Figure 1-3). This is why there is better charge injection causing a higher short-circuit current. The better performance of the (7,3) film over the (6,5) film can be explained by the small difference in band gap between the two structures. This difference in band gap translates to less competition for photons between the (7,3) carbon nanotubes and the dye since the absorption spectrum of the dye is removed from the absorption of the (7,3) chirality (see Chapter 5 for transmittance measurements). This results in a higher short-circuit current in the DSSC with the (7,3) anode. This difference in barrier height in the (7,3) structure also results in the shunt resistance being higher as well.

The shunt resistance is lowest for the mixed chirality film due to the presence of metallic tubes causing recombination between the photoanode and counter-electrode. As discussed in chapter 4, the purity of semiconducting tubes in the (7,3) & (6,4) films is lowest and thus there are more metallic tubes than in the rest of the chirality-specific films. This is correlated to the saturation dark current and the shunt resistance. The presence of metallic tubes allows for shunt pathways for the electrons to travel and increased opportunities for recombination. The film made up mainly of (6,4) & (6,5) chiralities has the highest purity of semiconducting tubes and therefore the highest shunt resistance and lowest saturation dark current. As identified previously in Chapter 5, the conductivity of the (6,4) & (6,5) film is also the lowest. This is not only due to the absence of metallic tubes, but also to the mix of the two chiralities in the same film creating electron traps. Both (6,5) and (7,3) films have a high semiconducting nanotube purity; however they are also the most conductive out of the chiral-specific films. So while their shunt resistance is lower (with an higher saturation dark current) than the (6,5) & (6,4) mixed film, there are less
shunt pathways and less recombination than the films with more metallic tubes but yet comparably conductive.

The ideality factor is about 2 in all of the anode materials in this section. The anode resulting in a DSSC performance closest to an ideal diode is the (7,3). None of the DSSCs tested produced an ideality factor between 1 and 2, due to the exposure of CNTs to the electrolyte and the increased amount of shunt pathways.

### 6.3 Cathode Performance of SWNT Film (Mixed and Single Chirality)

Since carbon nanotubes catalyze the electrolyte reaction so well, experiments were done to test how CNT films of mixed and single chirality performed as counter electrodes. As a reference cell, these films were tested and compared against the conventional platinum film. Similar to previous experiments, the photoanodes consist of the commercial P90 TiO₂ and was mixed into a paste and screen printed onto the FTO glass.

As illustrated in Figure 6-9 and recorded in Table 6-4, the short circuit current is highest in the cell with the platinum (Pt) counter-electrode due to the increased conductivity of the Pt. The conductivity of the mixed film is undermined by the presence of tubes with mixed band gaps and while the (6,5) has a fairly high purity of a single chirality, it has a lower conductivity in comparison to the Pt and mixed film. The result is a decreased rate of reduction of the tri-iodide to iodide, causing a slow electron regeneration of the dye. If the rate of injection of electrons from the dye to the TiO₂ is faster than the regeneration of the dye from the electrolyte, it will become more favourable for the dye to regenerate its electrons from the TiO₂ instead. This
hinders the generation of current in the photoanode of the cell and thus lowers the short circuit current.

**Figure 6-9:** I-V Curves of DSSCs taken under an irradiance of 1 sun. The DSSC with the cathode composed of platinum displayed highest fill factor (FF) and performed with the highest efficiency ($\eta$). The tail in the (6,5) counter-electrode is due to the high amount of semi-conducting tubes thus making the contact between the film and the FTO less ohmic. The result is the formation of two junctions seen by a change in the slope of the curve.

**Table 6-4:** DSSC performance parameters of devices with the cathode composed of platinum, mixed chirality carbon nanotubes, and (6,5) carbon nanotubes respectively.

