Nickel (II) phthalocyanine-multi-walled carbon nanotube hybrids as supercapacitors

by

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Supervisor: Dr. K.I. Ozoemena
Dedication

To my dear family who have stuck by me and supported me during the course of my studies.
Declaration

I declare that the dissertation, which I hereby submit for the degree Magister Scientiae in the faculty of Natural and Agricultural Sciences at the University of Pretoria is my own work and has not previously been submitted by me for a degree at this, or any other tertiary institution.

Alfred T. Chidembo
s27280552
Acknowledgements

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My sincere appreciation and gratitude go to my supervisor, Dr. Kenneth Ozoemena, for believing in me when no one else did. For giving me the opportunity to further my studies and mostly for the intellectual guidance and concern; it has truly been an exciting learning curve for me.

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To my friend Munyaradzi Mugadza, there are friends who stick closer than a brother, and you have done just that for me, ever since we were kids. Thanks a lot. To all my friends and colleagues, I thank you from the bottom of my heart.

Finally, I would like to thank the National research Foundation (NRF) for the financial assistance.
Abstract

The thesis examines for the first time the supercapacitive properties of three different nickel (II) phthalocyanine complexes, nickel (II) tetraamino phthalocyanine (NiTAPc), nickel (II) phthalocyanine and nickel (II) tetra tert-butyl phthalocyanine as

(a) nanocomposites with multiwalled carbon nanotubes and as
(b) an electropolymer supported on multiwalled carbon nanotube platform (MWCNT-polyNiTAPc).

The supercapacitive properties of nickel (II) tetraaminophthalocyanine (NiTAPc)/multi-walled carbon nanotube (MWCNT) nanocomposite films have been found to possess a maximum specific capacitance of 981 F g\(^{-1}\) (200 mF cm\(^{-2}\)), a maximum power density of 700 Wkg\(^{-1}\), a maximum specific energy of 134 Wh kg\(^{-1}\) and excellent stability of over 1500 charge-discharge continuous cycling. When compared to MWCNTs modified with unsubstituted nickel (II) phthalocyanine (MWCNT-NiPc) or nickel (II) tetra tert-butyl phthalocyanine (MWCNT-tBuNiPc), MWCNT-NiTAPc exhibited superior supercapacitive behaviour, possibly due to the influence of nitrogen-containing groups on the phthalocyanine. The MWCNT-polyNiTAPc electrode had a maximum (112 mF cm\(^{-2}\)) capacitance which was higher than that observed for MWCNT-polyNi(OH)TAPc (84.9 mFcm\(^{-2}\)) but lower that of the MWCNT-NiTAPc (200 mF cm\(^{-2}\)).
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<tbody>
<tr>
<td>AFM</td>
<td>Atomic force microscopy</td>
</tr>
<tr>
<td>Ag</td>
<td>Silver wire pseudo-reference electrode</td>
</tr>
<tr>
<td>Ag</td>
<td>AgCl</td>
</tr>
<tr>
<td>BET</td>
<td>Brunauer, Emmett and Teller</td>
</tr>
<tr>
<td>BPPGE</td>
<td>Basal plane pyrolytic graphite electrode</td>
</tr>
<tr>
<td>C.E</td>
<td>Counter electrode</td>
</tr>
<tr>
<td>CNT</td>
<td>Carbon nanotube</td>
</tr>
<tr>
<td>CV</td>
<td>Cyclic voltammetry</td>
</tr>
<tr>
<td>CV</td>
<td>Cyclic voltammogram</td>
</tr>
<tr>
<td>DMF</td>
<td>Dimethylformamide</td>
</tr>
<tr>
<td>EDLCS</td>
<td>Electrical double layer capacitors</td>
</tr>
<tr>
<td>FTIR</td>
<td>Fourier transform infrared</td>
</tr>
<tr>
<td>HOMO</td>
<td>Highest occupied molecular orbital</td>
</tr>
<tr>
<td>IHP</td>
<td>Inner Helmholtz plane</td>
</tr>
<tr>
<td>IR</td>
<td>Infrared</td>
</tr>
<tr>
<td>LUMO</td>
<td>Lowest occupied molecular orbital</td>
</tr>
<tr>
<td>MPc</td>
<td>Metallophthalocyanine</td>
</tr>
<tr>
<td>MWCNT</td>
<td>Multiwalled carbon nanotube</td>
</tr>
<tr>
<td>NiTAPc</td>
<td>Nickel (ii) tetraaminophthalocyanine</td>
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NiTAPc-poly = Electropolymerized Nickel (ii)
Tetraaminophthalocyanine film

NiTAPc-poly-OH = Electropolymerized Nickel (ii)
tetraaminophthalocyanine film in alkaline solution

tBuNiPc = Nickel (ii) tetra-tert-butylphthalocyanine
NiPc = Nickel (ii) phthalocyanine
PANI = Polyaniline
PTFE = Polytetrafluoroethylene
PPy = Polypyrrole
OHP = Outer Helmholtz plane
PVDF = Polyvinylidene fluoride
R.E. = Reference electrode
SEM = Scanning electron microscope
SWCNT = Single walled carbon nanotube
TBAP = Tetrabutylammonium perchlorate
UV-Vis = Ultraviolet-visible
W.E. = Working electrode
XRD = X-ray diffraction
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CHAPTER 1

INTRODUCTION


1.1 General Overview of Thesis

The past few years have seen an increased focus on developing sustainable energy devices that are environmentally friendly. Pollution from combustion of fossil fuels has triggered this massive attention on ‘green energy’. The three main electrochemical energy storage and conversion (or delivery) systems (i.e. supercapacitors, batteries and fuel cells) are currently attracting major research interests as they have been forecasted to drastically ease the current global energy crisis and reduce man’s dependence on fossil fuels (crude oil) for his main energy source.

Supercapacitors exhibit unique energy storage and conversion properties when compared to batteries and fuel cells, notably in terms of power density and fast charging and discharging. The electrode from which a supercapacitor is fabricated impacts heavily on its properties. A significant amount of research on carbon nanotubes and other carbon materials as potential supercapacitor electrodes has therefore been undertaken over the past few years. Hybrid capacitors that make use of CNTs and metal oxides or conducting polymers have also been probed for the same applications. However, no reports have been made on the use of phthalocyanines as supercapacitor electrode materials.
In this thesis, three types of nickel (II) phthalocyanines, unsubstituted nickel (II) phthalocyanine (NiPc), nickel (II) tetra-tert-butylphthalocyanine (tBuNiPc) and nickel (II) tetraaminophthalocyanine (NiTAPc) were interrogated with acid-functionalised multi-walled carbon nanotubes (MWCNTs), and their supercapacitive properties investigated using cyclic voltammetry, electrochemical impedance spectroscopy and galvanostatic charge/discharge experiments. This approach is expected to take advantage of the remarkable characteristics of the MWCNTs and MPc to make a novel and robust nanocomposite with higher capacitance compared to its constituents.

1.1.1 Aim of Thesis

The main aim of this thesis is to employ electrochemical techniques (cyclic voltammetry (CV), galvanostatic charge/discharge (CD) and electrochemical impedance spectroscopy (EIS)) to interrogate the supercapacitive properties of:

(i) NiTAPc/MWCNT, NiPc/MWCNT and tBuNiPc/MWCNT composites supported on a basal plane pyrolytic graphite electrode (BPPGE) platform and

(ii) Electrodeposited NiTAPc onto a BPPGE surface pre-modified with MWCNT.
This introduction section provides a general overview of supercapacitors, carbon nanotubes and metallophthalocyanines, electrochemical techniques (CV, CD and EIS), electrode modification strategies, spectroscopic and microscopic characterization of supercapacitor materials. In chapter two the procedure adopted for the experiment is provided. Chapter three discusses the results obtained.

The choice of nickel (II) phthalocyanine complexes supported on MWCNTs has been motivated by many factors such as: (i) cost effectiveness and ease of preparation of MPc complexes, notably NiTAPc, (ii) the high popularity of Ni complexes as supercapacitor materials compared to Fe and Co complexes, (iii) the ability to be stably immobilized onto basal plane pyrolytic graphite substrate via $\pi - \pi$ interaction, (iv) the high surface area, high conductivity and interconnectivity of CNTs that enhance capacitive properties.
1.2 Supercapacitors

Electrochemical capacitors also known as supercapacitors, ultracapacitors or power capacitors are energy devices that have high cycling stability, energy and power density.\(^1\) Supercapacitor and ultracapacitor are the two terms that are mainly used to describe these energy storage devices.

1.2.1 A Brief Historical Perspective

The first patent on the supercapacitor was granted to Becker at General Electric Corp. in 1957, in which they proposed a capacitor based on porous carbon material. This was the first electrical device using double-layer charge storage.\(^2\) Becker’s device was however never commercialized. SOHIO were the next big company to work on supercapacitors where they attempted to market their device using high surface carbon materials.\(^3\) Other companies such as NEC, ECOND and Panasonic also marketed their own devices with Panasonic introducing the “Goldcap” double layer capacitor in 1978 in Japan. In the 1990’s, Cap-XX, Nippon Chemi-con (NCC) of Japan and Nesscap were the major role players in manufacturing supercapacitors.

In 2002 Maxwell Technologies of the USA purchased the Montena Components of Rossens, Switzerland, an electrochemical capacitor
manufacturer making it one of the leading US producers of electrochemical capacitors. Today, Maxwell Technologies has become one of the major manufacturers of supercapacitors in the world. Perhaps, the greatest potential for growth and development of the supercapacitor lies in the automotive industry where hybrid electric vehicles (HEVs) are increasingly becoming popular as well as in memory protection of electronic devices.

1.2.2 Design and Operation Principle

Supercapacitors operate on the same principle as traditional capacitors although they make use of electrodes that have a much higher surface area thereby increasing the capacitance and energy. A supercapacitor is made of two electrodes that are separated by a thin insulator called a separator. The two electrodes are required to be electrochemically inert. The separator is made of porous material so as to facilitate the movement of the electrolyte ions into the pores.

An electrolyte is a material that provides conductivity between the positive and negative electrodes of a cell. The two electrodes, at some distance apart are immersed in an electrolyte. Energy storage is achieved by the separation of charges when a voltage is applied to the electrodes. Negative ions will diffuse to the positive electrode and the positive charges to the negative electrode. This is illustrated in Figure 1.1. The voltage applied forces the formation of an electrochemical double layer (Helmholtz
layer) at the inner surface of the electrodes where the charge is stored. The double layer has a thickness of about 5-10 Å.\textsuperscript{5} To explain the mechanism of charge storage at the electrode/electrolyte interface an area of one of the two electrodes will be examined in this section.

\textbf{Figure 1.1:} Schematic diagram showing the mechanism of a supercapacitor made of porous carbon material. Figure adapted from Kotz and Carlen.\textsuperscript{5} (b) Operating principle of an electrical double layer capacitor. Figure adapted from Frackowiak and Béguin.\textsuperscript{6}
Figure 1.2: Proposed model of the electrical double layer region under conditions where anions are specifically adsorbed. Figure adapted from Bard and Faulkner.⁷

An array of charged species and oriented dipoles existing at the metal-solution interface is called the electrical double layer. The electrical double layer forms at the electrode-electrolyte interface where the electrode surface accommodates one layer and the solution another as shown in Figure 1.2.⁴ The result is a separation of charge between the electrode surface and the solution. The double layer capacitance is generally a function of potential, which makes it different from a standard capacitor. The inner layer at the electrode surface known as the compact, Helmholtz, or Stern layer is composed of solvated molecules and other ions or molecules that are thought to be adsorbed. The inner layer can be
divided into two Helmholtz planes, the inner Helmholtz plane (IHP) and the outer Helmholtz plane (OHP).

From Fig. 1.2, the IHP is located at a distance $x_1$ and the OHP at a distance $x_2$. The OHP is the location for the maximal charge density. The IHP is the closest distance of approach of specifically adsorbed ions. As solvated ions interact with the charged metal only long-range electrostatic forces dominate, so that their interaction is essentially independent of the chemical properties of the ions. These ions are said to be nonspecifically adsorbed. The structure of the double layer can affect the rates of electrode processes in as much as it can be affected by the structure of the electrode and the electrolyte layer itself. Due to the different mechanisms of charge storage, different classes of supercapacitors have emerged. The situation in figure.1.2. can be represented by a simple circuit.

![Simple circuit showing the connection of electrodes in a two electrode system.](image)

**Figure 1.3:** Simple circuit showing the connection of electrodes in a two electrode system. Figure adapted from Winter and Brodd.