<table>
<thead>
<tr>
<th></th>
<th>$\eta$</th>
<th>FF</th>
<th>$I_{sc}$ (mA)</th>
<th>$V_{oc}$ (V)</th>
<th>$R_s$ ($\Omega$)</th>
<th>$R_{sh}$ ($\Omega$)</th>
<th>$I_o$ (A)</th>
<th>m</th>
<th>IPCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt</td>
<td>5.2%</td>
<td>71%</td>
<td>10.37</td>
<td>0.699</td>
<td>8.7</td>
<td>33792.1</td>
<td>$1.3 \times 10^{-8}$</td>
<td>1.67</td>
<td>27%</td>
</tr>
<tr>
<td>Mixed</td>
<td>3.7%</td>
<td>60%</td>
<td>8.88</td>
<td>0.697</td>
<td>20.0</td>
<td>55958.1</td>
<td>$1.1 \times 10^{-7}$</td>
<td>2.16</td>
<td>30%</td>
</tr>
<tr>
<td>(6,5)</td>
<td>0.73%</td>
<td>24%</td>
<td>7.25</td>
<td>0.417</td>
<td>111.1</td>
<td>111.0</td>
<td>$5.7 \times 10^{-5}$</td>
<td>2.35</td>
<td>15%</td>
</tr>
</tbody>
</table>
The counter-electrodes composed of Pt and mixed chirality films have comparable quantum efficiency (see Figure 6-10). The discrepancy between the quantum efficiency and the short-circuit current is due to the fact that the change in efficiency does not scale linearly with power. The comparable quantum efficiency values are expected between the DSSCs with Pt and mixed films because the parameter primarily depends on the performance of the photoanode, which is composed of only TiO$_2$. The DSSC with the (6,5) cathode is also made up of TiO$_2$ in the photoanode; however the quantum efficiency is quite low in comparison to the other two cells due to the significant difference in the short-circuit current.

![Quantum Efficiency performance of the DSSCs with platinum, mixed chirality films, and (6,5) CNT films. The maximum absorbance occurred at 515 nm with the mixed chirality counter-electrode device having the highest incident photon to conversion efficiency (IPCE) of 30%.

Figure 6-10: Quantum Efficiency performance of the DSSCs with platinum, mixed chirality films, and (6,5) CNT films. The maximum absorbance occurred at 515 nm with the mixed chirality counter-electrode device having the highest incident photon to conversion efficiency (IPCE) of 30%.}
The open-circuit voltage is similar between the mixed film and platinum counter-electrodes indicating good electron injection from the mixed tubes to the electrolyte. The value of the open-circuit voltage is close to the thermodynamic value of approximately 0.7 V, given by the difference between the Fermi level of the TiO$_2$ and the redox level of the electrolyte. The (6,5) however has a much lower open-circuit voltage then the other two cells which is likely due to the high amount of saturated dark current.

There are two kinds of series resistance acting on the counter electrode:

1. The Ohmic resistance.

2. The resistance in charge transfer which does not change linearly with the voltage, and is thus not ohmic. This resistance depends on the process of electron transfer from the counter electrode to the electrolyte. The slower the transfer, the lower the short-circuit current and open-circuit voltage.

The series resistance also affects the fill factor. As the series resistance increases, more power is dissipated causing the fill factor to lower and ultimately lower the short-circuit current. There is a difference in the series resistance of the mixed and (6,5) cathodes due to the presence of semi-conducting tubes and electronic impurities. Based on this, if the mixed film and the platinum are compared, it was observed that the mixed chirality counter-electrode has a higher series resistance; but not as high as the (6,5) only film.
**Figure 6-11:** I-V Curves measured in a dark environment. The (6,5) chirality film shows the largest increase in dark current with an increase in voltage while DSSCs assembled with platinum and mixed chiralities in the counter electrode have relatively comparable dark current behaviour.

The shunt resistance is lowest for the DSSC assembled with a (6,5) chirality counter-electrode due to the lack of compactness in the film. As explained previously in this chapter, the deposition method used makes it hard to control the uniformity of the film. This causes some of the tubes in the counter electrode to make contact with the FTO glass in the anode creating a short and decreasing the shunt resistance.

In Table 6-4 the mixed film has a higher shunt resistance than the (6,5) film despite the similar deposition technique of each of the films. Therefore in this case there must not be a short and yet the saturated dark current is higher than the platinum electrode (see Figure 6-11). Due to the presence of semiconducting tubes in the mixed film, the electron transfer between the counter electrode and the electrolyte is not as efficient as the platinum electrode and therefore does not
catalyze the electrolyte as effectively. Again this results in a reduced rate of reduction in the tri-iodide to iodide which then negatively effects the regeneration the dye. Therefore, regardless of the high shunt resistance in the mixed film DSSC, the decreased rate in which the tri-iodide becomes reduced makes it easier for the dye to take electrons back from the TiO$_2$ thereby increasing the saturation dark current.

![Graph](image.png)

**Figure 6-12**: $\ln(I)$-$V$ Curves of DSSCs in a dark environment. The cathode composed of the (6,5) chirality film shows a low shunt resistance and a high saturated dark current.

All of the carbon nanotube films used for the counter-electrode produced DSSCs performing with an ideality factor higher than 2. While this is not ideal for a diode or solar cell, it is not unusual in a DSSC. The cathode composed of the (6,5) film had the highest ideality factor due to the short between the film and the FTO anode increasing recombination. This is consistent with the high saturated dark current, and the low shunt resistance. Finally, the DSSC
with the platinum counter-electrode had the better ideality factor of between 1 and 2. This could again be attributed to the higher rate of reduction two the electrolyte.
Chapter 7

Conclusion and Future Work

Samples of HiPCO SWNTs of enriched chirality were prepared using the procedure originally outlined in a 2011 paper by Liu et al. and modified for this work in order to obtain a more efficient selection of high purity single chiralities. Enriched tubes of (6,5) and (7,3) at purities 93 and 86% were obtained in a single pass. The approach used was dependent on the ratio of gel to CNTs, which varied in each batch prepared. The procedure was adjusted for each batch. The versatility of this separation technique suggests the possibility of separating small diameter and semiconducting “homemade” CNTs. Research has suggested, through atomic force microscopy data, that after sonication there is little difference in length among the different (n, m) nanotubes; consequently, length difference of the CNTs was assumed not to be a factor in chirality separation.

The different chiralities were tested by preparing DSSC with CNT/ TiO₂ composites. CNTs deposited in a thin film layer configuration were also tested within a DSSC device. An improved performance in the film over the composite was observed and attributed to decreased competition with the dye, less exposure to the electrolyte, and increased behaviour as a Schottky barrier. Additionally there was an improvement in the performance of DSSCs with photoanodes of a single chirality. Using a single-chirality in the photoanode allowed us to tune the Schottky barrier height so that there was a larger difference between the LUMO of the dye and the conduction band in the TiO₂. This process promotes electron injection from the dye to the TiO₂. The end result was an increased forward bias and less recombination. Since the single chirality
films only absorbed light at a single wavelength and did not overlap the maximum absorbance of the dye, there was little competition between the single-chirality SWNT films and the dye. Future work includes measuring the work function of each of the films to better determine the type of heterojunction at the CNT/TiO$_2$ interface.

As there was a significant catalyzing effect from the CNTs on the electrolyte, films with both mixed and single-chirality CNTs were tested as the counter-electrode of the DSSC. The mixed film, with the presence of metallic species, performed well as the DSSC counter-electrode, achieving better quantum efficiency than platinum. However, the fill factor and short circuit current was less than that of the Pt electrode translating into decreased DSSC performance. To improve the behaviour of CNTs in the counter electrode, future work will include separation of metallic from semiconducting tubes to deposit only metallic tubes as a cathode film.

The work presented in this study required the deposition of a CNT film directly on FTO glass, treated with TiCl$_4$, followed by screen-printing of a TiO$_2$ layer on top of the deposited film. Later work will include the deposition of a thin blocking layer of TiO$_2$ to protect the nanotube film from recombination with the electrolyte. There are numerous ways this can be achieved including atomic layer deposition, metal-organic chemical vapour deposition, or spray pyrolysis. Using spray pyrolysis as a CNT deposition method may also produce more compact films of differing areas adapted to the dimensions of the TiO$_2$ film, reducing contact between the CNT film and the electrolyte.
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[40] Huang, S.; Cai, X.; Liu, J. *Journal of the American Chemical Society* 2003, 125, 5636–5637


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