$C_1$ and $C_2$ are the double layer capacitances of the anode and cathode respectively and $R_i$ is the internal resistance of the cell. Equation 1 best
describes the calculation of total capacitance for capacitors connected in series.

\[
\frac{1}{C} = \frac{1}{C_1} + \frac{1}{C_2} \tag{1}
\]

The amount of electrical charge stored in a capacitor is reported in Farads (F), pico Farads (pF) or microfarads (µF). The specific capacitance is usually expressed in F/g or F/cm². However, typical values for specific capacitance (F/cm²) are usually in the range of 15-50 µF/cm².¹ The capacitance, \( C \), is defined as the ratio between the stored charge, \( Q \), to the applied voltage.

\[
C = \frac{Q}{V} \tag{2}
\]

In a normal capacitor, capacitance, \( C \), is given by equation (3).

\[
C = \frac{A}{4\pi d} \tag{3}
\]

where \( A \) is the area of the plates in square metres and \( d \), the distance between the electrode plates in metres. \( C \) in this case has the units for length. In cases where the electrode plates are separated by a dielectric medium (\( \varepsilon = \text{Fm}^{-1} \)), the capacitance is given by equation (4) where the units for capacitance are in F.

\[
C = \frac{A\varepsilon}{4\pi d} \tag{4}
\]

To achieve high capacitance values for electrochemical capacitors, the area (A) of the electrodes has to be increased since the distance, \( d \) is
number of factors and few of these will be mentioned here. The capacitance of a two electrode system and a single cell electrode vary significantly by a factor of four.\textsuperscript{10} It is therefore imperative that in reporting these values, one states the type of electrode measurement used. The choice of electrolyte also has to be done carefully as the decomposition voltage of the electrolyte determines the voltage at which the supercapacitor operates (i.e. the operating voltage of a supercapacitor). Different electrolytes have been studied with aqueous electrolytes being restricted to a maximum voltage of 1 V due to the thermodynamic electrochemical window of water (1.23 V). The possibility of achieving higher voltages comes from organic electrolytes where the maximum working voltage is limited to 2.5-2.7 V.\textsuperscript{10} However, organic electrolytes increase the resistance and have lower conductivity. The amount of maximum usable power of supercapacitors that use organic electrolytes is low due to the high resistance. The energy, $E$ stored in a capacitor is directly proportional to the capacitance.

$$E = \frac{1}{2} CV^2 \text{ ................................................................. (5)}$$

In general, the power $P$ is the energy represented per unit time. From equation 5, high values of $C$ and $V$ automatically imply a higher energy.
1.2.3 **Energy and Power Densities: The Ragone Plot**

*Energy density* (expressed in Wh/kg) and *Power Density* (expressed in W/kg) are two terms that are mainly used in energy systems to compare energy contents and rate capabilities respectively. Both quantities have their density as a quantity per unit mass or per unit volume. A Ragone plot (Fig.1.4) has been developed to show the relationship between power and energy densities for various energy devices and also highlights the performance characteristics of these energy storage and conversion systems.

![Ragone Plot Diagram](image)

**Figure 1.4:** A simplified Ragone plot of specific power versus specific energy for various energy storage devices.⁴,⁵
Conventional capacitors from the diagram above show high power density but lower energy density when compared with batteries and fuel cells. This implies that batteries can store more energy than a capacitor but the delivery time of this energy is longer, hence a lower power density. A capacitor, although it stores less energy can quickly discharge and rapidly release large amounts of power resulting in a high power density. The importance of the supercapacitor as a link between traditional capacitors and batteries is therefore highlighted in the Ragone plot. Although a supercapacitor provides greater capacitance, it still falls short of reaching the energy densities of batteries and fuel cells. For this reason, this work will focus only on the supercapacitor in an attempt to improve the capacitance at the electric double layer and consequently the energy density of the device.

1.2.4 Relevance of Supercapacitors in Energy Devices

Supercapacitors serve as intermediate systems that bridge the power/energy gap between traditional capacitors (high power) and batteries (high energy). Supercapacitors, due to their long cycle life and stability, find numerous applications in car audio systems, photo and video cameras, solar lanterns, lightweight electronic fuses, starting power of fuel cells, memory protection of computer electronics etc. Furthermore, they are non toxic and are applicable in a large temperature range.
Supercapacitors have a shorter charging time and high power density compared with fuel cells and batteries.

A closer look at the Ragone plot clearly indicates the following points:

(a) Fuel cell is a high-energy system;
(b) Supercapacitor is high-power system; and
(c) Batteries possess intermediate power & energy properties;
(d) No single electrochemical energy storage system can match the internal combustion engine

To be able to match the performance characteristics similar to those of the internal combustion engine, the new trend is to combine these electrochemical energy storage systems in electronic devices and hybrid electric vehicles. For example, the diagram (Figure 1.5) below explains how this is done with supercapacitors and batteries.

![Diagram](image)

**Figure 1.5:** (a) A battery powered device where the EC supercapacitor provides power backup of the load in case of disconnection of the battery. (b) An AC-voltage powered device involving heavy switching currents. The EC protects the critical load (e.g. memory) from large voltage drops. Figure adapted from Kotz and Carlen.\(^5\)
1.2.5 Classification of Supercapacitors

Supercapacitors are classified into two different categories namely, (i) Electric Double Layer Capacitors (EDLCs) and (ii) pseudocapacitors. Recently, a third category has been added that incorporates hybrid capacitors. The operation mechanism of EDCLs involves the non-Faradaic separation of charges at the “double-layer” (i.e., electrode/electrolyte interface) while pseudocapacitors involve fast Faradaic, redox reaction of electroactive materials at the interface. Hybrid capacitors involve both mechanisms.

EDLCs make use of high surface area carbon materials such as activated carbon, carbon nanofibres, carbon aerogels and carbon nanotubes (CNTs). Pseudocapacitors employ metal oxides such as RuO$_2$, NiO$_2$ and MnO$_2$ and conducting polymers as electrode materials.$^4, 10$ An active research interest involves the integration of EDLC and pseudocapacitive materials to obtain high-performance supercapacitors (Hybrid capacitors) compared to the individual capacitor system.$^9$
Figure 1.6: General classes of supercapacitors. Figure adapted from Halper and Ellendogen.¹¹

1.2.5.1 Electric Double Layer Capacitors (EDLCs)

Activated carbons, carbon aerogels, carbon xerogels, carbon nanotubes, and carbon nanofibres have attracted a lot of interest from researchers in the supercapacitor industry. After the engineers at SOHIO
had used carbon as a material of choice for the manufacture of supercapacitor electrodes, it has become one of the most frequently used electrode material in different modifications. The investigation of carbon based electrodes for supercapacitors have largely been fuelled by the attractive properties of carbon.

Carbon is a low cost material that has an impressive surface area of about 2500 m$^2$/g ideal for supercapacitor electrodes. Its porous nature means that carbon based electrodes can store charge in the pores that exist in the electrode. The availability of pores in any electrode material for supercapacitors is of vital importance with the pore size and pore volume playing a pivotal role in charge storage. An et al. reported micropores (<2nm) and macropores (>50 nm) to be inappropriate for supercapacitor electrodes. Micropores provide a slow rate of electrolyte diffusion while the macropores are too large to retain hydrated ions. Mesopores are therefore ideal for storing charge in supercapacitors. According to Frackowiak et al. ionic conductivity relates the mobility of ions inside the pores. The rate of electrochemical accessibility is dictated by the mobility of ions inside the pores. Larger pores are easily accessible whereas smaller pores are difficult to access, making the movement of ions difficult. A network of resistors was therefore postulated by Frackowiak et al. due to the non-uniformity in resistance throughout the electrode material as a whole.
(a) **Activated carbons**

Activated carbons, in the activation process can be tailored in such a way that the distribution of pores, pore size and pore volume can be controlled. Activated carbons achieved by treatment with KOH have been reported to exhibit a high oxygen content that influences the electrochemical characteristics. Shi\(^{15}\) suggested the strong dependence of power density of activated carbons to the porous structure. Shi *et al.* went on to study the relationship between BET surface area, total pore volume, average pore size, pore volume and pore distribution and capacitance. They found that there is no linear relationship between the specific double layer capacitance and surface area. They also reported that the specific double layer capacitance per unit of external surface area is strongly dependant on the porous structure. Since activated carbon is usually in powder form, binders such as polyvinylidene fluoride (PVDF) \(^{16}\) and polytetrafluoroethylene (PTFE) \(^{17, 18}\) are necessary for making a paste that can be used as a practical electrode.

(b) **Carbon aerogels**

Carbon aerogels are highly mesoporous materials that consist of a continuous rigid solid framework and an open, continuous network of pores. Amongst their favorable characteristics are the good electrical
conductivity, controllable pore structure and highly useable surface area. These qualities make carbon aerogels good candidates for supercapacitor electrodes.

(c) Carbon nanotubes

Carbon nanotubes have emerged as leading materials for electrodes since their first use as supercapacitor electrodes was reported by Niu et al. The highly porous nature, chemical and thermal stability as well as high surface area give carbon nanotubes an edge over other materials being investigated as supercapacitor electrodes. The porous nature of carbon nanotubes allows easy access of ions from the electrolyte into the pores of the electrode. Both single walled carbon nanotubes (SWCNTs) and multiwalled carbon nanotubes (MWCNTs) have undergone immense research as potential electrode materials. Niu et al. reported the use of MWCNTs as supercapacitor electrodes with capacitance as high as 102 and 49 F/g at 1 and 100 Hz respectively in 38% \( \text{H}_2\text{SO}_4 \) as the electrolyte and a maximum specific capacitance of 113F/g. Frackowiak et al. reported a maximum capacitance value of 135 F/g. In both cases, the two electrode system was employed.

The capacitance of CNTs can be significantly enhanced by functionalizing them first. Functionalization involves the treatment of CNTs with nitric acid/ sulphuric acid mixture under harsh conditions resulting in
defects on the walls of the carbon nanotubes, an essential property for charging the double layer and also enhancing wettability of CNTs. By functionalizing the CNTs, surface carboxyl groups are added resulting in pseudocapacitance and a deviation from ideal double layer capacitive behaviour. Excessive oxidation however lowers conductivity due to the damage caused on the CNT structure.\textsuperscript{10} Apart from functionalization of CNTs, the use of binders such as PVDF and PTFE have been reported to increase specific capacitance of CNT based supercapacitors.\textsuperscript{18}

\textbf{1.2.5.2 Pseudocapacitors}

Pseudocapacitance has been described by Conway as arising from surface faradaic processes. In such instances, the charge, $Q$, passed during the oxidation and reduction process is continuously, and reversibly, dependant on the electrode potential.\textsuperscript{21} Electrosorption of H or metal adatoms and redox reactions of electroactive species are some of the processes associated with pseudocapacitance.\textsuperscript{6} The charging and discharging of a pseudocapacitor takes place through Faradaic processes involving electron transfer across the electrode-solution interface. The contribution of pseudocapacitance to the final capacitance of an electrode cannot be underestimated. However, cyclic life of the capacitor is reduced due to the redox processes that take place in the electroactive electrode material. Conducting polymers and metal oxides such as RuO$_2$ and other
oxides of transition metals are good examples of materials that give rise to pseudocapacitance.

(a) Metal Oxide based Supercapacitors

Research into possible supercapacitor electrode materials has intensified over the years and several materials have been investigated. Metal oxides e.g. MnO$_2$, IrO$_2$, RuO$_2$, and NiO have been studied as potential electrode for supercapacitor applications. The most popular novel metal oxides for supercapacitor electrodes are RuO$_2$ and IrO$_2$. These metals give rise to high capacitance values ranging from 720-760 Fg$^{-1}$ (three electrode system) and can be probed over a wider potential window compared with carbon materials. Although these materials exhibit favorable characteristics, they are very expensive and this limits their applications. As an alternative, inexpensive metal oxides such as MnO$_2$, NiO and Fe$_3$O$_4$ have been reported in literature.

The capacitance of metal oxides can be enhanced by mixing them with high surface area carbon materials to make nanocomposites which combine double layer capacitance and pseudocapacitance. Zheng et al. in their work, reported that MWCNTs greatly improved electrochemical properties, power density, specific capacitance and cycle performance of NiO. Recently, it has been discovered that metal hydroxides can achieve higher capacitance as compared to metal oxides.
(b) **Conducting Polymer Based Supercapacitor**

Polymeric materials such as polyaniline, polypyrrole (PPy) and polythiophenes are promising materials for supercapacitor electrodes. Conducting polymers have desirable characteristics such as

(i) high specific capacitance,

(ii) high conductivities,

(iii) “fast” charge-discharge processes,

(iv) good thermal and chemical stability

(v) high energy density and power density.

Polyaniline (PANI) is the most frequently used polymer as an electrode material for supercapacitors due to its ease of polymerization in aqueous media and high environmental stability. PANI can also exhibit different oxidation states giving rise to pseudocapacitance. Compared to carbon materials; PANI however has poor cycling stability. Capacitance values as high as 775 F/g have been reported by Gupta and Miura using the three electrode system.
The mechanism of operation of polymer based supercapacitors is shown in Figure 1.7. There are two polymer configurations, where one is negatively charged (n-doped) and the other is positively charged (p-doped). In their work on polymer based redox supercapacitors, Arbizzani et al.\textsuperscript{42} reported the limited use of n-doped compared to p-doped polymers. However a combination of n-doped and p-doped configurations as negative and positive electrodes respectively has been reported to be more promising.\textsuperscript{43} Conducting polymers have also been used to make composites with CNTs.\textsuperscript{40}


### 1.2.5.3 Hybrid Capacitors

Hybrid capacitors exploit the advantages of both pseudocapacitors and electrical double layer capacitors resulting in better performing supercapacitors that outperform both EDLCs and pseudocapacitors. The carbon materials provide a backbone with high surface area and favorable pore distribution while the pseudocapacitive material enhances the capacitance through faradaic reactions on the surface of the electrode.\(^44, 45\) Research in hybrid capacitors promises to overtake research in EDLCs and pseudocapacitors.

**(a) Composite hybrids**

The modification of CNTs with conducting polymers or metal oxides for supercapacitor electrodes has also been shown to improve capacitance.\(^36\) In such instances, the carbon nanotubes function as electron acceptors while the polymer acts as an electron donor. Supercapacitor studies of CNTs/conducting polymers have been well documented.\(^40, 46, 47\)

In this work, the best qualities of MWCNTs and metallophthalocyanines have been combined to form a hybrid capacitor with higher capacitance and remarkable cycling stability.
(b) **Asymmetric**

Asymmetric hybrid supercapacitors combine both faradaic and non-Faradaic processes to achieve high power and energy densities than EDLCs and pseudocapacitors. In asymmetric supercapacitors, the two electrodes are each made up of different material. Arbizzani, Mastragostino and Soavi\(^{48}\) reported the use of p-doped poly (3methylthiophene) as a positive electrode and activated carbon as a negative electrode. The hybrid capacitor outperformed the EDLCs. This arrangement affords the opportunity to exploit the favorable characteristics of the individual electrodes.

(c) **Battery-Type**

Asymmetric and battery-type capacitors operate in the same principle where two electrodes made up of different material are used. Battery-type supercapacitors however employ a battery electrode and a supercapacitor electrode. Such capacitors are a result of the need for storage systems which can maintain high cycle life, high energy density and fast charge characteristics. The most common materials for the battery electrode are metal hydroxides. Little research has been done on this type of
supercapacitor although it has been forecasted to bridge the gap between batteries and supercapacitors.\textsuperscript{49}
1.3 Carbon Nanotubes

The rediscovery of tubular carbon nanostructures\textsuperscript{50} in 1991 by Sumio Iijima sparked a lot of interest from researchers around the world. The first evidence of tubular nano-sized filaments similar to carbon nanotubes was first reported by the Russian scientists, L.V. Radushkevich and V.M. Lukyanovich in 1952.\textsuperscript{50a} Wiles and Abrahamson in 1978 are also reported to have discovered fibres on graphite anodes and these fibres are now known as carbon nanotubes.\textsuperscript{50b} Iijima, an NEC electron microscopist in Tsukuba, Japan, rediscovered carbon nanotubes in the soot of the arc discharge method. Carbon nanotubes exist in two main forms, multi-walled carbon nanotubes (MWCNTs) and single-walled carbon nanotubes (SWCNTs). Iijima claimed to have discovered MWCNTs in 1991 by Iijima and SWCNTs two years later\textsuperscript{51}. They reported MWCNTs to be hollow nanostructures made up of concentric cylinders of graphite that resembled graphitic shells with an adjacent shell separation of \( \sim 0.34 \) nm and diameters of \( \sim 1 \)nm. SWCNTs which are more similar to ideal fullerenes on the contrary, only have single graphitic shells.

Smalley and co-workers \textsuperscript{52} in their publication provided more insight into the synthesis of bundles of aligned SWCNTs with a yield higher than 70\%. Carbon nanotubes cannot be viewed in isolation but as structures related to fullerenes. The connectivity between these two interesting forms
of carbon lies in their functionality, with carbon nanotubes acting as nanowires and carbon fullerenes as quantum dots. Figure 1.8 shows a typical SEM image of normal carbon nanotubes.

![SEM image of carbon nanotubes](image)

**Figure 1.8:** SEM image showing normal carbon nanotubes. Figure adapted from Jiang *et al.*

### 1.3.1 Structure of carbon nanotubes

Carbon nanotubes are made up of sp$^2$ carbon atoms which are bonded together to make up graphite sheets which are rolled up to form carbon nanotubes as shown in figure 1.9. The roll-up vector determines the valuable electronic properties of the nanotubes. Carbon nanotubes are mainly synthesized using three different methods namely, arch-discharge,
laser ablation and chemical vapor deposition (CVD) and can exist in different atomic structures (figure 1.10).

**Figure 1.9:** Schematic of a 2D graphene sheet illustrating the folding of graphene sheet making up SWCNT. Figure adapted from Odom *et al.*

**Figure 1.10:** Illustrations of atomic structures (a) armchair and (b) a zigzag nanotube. Figure adapted from Popov.

### 1.3.2 Properties of carbon nanotubes

The field of carbon nanotube research is still growing and the amount of interest in this area has been phenomenal. Carbon nanotubes have unique properties such as the remarkable ability to either have conducting
or semiconducting properties. This behaviour stems from the chirality and diameter of the nanotubes.\textsuperscript{53} Additionally, carbon nanotubes have great thermal stability, high tensile strength and elasticity. Wong \textit{et al.}\textsuperscript{58} matched the theoretically predicted properties of carbon nanotubes with experimental data using atomic force microscopy (AFM). Knowledge of these properties enabled the scientists to link the materials with possible applications. In their study, Wong \textit{et al.} concluded that MWCNTs are able to store and absorb considerable energy.

Perhaps the most important property of carbon nanotubes that make them ideal supercapacitor electrodes is their high surface area. Peigney and co-workers\textsuperscript{59} reported the specific surface area of individual carbon nanotubes and bundles of carbon nanotubes as a function of tube diameter, the number of walls and the number of carbon nanotubes in a bundle. The functionalization of carbon nanotubes opens the way for research into their use as composites with other materials.

\subsection*{1.3.3 Applications}

The wide varieties of applications that carbon nanotubes receive are attributable to their remarkable properties. CNTs have the unique ability of promoting electron transfer in electrochemical reactions that has prompted their use in electrochemical sensing. The electrocatalytic activities of CNTs have also been probed thoroughly by researchers with molecules such as
NADH and H$_2$O$_2$ receiving a lot of attention$^{60-63}$. Ozoemena et al. have recently reported the use of carbon nanotubes – metal phthaloaycyanine composites for the detection of dopamine $^{64}$ and electrocatalytic detection of V-type nerve agent sulfhydryl hydrolysis products$^{65}$. The electro-reduction of ferricyanide and the oxidation of NADH, epinephrine and nor-epinephrine using MWCNT abrasively loaded onto basal plane pyrolytic graphite electrodes was performed by Moore et al. in 2004$^{65a}$. MWCNT when grown using metal catalysis usually result in the nanotubes containing iron impurities. The metal impurities were however shown to aid the electrocatalysis of hydrogen peroxide at MWCNT-modified electrodes by Banks et al$^{65b}$.

The use of carbon nanotubes and nanostructured carbons in energy storage devices has been reported by Frackowiak and Béguin$^{66}$. Three important energy devices, lithium batteries, supercapacitors and fuel cells all make use of carbon nanotubes. In lithium – ion batteries, CNTs find their place in the construction of the negative electrode where MWCNTs play a pivotal role in the formation of a double layer$^{67}$. Good electronic conductivity make CNTs good materials for electrode construction for the storage of hydrogen where SWCNTs are believed to store as much as 110 mAhg$^{-1}$ of hydrogen electrochemically$^{68}$.
1.4 Metallophthalocyanines (MPCs)

The name Phthalocyanine was conceived by Linstead. The prefix phthal comes from the Greek word naphtha (rock oil) and cyanine (blue). Phthalocyanines (Pcs) are 18\(\pi\) – electron aromatic macrocycles that have a characteristic blue green colour that makes them materials of choice for making dyes. Pcs were discovered by accident in 1928 at the Grangemouth plant for Scottish Dyes Ltd. The glass-lined vessel containing reagents for the preparation of phthalimide from phthalic anhydride cracked resulting in the formation of a blue-green material. Two employees, Dandridge and Dunsworth, studied the material and discovered the material to be remarkably stable and insoluble. In 1929, a patent was granted to Dandridge and Dunsworth on the preparation and properties of the material.\(^6\),\(^7\)

The possibility of attaching around seventy metal ions in the central cavity of the phthalocyanine moiety makes the redox chemistry of these molecules very interesting to study. Reduction and oxidation reactions of MPC complexes either take place at the central metal atom, in the phthalocyanine ring itself or both. MPCs that have Zn and Cu as the central atoms have been reported by Ozoemena as having no accessible d-orbitals. This means that any redox reactions that occur will only involve the macrocyclic ring.\(^7\)
Figure 1.12: Molecular structure of nickel (II) phthalocyanine studied in this thesis where R = H (NiPc), R = -NH$_2$ (NiTAPc), R = -t Butyl (tBuNiPc).

1.4.1 Applications

Phthalocyanines owe their wide variety of applications to their properties such as insolubility not only in water but other organic solvents as well. The macromolecules furthermore have exceptional stability. Over the past years, phthalocyanines have been used in the dye industry. With the increase in technology, MPcs have been used as industrial catalysts as light absorbing materials for compact discs,\textsuperscript{72} as photoconducting materials in photovoltaic cells\textsuperscript{73}, in liquid crystal displays, sensors\textsuperscript{74} and fuel cells.\textsuperscript{75}
CHAPTER TWO

REVIEW OF METHODS OF CHARACTERIZATION OF SUPERCAPACITOR MATERIALS
2.1 Physico-chemical Characterisation of Supercapacitor Materials

2.1.1 X-Ray Powder Diffraction

X-Ray powder diffraction is a technique that has experienced significant growth over the past thirty years. The development of the Rietveld method has catapulted development of this technique to where it is today. In material sciences, some materials are only present in powder form. To elucidate their structures, techniques such as powder diffraction have had to be employed thereby playing a pivotal role in unlocking the structure of materials. X-ray diffraction finds its primary use in the fingerprint characterization of crystalline materials and the determination of their structure.

Any crystalline substance has a unique X-ray powder pattern which can be known as its “fingerprint”. The technique is also used for the determination of the bonding and packing of atoms in crystals. The area under the peaks gives important information about the quantity of each phase in a sample.
The X-ray radiation most commonly used is that emitted by copper, whose characteristic wavelength for the K radiation is \( =1.5418\,\text{Å} \). Diffraction occurs in every possible orientation of 2 theta when the incident beam strikes powder sample. Electrons in the atoms of the powder sample oscillate with the same frequency as the incident beam. Destructive interference and constructive interference are experienced when the waves are out of phase and in phase respectively. Upon diffraction, the beam may be detected by using a moveable detector, which is connected to a chart recorder. The counter is set to scan over a range of 2theta values at
a constant angular velocity. Routinely, a 2θ range of 5 to 70 degrees is sufficient to cover the most useful part of the powder pattern. The particle size is best estimated by the Debye-Scherrer equation,

$$L = \frac{K_s \lambda}{B \cos \theta} \quad \text{(9)}$$

Where $L$ is the average particle size, $K_s$ is the scherrer’s constant, $\lambda = 1.54056 \text{ Å}$, $B$ is the width of a strong peak in radians and $\theta$ the corresponding Bragg angle.\(^{79}\) For supercapacitor electrode materials, broad peaks are ideal as these imply the amorphous nature of the material and consequently a higher surface area. The peak height observed in the XRD pattern is related to the crystal size of the material under investigation.\(^{80}\)

![Figure 2.2](image)

**Figure 2.2:** Typical X-ray diffraction pattern for polyaniline nanowires studied as potential supercapacitor electrodes. Figure adapted from Gupta and Miura.\(^{34}\)

In this work, X-ray diffraction provided useful information regarding the presence of Nickel metal in the hybrids.
2.1.2 Infra-Red Spectroscopy

Infrared spectroscopy (IR) has become a well accepted technique for the study of surface reactions. Its huge success as an analytical tool has been largely due to its ability to examine adsorbed molecules on solid surfaces. IR makes use of infrared radiation to fingerprint a sample. Infrared radiation is transmitted through the sample and upon hitting it; some of the radiation is adsorbed while some of it passes through the sample. A spectrum is then recorded in the form of a plot of absorbance versus wavenumber. This results in a molecular fingerprint being created as no two samples can produce identical spectra.

Fourier Transform Infrared Spectroscopy (FTIR) can therefore provide a wealth of information regarding the identity and quality of a sample in addition to the components in a mixture. Some of the advantages of FTIR over other dispersive techniques include; high sensitivity due to the high signal-to-noise ratio, high rate of data generation, simplicity and reduced heating effect of the beam. For functionalized carbon nanotubes, stretching absorption peaks of importance are mainly the C=O, C-O ester groups usually observed around 1700 and 1000 cm$^{-1}$. Phthalocyanines on the other hand are usually characterized by UV-Vis spectroscopy as discussed in the following section.
2.1.3 UV-Vis Spectroscopy

UV-Vis spectroscopy is one of the oldest and important analytical techniques used for the quantitative determination of a variety of species. The technique finds a lot of applications in many areas such as physics, chemistry and biochemistry. The technique is popularly used for phthalocyanines where the B and Q band are thoroughly analysed. This is discussed in more detail in chapter 3.

The Bouguer-Lambert-Beer law forms the mathematical-physical basis of light-absorption measurements on gases and solutions in the UV-Vis and IR region. Beer’s law is represented as follows

\[ A = -\log T = \log \frac{P_0}{P} = \varepsilon bc \] ............... (10)

Where \( A \) is the absorbance, \( P_0 \) and \( P \), the incident radiant power and transmitted radiant power respectively, \( \varepsilon \) is the molar absorptivity, \( b \), the path length of the sample and \( c \), the concentration of absorber. A monochromatic beam of radiation with power \( P_0 \) is passed through a homogeneous medium. As the beam is transmitted through the box, some \( n \) ions, molecules or atoms absorb the radiation thereby decreasing the power of the incident ray to \( P \). The full derivation of Beer’s law is not shown in this work but Strong discussed this in detail. Typical solvents used in UV-Vis spectroscopy are \( n \)-heptane, water and trifluoroethanol, usually transparent from ca. 800 nm in the UV-Vis region.
2.1.4 Scanning Electron Microscopy

The Scanning Electron Microscope (SEM) is a powerful and versatile tool used for the imaging of surfaces. This is particularly important in the field of nanotechnology and supercapacitor studies where carbon nanotubes are frequently used. The SEM uses electrons instead of light in producing an image. The instrument allows for the observation of materials on a very small scale ranging from nanometers to micrometers. An ability to produce three-dimensional images is one of the most outstanding capabilities of the SEM apart from the high resolution, higher magnification and ease of sample observation. A large depth of field of the SEM allows for the focusing of larger areas of the specimen at one time resulting in three-dimensional images.

Although SEM is the most widely used technique in surface imaging, the technique has a lower sensitivity to surfaces when compared with other imaging techniques. When an electron beam is focused on a sample, the electron beam causes the emission of secondary electrons, X-rays, backscattered electrons and photons of various energies. The secondary electrons are of particular interest as these result in the formation of an image. The X-rays upon analysis can yield vital elemental information. The electron beam can be focused on one position or moved in a raster across the surface of the specimen to form images. SEM provides useful information on supercapacitor electrodes. The porous
structure of electrodes and alignment of carbon nanotubes are observed using this imaging technique. Such structures are essential for the storage of charge in the electrical double layer.

### 2.1.5 Transmission Electron Microscopy (TEM)

TEM uses the same principle as that of a light microscope. However, instead of light, TEM uses electrons. An electron beam with uniform current density is directed onto a specimen. As electrons interact with the thin specimen, an image is recorded on an imaging device. The amount of power used to generate electrons allows for higher magnification and better resolution.

An electron gun is responsible for producing the electrons that are irradiated onto the specimen. Two condenser lenses are responsible for focusing the beam into a pinpoint beam. A condenser aperture then restricts the beam by knocking out high angle electrons. As the beam comes into contact with the specimen, part of it is transmitted and focused by the objective lens forming an image. The image is then passed through the intermediate and projector lenses which enlarge it before it hits the phosphor screen where it can be seen. The image contrast is due to the thickness of the specimen. Darker areas on the image correspond to fewer electrons being transmitted through due to the thicker or denser nature of the material at that point. Lighter areas of the image are a result of more
electrons being transmitted through the material in areas that are thinner or less dense.\textsuperscript{75}

TEM provides information on the diameter and length of carbon nanotubes. The method also allows the user to observe the effect of functionalizing carbon nanotubes for supercapacitor electrodes. The activation and functionalization of carbon nanotubes results in shorter nanotubes which can be observed on the images provided by TEM and consequently a higher surface area is ideal for supercapacitor electrodes.\textsuperscript{86}

2.1.6 Atomic Force Microscopy (AFM)

AFM is an imaging technique that relies heavily on the forces that exist between the tip and the sample under investigation. In AFM, a tip is fixed to the end of a flexible cantilever. As the tip moves across the surface of the sample, changes in surface height are measured by a laser beam in the quadrant photodetector.\textsuperscript{81}

At the same time forces between the tip and the surface are detected by the tip. The different types of forces that can be encountered are as follows:

- Van der Waals Forces – these forces are present in all materials and play a major role in imaging by AFM.
• Electrostatic force- as ions are brought closer to each other, the attractive forces between them increase sharply.

• Adhesive forces – As AFM tips are used over and over again, they lose their shape and sharpness resulting in a greater contact area between the tip and the sample. This leads to the presence of “adhesive forces”.

• Double Layer forces – A positively charged layer called an “ionic atmosphere” can be formed at the solid – liquid interface when imaging is performed in an aqueous medium.\(^8\) In the case of supercapacitor electrodes, it is essential to have information on the topology of the electrode material. Surface roughness can be detected using AFM and this is particularly important when coupled with electrochemical impedance spectroscopy (EIS) data to model the circuit suitable for a supercapacitor electrode.
2.2 Electrochemistry: Characterization of Supercapacitor Electrodes

Electrochemistry is the interplay between chemistry and electricity where reactions involving electrical currents and potential are dealt with.\(^\text{88}\) Electrochemical reactions are invariably oxidation/reduction reactions carried out under special circumstances. Oxidation and reduction can simply pertain to the donation of electrons by one atom to the other. The two always occur together. When oxidation or reduction occurs, electrons carry electrical currents which can be used to carry out useful work like lighting a bulb.\(^\text{89}\) Electrochemical reactions take place between the electrode surface and the solution. Studies in supercapacitor development all focus mainly on the type of reactions taking place at the electrode-solution interface. Electrochemistry finds numerous applications in developing sensors, corrosion studies, environmental protection and development of energy devices such as fuel cells and batteries. There are many electrochemical techniques that exist but in this work, cyclic voltammetry, electrochemical impedance spectroscopy and galvanostatic charge-discharge will be discussed.
2.2.1 Cyclic Voltammetry (CV)

Cyclic voltammetry (CV) is a potentiodynamic experimental method and probably the simplest and most versatile electroanalytical technique used by electrochemists for the study of electroactive species. A number of methods have in the past been employed for the measurement of reaction kinetics but CV has stood out due to the ease of measurement and its effectiveness in observing redox behaviour over a wide potential window. The first step in characterizing any electroactive material is to perform cyclic voltammetry over the chosen potential window.

(a) Instrumentation

In a three electrode system, a working electrode is immersed in the electrolyte, a potential is applied to it and the resulting current recorded, yielding a cyclic voltammogram. A three electrode cell consists of a working electrode, a reference electrode (usually SCE or Ag/AgCl) and an auxiliary or counter electrode (platinum) as shown in figure 2.15.
Figure 2.3: Schematic diagram of a three electrode cell used in this work.

A potentiostat applies a potential to the working electrode which is measured against the reference electrode.\textsuperscript{90} The auxiliary electrode generates the current required to sustain electrolysis at the working electrode. \textsuperscript{91} An excitation signal which is presented as a linear potential scan with a triangular waveform is passed between the two electrodes (figure 2.4). A cyclic voltammogram is obtained by measuring the current at the working electrode where the reaction takes place during the potential scan.\textsuperscript{92} The voltammogram is a display of current versus potential (figure 2.5).
Figure 2.4: A typical cyclic voltammogram excitation signal recorded for a reversible single electrode transfer reaction is in a solution containing only a single electrochemical reactant. Figure adapted from Kissinger and Heineman.\textsuperscript{90}

(b) **Fundamentals of cyclic voltammetry**

Nicholson and Shain have contributed significantly to the understanding of cyclic voltammetry and the development of the technique.\textsuperscript{93} Nicholson and Shain performed quantitative simulations of CV in 1964 and the field of electrochemistry has never been the same since then. Perhaps the easiest and simplest explanation of how cyclic voltammetry operates has been given by Kissinger and Heineman.\textsuperscript{90} They reported how a typical cyclic voltammetry experiment progresses using 6
mM K₃Fe(CN)₆ in KNO₃. A typical cyclic voltammogram they obtained is shown in Figure 2.5.

**Figure 2.5:** Typical cyclic voltammogram for a redox active species, K₃Fe(CN)₆ in 1 M KNO₃ versus SCE. Figure adapted from Kissinger and Heineman.

The scan was performed over a potential window of -0.2 to 0.8 V. At 0.8 V, Fe(CN)₆³⁻ is not electrolyzed, upon the switching on of the electrode. The forward scan begins at this potential moving towards more negative potentials to reduce Fe³⁺(CN)₆³⁻ as shown at point (b). The chemical reaction at (b) is shown below.

\[
\text{Fe}^{III}(\text{CN})_{6}^{3-} + e \rightarrow \text{Fe}^{II}(\text{CN})_{6}^{4-} \ldots \ldots (11)
\]
From $b-d$ the current increases rapidly until the concentration of Fe$^{\text{III}}$(CN)$_6^{3-}$ diminishes and that of Fe$^{\text{II}}$(CN)$_6^{4-}$ increases causing the current to reach a peak ($d$). The current rapidly decreases ($d-g$) as the electrode becomes an oxidant. The reverse scan then begins, this time with Fe$^{\text{II}}$(CN)$_6^{4-}$ being oxidized back to Fe$^{\text{III}}$(CN)$_6^{3-}$ in the process

$$\text{Fe}^{\text{II}}(\text{CN})_6^{4-} \rightarrow \text{Fe}^{\text{III}}(\text{CN})_6^{3-} + e \quad \text{.........(12)}$$

An anodic current ($i-k$) results and as concentration of Fe$^{\text{II}}$(CN)$_6^{4-}$ decreases, another peak is observed ($j$). The solution surrounding the electrode is now deficient of Fe$^{\text{II}}$(CN)$_6^{4-}$ but abundant in Fe$^{\text{III}}$(CN)$_6^{3-}$ completing the cycle and ready to start a new one. A complete voltammogram is therefore recorded. A cyclic voltammogram is characterized by some important parameters such as cathodic ($E_{pc}$) and anodic ($E_{pa}$) peak potentials, the cathodic ($i_{pc}$) and anodic ($i_{pa}$) peak currents, the cathodic half-peak potential ($E_{p/2}$) and half wave potential ($E_{1/2}$).

Two limiting cases exist where CV is employed. The electrode processes can either be reversible or irreversible. All reversible voltammograms have the typical shape as that shown in Figure 2.5. A reversible voltammetry can be characterized by ‘fast’ electrode kinetics in relation to the rate of mass transport to the electrode. Reversible electrochemical behaviour also implies that the Nernstian equation (13) is obeyed on every point of the voltammogram.
\[ E = E^0_f(A/B) + \frac{RT}{F} \ln \frac{[B]_0}{[A]_0} \]  

((13))

Where \( E \) is the applied potential, \( E^0_f \) the formal electrode potential and \([A]_0\), \([B]_0\) the concentrations of reactant and product respectively. \( R \) is the gas constant, \( T \), the absolute temperature and \( F \), Faraday’s constant.

Other characteristics for a reversible reaction are that \( \Delta E = E_{pc} - E_{pa} = 59 \) mV (for single electron reactions) is independent of scan rate (v).

The ratio of \( i_{pc} \) to \( i_{pa} \) is equal to 1.94

Irreversible reactions are due to slow electrode kinetics at the electrode-electrolyte interface. \( \Delta E \) varies with scan rate and the reverse peak is absent for an irreversible reaction. Quasi-reversible reactions are characterized by a shift in \( \Delta E \) with shift in scan rate and \( i_p/v^{1/2} \) is independent of scan rate.

Cyclic voltammetry is probably one of the most useful techniques in electrochemistry that should be performed in the preliminary stages of any electrochemistry project. Its ease of operation, good sensitivity and fast generation of results makes it a popular tool for qualitative analysis of electrode reactions. However, cyclic voltammetry should be used in conjunction with other electrochemical techniques to provide a full picture of both qualitative and quantitative diagnosis of electrode reactions. Ideal capacitor behaviour is characterized by a rectangular cyclic voltammogram. Electrode materials that exhibit pseudocapacitance deviate from the rectangular shape resulting in a distorted voltammogram where reversible peaks are identified as illustrated in figure 2.6.
Figure 2.6: Typical voltammograms for electrochemical capacitors. Figure adapted from Frackowiak and Béguin. 9 9

2.2.2 **Galvanostatic Charge/Discharge (CD)**

The Galvanostatic charge/discharge (CD) method is probably the most dependable electrochemical method that can be used for calculating specific capacitance of electrodes. 61 A typical constant charge/discharge profile of an ideal supercapacitor is characterized by linear charging and discharge profiles as shown in figure 2.7. Furthermore, a mirror image of the charging curve and the discharging curve indicates a typical supercapacitor. In contrast, a supercapacitor that is characterized by pseudocapacitance exhibits non-linear charge and discharge profiles. 95
**Figure 2.7:** Typical constant charge/discharge profile for nafion based supercapacitor. Figure adapted from Lakatos- Varsanyi and Hanzel.\(^{96}\)

The concept of Galvanostatic charge/discharge is analogous to that of cyclic voltammetry although the latter results in an \(E-i\) plot. An excitation signal is applied to the system and causes the potential to scan from the starting potential to the final potential and back again resulting in the first scan. Different currents can be chosen resulting in different current densities that can be used to calculate specific capacitance. The potential window used in Galvanostatic charge/discharge experiments should always be consistent with that used in cyclic voltammetry.

Galvanostatic charge/discharge experiments are used to calculate specific capacitance, specific energy and specific power. Ragone plots are then plotted after calculating the values of specific power against specific energy using the formulae.
\[ SC(F / g) = \frac{[i(A) \times \Delta t(s)]}{[\Delta E(V) \times m(g)]} \] ........................................ (14)

\[ SE(Wh kg^{-1}) = \frac{[i(A) \times t(s) \times \Delta E(V)]}{m(kg)} \] ......................... (15)

\[ SP(W kg^{-1}) = \frac{[i(A) \times \Delta E(V)]}{m(kg)} \] ......................... (16)

Where \( I \) is the discharge current in ampere, \( \Delta t \) is the discharge time in seconds, \( \Delta E \), the potential difference in volts and \( m \), the mass of the electroactive material in grams. For specific energy and specific power the mass, \( m \) is in kg.\(^97\) The energy deliverable efficiency (\( \eta \ \% \)) can be calculated using equation 17.

\[ \eta (\%) = \frac{t_d}{t_c} \times 100 \] ........................................ (17)

where \( t_d \) and \( t_c \) are discharge time and charging time, respectively.

### 2.2.3 Electrochemical Impedance Spectroscopy (EIS)

Electrochemical impedance spectroscopy has been branded one of the most powerful tools in electrochemistry.\(^46\) It is the study of the variation of total impedance of an electrochemical cell with frequency of a small-amplitude AC perturbation. EIS can be applied to a host of areas of study such as corrosion studies\(^98, 99, 100\), organic coatings\(^101, 102\), energy storage devices,\(^103, 104\) sensors\(^105, 106\) and the study of liquid-liquid interfaces.\(^107\)
EIS provides a wealth of information when it comes to the study of interfacial characteristics and pseudocapacitance of the electrode.

**(a) Fundamentals of Impedance Spectroscopy**

Ohm’s law gives a simple relation between dc-potential \( E \) and dc-current \( I \)

\[
R = \frac{E}{I} \quad \text{................................................................. (18)}
\]

Ohm’s law is only limited to one circuit element which is an ideal resistor. When performing experiments, other elements come into existence and these have to be considered as well. The expression above changes when an ac-signal is applied to a cell.

\[
Z(\omega) = \frac{\Delta E(\omega)}{\Delta I(\omega)} \quad \text{................................................................. (19)}
\]

Where \( Z(\omega) \) is the complex impedance. \( \Delta E(\omega) \) is the change in potential and \( \Delta I(\omega) \), the change in current.

Electrochemical impedance is usually measured by applying an AC potential to an electrochemical cell and measuring the current through the cell. A sinusoidal potential excitation (figure 2.8(a)) is applied to the electrode interface making the cell’s response pseudo-linear. The current response leads or lags the voltage by a phase angle of \( \theta \).
The capacitance of the electrode is given by

\[ Z = \frac{1}{j\omega C} \] ................................................................. (20)

The total impedance is given by the following equation that incorporates both the real and imaginary components of the impedance. The angular frequency \( \omega \) is expressed as \( 2\pi f \).

\[ Z(\omega) = Z'(\omega) + jZ''(\omega) \] ......................................................... (21)

\( Z \) is the total impedance, \( Z' \), the imaginary impedance and \( Z'' \) the real impedance.\(^{109}\) \( Z \) is therefore a vector quantity that is plotted in the plane with either polar or rectangular coordinates (figure 2.8 (b)) with \( \theta = \tan^{-1}(Z''/Z') \). The phase difference, \( \theta \), between the voltage and the current is equal to zero for a pure resistor and 90° for a pure capacitor.\(^{110}\)
(b) **Data representation**

The relationship between the impedance and frequency is very useful and can be displayed in various ways. A Bode plot is a plot of $\log |Z|$ and $\theta$ against $\log \omega$ (figure 2.9 (b)). Bode plots are very useful and give more explicit information regarding frequency. A *Nyquist plot*, displays $Z_{\text{Im}}$ vs. $Z_{\text{Re}}$ for different values of $\omega$ (figure 2.9 (a)). Both these representations will be dealt with in more detail in chapter 3.

Figure 2.9: (a) Nyquist plot for a supercapacitor made of activated carbon and numerical data fitting Figure adapted from Qu.\(^8^0\) (b) Bode $|Z|$ and Bode angle plots of a typical supercapacitor. Figure adapted from Du and Pan.\(^1^1^1\)
(c) **EIS Fitting**

The Kramers-Kronig (K-K) test is ideally the first test that one should undertake to determine whether EIS experimental data is stable in time and linear. The test checks whether data points recorded at the beginning of an experiment are in agreement with those recorded at the end. A few assumptions have been employed to check the validity of the experimental data. These assumptions are (i) that the impedimetric response is only related to the excitation signal; (ii) that the impedimetric response is linear (or the perturbation is small, e.g., < 10mV, for non-linear systems; (iii) that the system does not change with time, say due to ageing, temperature changes, non-equilibrium conditions, etc and (iv) that the system is finite for all values of $\omega$, including zero and infinity.\textsuperscript{112} The K-K test results in a value of pseudo $\chi^2$. Bad fit results in a high value of $\chi^2$ indicating the corruption of the data recorded. An equivalent series that fits the experimental data will therefore not be found. Values of $\chi^2$ lower than $10^{-6}$ are considered to be good while those higher than $10^{-4}$ are considered to be bad. The relative error and $\chi^2$ functions are very important and accurate ways of establishing the relationship between the experimental and the modeling data.

EIS data is commonly analyzed by fitting it to an equivalent electrical circuit model as shown in Fig 2.10. Most of the circuit elements in the model are common electrical elements such as resistors, capacitors, and
inductors. To be useful, the elements in the model should have a basis in the physical electrochemistry of the system. The simplest circuit used is the Randles circuit.

![Randles equivalent circuit](image)

**Figure 2.10:** Randles equivalent circuit depicting an ideal situation.

The Randles circuit is made up of (i) the electrolyte resistance between working and reference electrodes $R_s$, (ii) the double layer capacitance ($C_{dl}$), and (iii) the Faradaic impedance consisting of charge-transfer resistance $R_{ct}$ at the working electrode-electrolyte interface and the Warburg Impedance $Z_w$, which reflects the influence of the mass transport of the electroactive species on the total impedance of the electrochemical cell. The charge transfer resistance is closely associated with the surface area of the electrode and its conductivity. A summary of all the circuit elements is given in the table 2.1.
Table 2.1: A summary of some of the circuit elements used in the description of equivalent circuits.

<table>
<thead>
<tr>
<th>Element</th>
<th>Symbol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Charge transfer Resistance</td>
<td>$R_{CT}$</td>
</tr>
<tr>
<td>Solution Resistance</td>
<td>$R_S$</td>
</tr>
<tr>
<td>Capacitance</td>
<td>$C$</td>
</tr>
<tr>
<td>Inductance</td>
<td>$L$</td>
</tr>
<tr>
<td>Constant Phase Element</td>
<td>$Q$</td>
</tr>
<tr>
<td>Warburg Diffusion</td>
<td>$W$</td>
</tr>
</tbody>
</table>

For practical situations where the electrode surface is not smooth, the modified Randles circuit is applied wherein one or both elements ($C_{dl}$ and $Z_w$) are replaced by a constant phase (CPE) element representing pseudocapacitance.

The impedance of CPE ($Z_{CPE}$) is defined as \(^{113,114}\)

$$Z_{CPE} = \frac{1}{[Q(j\omega)^n]} \quad \quad \quad \quad \quad \quad \quad \quad \quad \quad (22)$$

where $Q$ is the frequency-independent constant relating to the surface electroactive properties, $\omega$ is the radial frequency, the exponent $n$ arises from the slope of log $Z$ vs log $f$ (and has values $-1 \leq n \leq 1$). If $n = 0$, the CPE behaves as a pure resistor; $n = 1$, CPE behaves as a pure capacitor, $n = -1$ CPE behaves as an inductor; while $n = 0.5$ corresponds to Warburg impedance ($Z_w$) which is associated with the domain of mass transport control arising from the diffusion of ions to and from the electrode|solution interface. Generally speaking, CPE occurs by several factors such as (i)
the nature of the electrode (e.g., roughness and polycrystallinity), (ii) distribution of the relaxation times due to heterogeneities existing at the electrode/electrolyte interface, (iii) porosity and (iv) dynamic disorder associated with diffusion.\textsuperscript{115}

EIS, like any other technique, has its advantages and limitations. The technique is very sensitive, non-destructive and accurate but less understood. Furthermore it is used in situ and has a fast rate of generating results. However it is not a stand alone technique and needs to be complimentsed by other techniques such as cyclic voltammetry. The interpretation of data can also prove to be intricate. In conclusion, the three electrochemical techniques discussed above when coupled together; provide useful information for probing supercapacitor electrodes.

\textbf{2.2.4 Chemically modified electrodes (CMEs)}

There are many well documented methods of electrode modification, which include:

(i) \textbf{Dip-dry or Immersion}. One of the simplest methods of modifying an electrode is to dip the electrode surface into a solution of a polymer/modifier. The electrode is then withdrawn from the modifier’s solution and the solvent is allowed to dry. The main drawback of this method is that the amount of material on the electrode cannot be controlled.
(ii) **Drop-dry.** A known amount of the electroactive material is deposited onto the electrode surface and allowed to dry.

(iii) **Spin-Coating.** The electrode modifier upon deposition is evaporated by spinning the electrode on a spin coater. This method is used mainly in the semiconductor industry. A uniform film thickness is obtained.

(iv) **Electrodeposition or potential cycling.** This method involves the coating of the electrode surface by electrochemically running voltammetric scans in the solution of the modifier (i.e. not the electro-active probe).

(v) **Covalent attachment.** In this method, the formation of covalent bonds between specific functional groups on the electrode surface and the electroactive material is the main objective. It is essential to use a coupling agent to achieve the desired interactions. It is possible to form multimolecular layers at the electrode surface through this method.

The choice of the method to use to modify an electrode depended on the molecule to be attached on the surface and the type of surface. In this
thesis, drop dry method and electrodeposition were adopted because of their several advantages for the work proposed in this thesis.
CHAPTER THREE

EXPERIMENTAL
3.1 Materials and Reagents

Multi-walled carbon nanotubes (MWCNT, purity > 90 %, 110-170 nm in diameter, 5-9 µm in length) obtained from Sigma Aldrich were acid-functionalised (converted mainly into carboxylated) by undergoing very stringent acid treatment following the established methods.¹¹⁶,¹¹⁷ 1 g of MWCNTs in 140 ml of 2.6 M HNO₃ was refluxed for 48 hours. The carbon nanotube sediments were separated from the reaction mixture, washed with distilled water, and sonicated in a concentrated H₂SO₄/HNO₃ mixture (3:1 ratio, 98% and 70% purity, respectively) for 24 hours. Thereafter, the sediments were washed with distilled water, stirred for 30 minutes in a H₂SO₄/H₂O₂ mixture (4:1, 98% and 30% purity, respectively) at 70 °C, and washed again with deionised water. Finally, the purified paste-like MWCNTs were then oven-dried at 50 °C for 48 hours.

Nickel (II) tetraaminophthalocyanine (NiTAPc) was synthesised and characterised following the well-known procedure introduced by Achar and Jayasree.¹¹⁸ For a comparative study, unsubstituted nickel phthalocyanine (NiPc) was purchased from Aldrich. Nickel (II) tetra-tert-butyl phthalocyanine (tBuNiPc) was obtained from its metal-free precursor, tetra-tert-butyl phthalocyanine (tBuPc) kindly donated by Prof T. Fukuda, Graduate School of Science, Tohoku University, Japan. Introduction of nickel into the core of the tBuPc was achieved using a
similar metallation process described before. Briefly, this involved refluxing a DMF solution containing a mixture of H$_2$BuPc and excess of nickel acetate for about an hour. The DMF was then evaporated and the dark green residue purified using a short silica gel column (CHCl$_3$ + small percentage of methanol as eluent) to afford the desired NiTTBPc.

As expected, the formation of the nickel complex was easily established UV-visible spectrophotometrically in CHCl$_3$ by observing the total collapse of the split Q-bands (663 and 696 nm) of the metal-free tBuPc complex ($D_{2h}$ symmetry) to the single Q-band (668 nm) of the metal, NiTTBPc, complex ($D_{4h}$ symmetry). Ultra-pure water of resistivity 18.2 MΩcm was obtained from a Milli-Q Water System (Millipore Corp., Bedford, MA, USA) and was used throughout the experiments.
3.2 **Equipments and methods**

Field emission scanning electron microscopy (FESEM) images were obtained from JEOL JSM 5800 LV (Japan) while the energy dispersive x-ray spectra were obtained from NORAN VANTAGE (USA) at the Microscopy and Microanalysis Laboratory of the University of Pretoria. Transmission Electron Microscopy was performed on the JEOL JEM-2001 F/HT transmission electron microscope. Atomic Force Microscopy (AFM) experiments were performed with AFM 5100 System (Agilent Technologies, USA). TGA was performed with a Mettler Toledo A851 (Switzerland). Simultaneous TGA/DTA data machine was used for thermal and gravimetric analyses. Powder samples (ca. 10mg) were placed in an open 70 µL alumina pan and heated from 25-1000°C at a scan rate of 10°C / min in nitrogen atmosphere. The Perkin-Elmer GX 2000 FTIR spectrometer attached to the Perkin-Elmer Auto Image Microscope system equipped with liquid Nitrogen cooled MCT detector was used for FTIR analysis. UV-Vis spectroscopy was performed using the Cary 300 UV-Vis spectrophotometer driven by the Varian software 3.0.

The samples were prepared for XRD analysis using a back loading preparation method. XRD experiments were performed using a PANalytical X’Pert Pro powder diffractometer with X’Celerator detector and variable divergence and receiving slits with Fe filtered Co-Kα.
radiation. The samples were scanned at the required 2θ angle ranges. The phases were identified using X’Pert Highscore plus software.

Electrochemical experiments were carried out using an Autolab Potentiostat PGSTAT 20 (Eco Chemie, Utrecht, Netherlands) driven by the GPES software version 4.9. Electrochemical impedance spectroscopy (EIS) measurements were performed with Autolab Frequency Response Analyser (FRA) software between 100 kHz and 10 mHz using a 5 mV rms sinusoidal modulation. Bare or modified BPPGE disk (d = 5mm in Teflon) was used as the working electrode and was fabricated at the Chemistry Department of the University of Pretoria from BPPG plate (Le Carbone, Sussex, UK). Electrical contact with the disk was maintained through an inserted copper wire held in place with conducting silver varnish L 100 (Kemo® Electronic, Germany). An Ag/AgCl in saturated KCl and platinum wire was used as reference and counter electrodes, respectively. All solutions were bubbled with nitrogen for 5 minutes prior to each electrochemical experiment. Experiments were performed at 25 ± 1 °C.
3.3 Preparation of Nickel Phthalocyanine -MWCNT composites and electrode fabrication

Prior to modification with NiTAPc, MWCNTs or MWCNT-NiTAPc, the BPPGE surface was cleaned by gentle polishing on a p1200C Norton carborundum paper (Saint-Goban Abrasives, Isando, South Africa) followed by cleaning with cellotape process of removing graphite layers and finally rinsing in acetone to remove any adhesives. All modifiers were prepared in dry distilled DMF and used within 48 hours of preparation. MWCNT-NiTAPc composite was obtained by grinding equal amounts of MWCNT-COOH and NiTAPc (1:1 mass/mass ratio) using mortar and pestle for about 5 minutes. The mixture was then dissolved in DMF (1 mg MWCNT-NiTAPc mixture/1ml of DMF) and ultrasonicated for 30 minutes. The drop-dry method was then employed to load the MWCNT-NiTAPc hybrid onto the BPPGE surface by placing about 20 μL drop of the mixture onto the BPPGE surface and dried in the oven at ~ 50 °C for 5 minutes. The same procedure was repeated with tBuNiPc and NiPc.
**Scheme 3.1:** Schematic representation for the synthesis of the MWCNT-NiTAPc nanocomposite and electrode preparation using the drop-dry method.
3.4 Preparation of poly-Nickel Phthalocyanine -MWCNT composites and electrode fabrication

The BPPGE was placed into a solution containing 1 mM NiTAPc and 0.1 M tetra-n-butylammonium perchlorate (TBAP) which acted as a supporting electrolyte. The NiTAPc film was deposited onto the electrode by cycling under continuous cycling conditions of -0.2 to 0.9 V at 100 mVs$^{-1}$ for 40 cycles.$^{120}$ The resulting electrode (BPPGE-polyNiTAPc) was then washed with acetone and distilled water and dried in an oven at 50 °C. The BPPGE-MWCNT electrode was obtained by first drop-drying 20 µL of MWCNT dissolved in DMF and allowed to dry in an oven at 50 °C for ca. 10 minutes. The resulting electrode was then placed in a solution of DMF/NiTAPc and cycled from -0.2 to 0.9 V at 100 mVs$^{-1}$ for 40 cycles. The electrode was then rinsed and washed using acetone and distilled water to obtain the desired BPPGE-MWCNT-polyNiTAPc electrode.

The BPPGE-MWCNT-polyNiTAPc was then transferred to a fresh solution of 0.1 M NaOH and cycled in the potential window of -0.2 to + 0.8 V for 50 scans at 100 mVs$^{-1}$ to obtain the BPPGE-MWCNT-polyNi(OH)TAPc electrode. Well defined peaks appear at around 0.4 V are observed. As the cycling continues, the peaks grow steadily until they reach a steady state where no further growth is observed. This is typical of cyclic voltammograms recorded for electroformed nickel macrocyclic-based films.
in alkaline solution and shows that the film has been formed as a result of the anodic polymerization of the macrocyclic complex.

It has been postulated that the polymerization of nickel macrocyclic complexes in alkaline media is very similar to that of the nickel hydroxide electrode.\textsuperscript{121} The nickel-nitrogen tetra coordination of the complex is thought to be lost in the polymerization process. Another postulate by Roslonek and Taraszewka proposes that upon electropolymerisation, the complexes are attached \textit{via} oxo-bridges.\textsuperscript{122} Capacitance studies for the BPPGE-polyNiTAPc, BPPGE-MWCNT-polyNiTAPc, BPPGE-polyNi(OH)TAPc and BPPGE-MWCNT-polyNi(OH)TAPc electrodes were all performed in 1 M H\textsubscript{2}SO\textsubscript{4}. 
CHAPTER FOUR

RESULTS AND DISCUSSION
4.1 Physico-chemical Characterization

4.1.1 X-Ray Diffraction

Carbon nanotubes are made up of graphite sheets rolled up together as already discussed in chapter one. Broader diffraction peaks observed in the XRD patterns (figure.4.1) for the materials indicate the amorphous nature of the four materials implying a small grain size. A decreased grain size consequently contributes to the high surface area of the materials studied. It is a well known fact that materials required for capacitance studies should have a high surface area.⁴ A typical powder diffraction for functionalized carbon nanotubes and normal MWCNTs is shown in figure.4.1 (a). The normal MWCNTs and functionalized MWCNTs show very similar structures with a strong peak (100) at 31° and a weak peak (004) at 51°. The peaks indicate that the MWCNTs consist of small domains of ordered graphene sheets. From the similarities in the structures of the two forms of MWCNTs we deduced that the crystal structures of the two are the same. Functionalization therefore did not alter the novel structure of the CNTs as has also been previously reported by Jiang et. al.¹²³
**Figure 4.1:** X-ray diffraction analyses for the functionalised MWCNT, Ni (II) phthalocyanines and MWCNT-Ni (II) phthalocyanine nanocomposites
Qu suggested the dependency of specific capacitance of the electrode material on the crystal orientation of the surface.\textsuperscript{80} From the XRD patterns; the orientation for the graphite sheets has not been altered by the functionalization of the MWCNTs even when coupled with the phthalocyanines. The MWCNT-NiPc, MWCNT-NiTAPc and MWCNT-tBuNiPc nanocomposites clearly have different XRD patterns from the MWCNTs. However, the contribution of the MWCNTs is significant. The strong peak at $31^\circ$ is evident in all the XRD patterns. For MWCNT-NiPc, peaks due to the NiPc are clearly visible. A strong peak at $7.9^\circ$ was observed. Other peaks at $2\theta = 11.3^\circ$ and $18.4^\circ$ were also observed for NiPc. The same peaks were also observed in the hybrid although at lower intensity. This phenomenon indicates a change in polycrystalline structure resulting in an amorphous compound.\textsuperscript{124}

For the MWCNT-COOH and MWCNT-NiTAPc, the XRD patterns show that there is very little order between the graphene sheets.\textsuperscript{125} The XRD patterns of the neat MWCNT and NiTAPc slightly differ from the MWCNT-NiTAPc (see figure. 4.1 (c)) hybrid confirming the formation of a new compound. The main peak at $8.75^\circ$ for NiTAPc shifted to $7.44^\circ$ for MWCNT-NiTAPc indicating a change in crystallinity and interplanar spaces. The tBuNiPc exhibits a strong peak is at $2\theta = 5.8^\circ$. This peak shifts to $2\theta = 10.2^\circ$ with a significant loss in intensity. This implies the formation of a new compound (MWCNT-tBuNiPc) that is more
amorphous in nature with smaller grains. This information is in accordance with HRSEM images.

### 4.1.2 TGA Analysis

In order to obtain important information on the thermal stability of the MWCNT-COOH, MWCNT-MPc and MPc complexes, TGA was performed. The curves in figure 4.2 show that the MWCNT-NiTAPc, MWCNT-tBuNiPc, tBuNiPc and NiTAPc initially lose weight at 100°C. This phenomenon can be attributed to the desorption of physically adsorbed water from the complexes. The complexes lose about 10% of the water content compared with 3% for MWCNT-COOH. Further decomposition of MWCNT-NiTAPc complexes is observed at about 540 °C where the weight of the complex decreases more rapidly compared with the MWCNT-COOH. The same behaviour is observed for the bulk MPcs.

The TGA analysis shows that the MWCNTs are very stable. After heating to about 500 °C, only 10% of the original weight had been lost due to pyrolysis. The Phthalocyanines however disintegrated more than the MWCNTs with NiPc losing ca. 90% of the mass after heating to 800 °C. The larger tBuNiPc and NiTAPc molecules lost significant weight only after heating up to 1000 °C although the initial pyrolysis occurred at 400 °C and 450 °C respectively, higher than that of the MWCNT hybrids. From the TGA curves, the hybrids all started to disintegrate at
lower temperatures as compared with the MWCNTs and the constituent Nickel phthalocyanines. The low decomposition temperature of the MWCNT-MPc hybrids is due to the increased internal energy caused by the structural twist of the planar MPcs to match well with the cylindrical structure of the MWCNTs via π-π interactions.\textsuperscript{126}
Figure 4.2: TGA analyses for the functionalised MWCNT, Ni (II) phthalocyanines and the MWCNT-Ni (II) phthalocyanine nanocomposites. (a) MWCNT, NiTAPc and MWCNT-NiTAPc, (b) MWCNT, NiPc and MWCNT-NiPc, (c) MWCNT, tBuNiPc and MWCNT-tBuNiPc.
4.1.3 UV-Vis Spectroscopy

The UV-visible spectrum observed for phthalocyanines originates from molecular orbitals within the aromatic $18\pi$ electron system and from overlapping orbitals on the central metal atom$^{127}$
Figure 4.3: Electron absorption spectra of the MWCNT, MWCNT-Ni (II) phthalocyanine composites and Ni (II) phthalocyanine in DMF. (a) MWCNT, NiPc and MWCNT-NiPc, (b) MWCNT, NiTAPc and MWCNT-NiTAPc, (c) MWCNT, tBuNiPc and MWCNT-tBuNiPc.
The spectra of MPc complexes are due to electronic transitions from the highest occupied molecular orbital (HOMO) ($\pi$) to the lowest unoccupied molecular orbital (LUMO) ($\pi^*$). Phthalocyanines have two characteristic bands, the Q band and the B (or Soret) bands. The Soret band is due to an $a_{2u}(\pi)-e_g(\pi^*)$ transition and the Q-band to an $a_{1u}(\pi)-e_g(\pi^*)$ transition.\textsuperscript{128,129} The Q band appears in the region around 600-700 nm in the far-red end of the visible region. The origins of this band according to Davidson are due to $\pi-\pi^*$ transition on the phthalocyanine macrocycle.\textsuperscript{130}

The B (or Soret) band appears in the region of 300-500 nm. The main suggestion of the large differences occurring in the absorption spectra of the phthalocyanines in this region indicate the presence of a d-band associated with the central metal atom. Upon close examination of figure 4.3.(a) the presence of three peaks in the UV region and a characteristic splitting of the peaks in the visible region is revealed. Due to the electronic interactions between the MWCNTs and the NiPc, the broad peaks in the Q band are slightly shifted towards the blue region. Figure 4.3 (b), shows a peak at 307 nm in the UV region. The other prominent peak, the Q band is observed at 727 nm in the visible region and a shoulder peak at 653 nm. This peak has been explained by Davidson in terms of $\pi-\pi^*$ excitation between bonding and antibonding molecular orbitals.\textsuperscript{130}
Upon mixing the phthalocyanine and MWCNT, the broad and most prominent peak in the visible region is reduced significantly to a point where it is almost non-existent. In figure 4.3 (b) the broad peak at 436 nm disappears completely in the MWCNT-NiTAPc spectrum. In the Soret band, the peak at 307 nm shifts to 273 nm and is also reduced in intensity. These results can be explained by the $\pi-\pi$ interactions between the MWCNTs and the conjugated system of the Pc backbone. According to Jaegfeldt et al.\textsuperscript{131} highly aromatic compounds absorb strongly to the basal plane of graphite via $\pi-\pi$ stacking. The adsorption of species on to the basal plane of graphite becomes stronger as the number of aromatic rings increases. Chen and co-workers\textsuperscript{132} reported the similarity in these strong interactions between polycyclic molecules and carbon nanotubes.

\subsection*{4.1.4 FTIR Analysis}

The IR spectrum of the MWCNT-COOH clearly show the presence of COOH groups due peaks at 1373 cm\textsuperscript{-1} and stretching modes of C=O at 1618 cm\textsuperscript{-1}.\textsuperscript{133} The broad band at 3406 cm\textsuperscript{-1} is due to the $-\text{OH}$ stretching vibration. The origin of this peak is attributed to the adsorbed water in the MWCNT pores. Smaller peaks observed at 2847 and 2915 cm\textsuperscript{-1} respectively attributed to the stretching vibrations of the $-\text{CH}_2$ and $-\text{CH}_3$ groups.\textsuperscript{134} On mixing with the Pcs, some peaks from the
Phthalocyanines are absent in the NWCNT-MPc spectrum especially at wavelengths lower than 1500 cm⁻¹. All three phthalocyanines have notable peaks in the range of 1080-1095 cm⁻¹ corresponding to the stretching mode of the C-N bond. The peaks for the C=N bond are also visible at 1481, 1421 and 1454 cm⁻¹ for NiTAPc, NiPc and tBuNiPc respectively.
Figure 4.4: FTIR spectra for MWCNT, MWCNT-Ni (II) phthalocyanine composites and Ni (II) phthalocyanine. (a) MWCNT, NiTAPc and MWCNT-NiTAPc, (b) MWCNT, NiPc and MWCNT-NiPc, (c) MWCNT, tBuNiPc and MWCNT-tBuNiPc.
The tBuNiPc spectrum exhibits a strong peak at 2950 cm\(^{-1}\) due to the C-H stretching modes of the tert-butyl group. In the NiTAPc molecule, notable peaks are observed at 3304 and 3192 cm\(^{-1}\) which could be due to the antisymmetric and symmetric vibrations of the NH\(_2\) groups. In the MWCNT-NiTAPc spectrum, the peaks are hidden by the stronger peak at 3462 contributed by the MWCNTs.\(^{135}\)

**4.1.5 SEM characterisation**

CNTs are well known to aggregate together and form bundles due to Van der Waals forces. After functionalization, the MWCNTs are well dispersed and form a unique pore structure due to the spaces that exist between the entangled CNT mats.\(^{136}\) This is clearly evident from all the pictures in figure 4.5. Figure 4.5 (a) shows the FESEM images of the MWCNT-NiTAPc nanocomposite. Figure 4.5 (a) (inset) clearly reveals the attachment (to a certain extent, via \(\pi-\pi\) interactions) of the NiTAPc nanoaggregates (50 – 100 nm size) on the nanofibril-like structure of the acid-functionalised MWCNTs.
**Figure 4.5:** Typical SEM images of (a) functionalized MWCNTs on basal plane graphite, (b) MWCNT-NiPc (c) MWCNT-NiTAPc (Inset is a slightly magnified portion showing that some of the NiTAPc molecules are first aggregated before attaching themselves onto the cylindrical structure of the MWCNTs.) and (d) MWCNT-tBuNiPc.

The FESEM image of the MWCNT-NiTAPc is different from that reported by Wang *et al.*\textsuperscript{126} of a metal-free phthalocyanine (tetra-*tert*-butyl...
phthalocyanine, \textit{tBuPcs}) species that were uniformly and non-covalently attached to the CNTs via \(\pi-\pi\) interactions. The authors noted that when the \textit{tBuPcs} was metallated with copper to form the copper (II) tetra-\textit{tert-buty}l phthalocyanine (\textit{tBuCuPcs}), these \(\pi-\pi\) interactions were hindered, which they attributed to the destruction of the aromaticity of the conjugated \textit{tBuCuPcs} backbone due to the formation of the coordinate bond between copper and the nitrogens of the core \textit{tBuPcs}. This explanation may partly hold for our MWCNT-NiTAPc, but their non-covalent attachment cannot be ruled out as other workers\textsuperscript{137} have also proved this to be possible with nanocomposites of SWCNTs/metallated porphyrins (compounds structurally-related to phthalocyanines).

\textbf{4.1.6 TEM images}

The HRTEM images in Fig. 4.6 show the effect of functionalization on the length of the CNTs. Functionalization cuts the CNTs into a smaller size thereby enhancing the surface area of the nanostructures. Figures 4.6 (c) and (d) show the interior empty structure of the MWCNTs with the phthalocyanine molecules attached to the walls of the CNTs appearing as little black dots.
**Figure 4.6:** TEM images of (a) pristine MWCNTs (b) acid – cut / functionalised MWCNTs (c) and (d). MWCNT-NiTAPc nanocomposite.
4.2 Electrochemical Characterization

4.2.1 Cyclic Voltammetry

Cyclic voltammetry is a fast and useful method when it comes to the evaluation of electrochemical capacitor properties of electrode materials. The results shown in the diagrams below indicate the capacitive nature of the CNT-phthalocyanine nanocomposites. The electrochemical properties of the hybrids and their constituents were studied by cyclic voltammetry in 1 M H$_2$SO$_4$ aqueous solution.

![Cyclic Voltammetry Diagram]

Figure 4.7: Comparative voltammetric evolutions for MWCNT-NiTAPc with different ratios of CNTs: MPc (i) 1:5, (ii) 1:2, (iii) 5:1, (iv) 2:1 and (v) 1:1 at 50 mVs$^{-1}$. 
Preliminary studies to show effect of varying amounts of MWCNTs and NiPc were performed. A ratio of 1:1 (MWCNT:MPc) proved to offer more capacitance when compared to other ratios. Equal masses of MPc and MWCNTs were therefore mixed and sonicated to produce the nanocomposites used for capacitance studies throughout this work.

Figure 4.8: (a) Overlaid cyclic voltammograms comparing a bare BPPGE (i), BPPGE-NiTAPc (ii), BPPGE-MWCNT (iii) and BPPGE-MWCNT-NiTAPc (iv) at 50 mV s\(^{-1}\). (b) Expansion of the voltammograms of the (i) to (iii). Electrolyte = 1 M H\(_2\)SO\(_4\).

The overlaid voltammograms in figure 4.8 show that the MWCNT-NiTAPc nanocomposite exhibits larger capacitance compared to the other electrodes. In acidic medium, pseudocapacitance arises from the redox reactions of the functional groups on the surface of the materials. In the
case of acid pretreated MWCNTs these are likely to correspond to the two-electron, two-proton redox process of surface quinones, which are also introduced as surface oxo-groups during acid pretreatment of MWCNTs.\textsuperscript{138, 139, 140} These surface quinones give rise to two small, ill-defined, quasireversible redox waves observed at ca. 0.2 V vs SCE at low pH, corresponding to both ortho- and para-quinone surface groups.

The larger pseudocapacitance of the MWCNT-NiTAPc nanocomposite compared to that of the MWCNTs or NiTAPc alone, suggests a synergy between these two \(\pi\)-electron species. This positive synergy may have arisen from depositing electroactive-NiTAPc species onto a high conducting and high surface area support (the MWCNTs) resulting in greater double-layer charging and the large Faradaic response. In fact, a closer look at figure 4.8 (a). shows that those peaks observed for the MWCNTs (figure 4.8 (b)) are also present (as weak peaks) at the MWCNT-NiTAPc voltammogram (figure 4.8 (a)). Thus, the broad redox couple (indicative of pseudo-capacitance) of the MWCNT-NiTAPc nanocomposite (figure 4.8 (a)) at around 0.2 V is related to both the oxygen functionalities of the acid-functionalised MWCNTs as well as the Ni(II)/Ni(III) redox process of the NiTAPc species (equation 23):

\[
\text{Ni(II)TAPc} \leftrightarrow \text{Ni(III)TAPc} + e^- \quad \text{.........................} \quad (23)
\]
The participation of the nitrogen-containing groups of the NiTAPc in the electrochemistry may not be completely ruled out. For example, redox chemistry corresponding to immine like groups within the phthalocyanine structure, which is analogous to the quinonyl redox process discussed above, may also be a contributory factor to the pseudocapacitance observed in figure 4.8.

\[ C^* = \text{NH} + 2e^- + 2H^+ \leftrightarrow C^*H - \text{NH}_2 \]  \hspace{1cm} (24)

According to Hulicova-Jurcakova et al.\textsuperscript{141} pyrrol and pyridinic nitrogens are electroactive and can enhance supercapacitance. This is because pyrrol or pyrrol-like nitrogen (-NH) improve charge mobility in a carbon matrix by their ability to introduce electron-donor characteristics and enhancing the carbon catalytic activity in electron-transfer reactions, while the pyridinic nitrogen (=N) can provide a lone pair for conjugation with the \(\pi\)-conjugated rings. Although amines bonded to the carbon surface are not expected to affect the electron-donor properties of carbon materials.\textsuperscript{141} The contribution to the overall capacitance by the oxygen-containing carboxyl, carbonyl and phenol groups of the acid treated MWCNTs cannot be ignored as reported by Frackowiak.\textsuperscript{6,9}

It is a well-established knowledge to the phthalocyanine researchers and electrochemists that the electropolymers of the metallo-tetra-aminophthalocyanine (MTAPc) complexes, notably NiTAPc, can easily be formed on any carbon electrode surface (for electrocatalysis and sensing).
using cyclic voltammetry in an appropriate organic solvent (such as DMF) in the presence of a supporting electrolyte salt (e.g., tetrabutyl ammonium phosphate, TBAP). The formation of such MTAPc electropolymers are known to be due to the electroactivity of these peripherally substituted amino groups.$^{142, 143}$

At higher scan rates (figure 4.9 (a)), the observed peak-to-peak separation ($\Delta E_p$) increases slowly (ca. 100 mV) and the ratio of anodic and cathodic peak current is almost unity, indicative of fast, electrochemically reversible electron transfer kinetics. A plot of peak (anodic and cathodic) current vs scan rate was linear (figure 4.8 (b)), indicative of a surface bound redox system. Note that despite that we did not employ any binding substance in the fabrication process, the thin film of the MWCNT-NiTAPc nanocomposite on the BPPGE surface still exhibited excellent electrochemical stability, possibly due to the ability of the BPPGE to hold MWCNT-NiTAPc species via $\pi-\pi$ interactions.
Figure 4.9: (a) Typical cyclic voltammetric evolutions of BPPGE-MWCNT-NiTAPc at varying scan rates 100, 200, 300, 400, 500, 600 and 700 mVs\(^{-1}\) (inner to outer). (b) Plot of peak current (anodic and cathodic) vs scan rate. Electrolyte = 1 M H\(_2\)SO\(_4\).

At high scan rates the cyclic voltammograms of the three nanocomposites project a clear picture of the differences in capacitance. It is interesting to note that the nanocomposites retain their shape even at high scan rate.
The cyclic voltammograms are not symmetrical about the zero current axis and their deviation from perfect rectangular shapes is due to pseudocapacitance contribution by the phthalocyanine. It is however interesting to note that the cyclic voltammogram of the MWCNT-tBuNiPc is significantly smaller than that of the other two nanocomposites. The resulting low capacitance of the MWCNT-tBuNiPc hybrid is probably due to the presence of the large t-Butyl groups. The large alkyl groups reduce the hydrophilicity of the hybrid in DMF. More importantly, the protons are hindered from accessing the MWCNT support. The cyclic voltammograms for the MWCNTs did not exhibit the typical rectangular shape expected for an ideal EDLC. This behaviour should certainly be
expected considering the long digestion period employed in the functionalisation of the MWCNTs.

### 4.2.2 Impedance spectroscopy characterisation

Unlike cyclic voltammetry or galvanostatic discharge methods, electrochemical impedance spectroscopy is the least reliable and accurate technique for determining the supercapacitive properties of materials, especially those related to conducting polymeric substances.\textsuperscript{144-149} Figure 4.11 shows typical comparative Nyquist plots of the electrodes.

**Figure 4.11:** Comparative Nyquist plots of BPPGE-NiTAPc (i), BPPGE-MWCNT (ii) and BPPGE-MWCNT-NiTAPc (iii) in 1 M H\textsubscript{2}SO\textsubscript{4} at 0.30 V, between 100 kHz and 1 Hz. Inset: The expanded portion of the high frequency region, showing the fitted line for the BPPGE-MWCNT-NiTAPc.
The low-frequency differential capacitance \( (C_d) \) for each of the electrodes can be obtained from the slope \((1/2\pi C_d)\) of the plot of the imaginary component of the impedance versus the reciprocal of the frequency (i.e., \(-Z''\) vs \(1/f\)).\(^{149-151}\) The values of the specific capacitances for the three modifiers were less than 500 F g\(^{-1}\), indicative of strong deviation from the data obtained from the CV and galvanostatic methods. At 40 \(\mu\)g (~2 mg cm\(^{-2}\)) loading mass fixed at 0.3 V, NiTAPc, MWCNT and the MWCNT-NiTAPc nanocomposite had capacitances of 13, 64 and 340 F g\(^{-1}\) respectively. This strong discrepancy has also been observed by several workers for conducting polymeric substances,\(^{144-149}\) and the origin of which has long been a subject of some controversy for more than two decades. For example, Murray and co-workers\(^{144}\) have suggested the involvement of some physico-chemical heterogeneities; Tanguy and co-workers\(^{27}\) attributed such phenomenon to “deeply trapped” counter ions which remain immobile during impedance experiment; while Kalaji and Peter\(^{148}\) described it as the consequence of slow conformational changes occurring in the polymer network.

Later, Ren and Pickup\(^{149}\) performed further experiments that claimed to support the arguments on conformational changes. It should be noted that recent workers have also observed this discrepancy and attributed it to “redox-switching hysteresis” associated with conducting polymers,\(^{152}\) or the hindrance that accompanies the penetration of alternating current into
the bulk electrode. As also observed in figure 4.17, the SC value of the different mass loading of the MWCNT-NiTAPc decreases significantly from the lowest mass loading (≈ 586 F g⁻¹ at 20 µg) to the highest mass (≈ 180 F g⁻¹ at 80 µg). Thus, it is very likely that one or more of these interpretations by other workers may also hold in our case, especially considering that thick films are prone to inhibiting facile current flow and exacerbating the trapping and/or immobility of ions. Since this behaviour has mainly been reported for the conducting polymers, it suggests that the MWCNT-NiTAPc nanocomposites exhibit some properties of conducting polymeric substances.

Despite the paucity of EIS in establishing reliable data on the supercapacitive data, it still remains a powerful technique for providing valuable insights into processes that occur at the electrode/solution interface. As seen in figure 4.11, the BPPGE-MWCNT shows near-vertical line as expected for pseudocapacitor. The BPPGE-MWCNT-NiTAPc (inset) shows smaller incomplete semi-circle in the high frequency region (describing the ESR or charge transfer resistance across the electrode/electrolyte interface), compared to the BPPGE-NiTAPc, followed by a near-vertical line at low frequency region, which is typical of capacitive behaviour.

The transition point between the high frequency and low frequency component, referred to as the “knee” or “onset” frequency (f₀) describes
the maximum frequency at which the capacitive behaviour is dominant, and is a measure of the power capability of a supercapacitor; the higher the $f_o$ the more rapidly the supercapacitor can be charged and discharged or the higher the power density that can be achieved from the supercapacitor.\textsuperscript{154,155} The values of the $f_o$ were about 4 Hz for NiTAPc and MWCNTs, and 720 Hz for the MWCNT-NiTAPc. Accordingly, the higher $f_o$ value of the MWCNT-NiTAPc confirms its higher power property compared to the individual NiTAPc and MWCNTs, which is consistent with the comparative CV data shown in figure 4.2.2 (a). The reciprocal of the $f_o$ is the response time of the capacitor. Note that the $f_o$ of MWCNT-NiTAPc is about 1.4 ms, suggesting that most of its stored energy is still accessible at frequencies as high as 720 Hz. Most commercially available supercapacitors, including those specifically designed for higher power applications operate at frequencies less than 1 Hz.\textsuperscript{156}

The impedimetric behaviour of MWCNT-NiTAPc could be satisfactorily modelled by the electrical equivalent circuit comprising the modified Randles circuit with one voigt RC element (Figure 3.12), involving a solution resistance ($R_s = -11.7$ $\Omega$), a double-layer capacitance ($C_{dl} = 95$ nF), an electron-transfer resistance ($R_{ct1} = 16.6$ $\Omega$ and $R_{ct2} = 18$ $\Omega$) and constant-phase elements ($CPE_1 = 10.5$ mF with $n_1 = 0.95$ and $CPE_2 = 33.6$ mF with $n_2 = 0.55$).
As already stated in chapter one, the impedance of CPE is defined as

$$Z_{CPE} = \frac{1}{[Q(j\omega)^n]} \tag{22}$$

The CPE$_1$ that replaced the diffusion impedance ($Z_w$) in the ideal Randles circuit is associated with a high $n_1$ value (0.95), describing the high porous nature of the electrode. The $n_2$ value (0.55) is approximately equal to the ideal Warburg behaviour of $n = 0.5$, describing the diffusion of ions through the thin film. From the Bode plot ($\log |Z|$ vs. $\log f$ (figure 4.13), the high frequency region yielded a slope of close to zero, characteristic of pure resistor, while the low frequency region yielded a slope of $<1.0$, characteristic of pseudocapacitor.
In general, these data clearly suggest that the impedance of this electrode varies from a pure resistor at high frequency to pseudocapacitor at low frequency region. Surprisingly, all attempt to replace the ideal $C_{dl}$ with a CPE (a real application situation) in the modeling circuit proved unsuccessful. As has elegantly been described recently by Orazem and Tribollet, the behaviour occurs as a result of distribution of time constants along either the area of the electrode surface (involving a 2-dimensional aspect of the electrode) or along the axis normal to the electrode surface (involving a 3-dimensional surface).
It is noteworthy that a 2-D distribution presents itself as an ideal RC behaviour, meaning that impedance measurements are very useful in distinguishing whether the observed global CPE behaviour is due to a 2-D or 3-D distribution or both. Thus, we may conclude that the observed impedimetric behaviour of the BPPGE-MWCNT-NiTAPc likely involve time constant distributions occurring along the area of the electrode as well as along the axis normal to the electrode. Also, from the other type of Bode plot (i.e., –phase angle ($\theta$) vs. log $f$, (figure 4.13), the $\theta$ was about $80^\circ$, which is less than the $90^\circ$ expected of ideal capacitive behaviour, thus further confirming the presence of CPE and pseudocapacitive nature of the BPPGE-MWCNT-NiTAPc.

Impedance spectroscopy was also performed for the MWCNT-tBuNiPc and the unsubstituted MWCNT-NiPc nanocomposites. The overall picture was the same as that for the MWCNT-NiTAPc hybrid where they all exhibit a shape characteristic of porous electrodes. The diagrams in figure 4.14 show the Nyquist and Bode plots for the MWCNT-NiPc nanocomposite. The phase angle for the hybrid is ca. $80^\circ$ indicative of an almost ideal supercapacitor. The synergy between the MWCNT and unsubstituted NiPc (figure 4.14 (b)) and tBuNiPc is also observed in the comparative Nyquist plots where the Pc and MWCNT contribute to a near-vertical line at low frequencies. The result is best shown by the Bode plots where the phase angle for the hybrids is close to $90^\circ$.
implying that the electrode reaction is dominated by double layer capacitance.

**Figure 4.14:** (a) Comparative Nyquist plots of BPPGE-NiPc (i) BPPGE-MWCNT-NiPc (ii) and BPPGE-MWCNT (iii) in 1M H$_2$SO$_4$ at 0.30 V, between 100kHz and 10mHz. (b) Bode plots for fitted BPPGE-MWCNT-NiPc data in 1M at 0.30 V. Comparative Nyquist plots of BPPGE-tBuNiPc (c) and Bode plots for the BPPGE-MWCNT-tBuNiPc (d)
4.2.3 Galvanostatic Charge/Discharge

The three hybrid supercapacitor electrodes were all tested under galvanostatic conditions. The electrodes were subjected to constant current charge/discharge cycling in the voltage range of -0.2 to 0.5 V at different current densities of 1 A g\(^{-1}\) to 30 A g\(^{-1}\). It is well documented that galvanostatic discharge is the most reliable and accurate method for evaluating the supercapacitance of electrodes.\(^{153,158,159}\) Figure 4.15 shows typical galvanostatic charge / discharge measurements obtained for the BPPGE-MWCNT (i) and BPPGE-MWCNT-NiTAPc (ii) at a current density of 1 A g\(^{-1}\); first charged to 0.5 V and then discharged to -0.2 V. The specific capacitance (SC), specific power density (SP) and specific energy (SE) based on the active materials (MWCNT, or NiTAPc or MWCNT-NiTAPc) were easily estimated from the discharge process using the established equations. (equations 14,15 and 16 from chapter 1)

\[
SC(F/g) = \frac{[i(A) \times \Delta t(s)]}{[\Delta E(V) \times m(g)]} \quad \text{.......................... (14)}
\]

\[
SE(Whkg^{-1}) = \frac{[i(A) \times t(s) \times \Delta E(V)]}{m(kg)} \quad \text{.......................... (15)}
\]

\[
SP(Wkg^{-1}) = \frac{[i(A) \times \Delta E(V)]}{m(kg)} \quad \text{.......................... (16)}
\]
It is well known that in an aqueous electrolyte, capacitance is predominantly determined by the hydrophilicity of the electrode materials, the stronger the affinity of the electrode materials; the higher the double layer capacitance.\textsuperscript{160} The high specific capacitance recorded for the MWCNTs should perhaps not be surprising considering the stringent acid-treatment procedure adopted in this work to functionalise our MWCNTs. For example, by merely stirring MWCNTs in 3:1 mixture of \( \text{H}_2\text{SO}_4/\text{HNO}_3 \) for just an hour, Kim \textit{et al.}\textsuperscript{160} reported the capacitance of MWCNTs immobilized onto a Ni mesh using PVDF solution in N-methylpyrrolidine as binder (and a 3-electrode system) to be 51.3 F g\textsuperscript{-1} in 1 M \( \text{H}_2\text{SO}_4 \). On the other hand, Frackowiak and co-workers\textsuperscript{9} did not

\textbf{Figure 4.15:} Typical charge-discharge cycles obtained at the BPPGE-MWCNT (i) and BPPGE-MWCNT-NiTAPc (ii) at 1 Ag\textsuperscript{-1}. Supporting electrolyte = 1 M \( \text{H}_2\text{SO}_4 \).
employ the long functionalisation times as used in our case, however, they reported values from 70 to 120 Fg\(^{-1}\) in 6 M KOH for a 2-electrode system and using PVDF binder.

Therefore, if we consider that we did not employ organic binders (which, to a certain extent, can impact negatively on the flow of ions and hence on the observed capacitance) and that capacitance value of a 2-electrode system is about 4 times less than for a 3-electrode system, the values reported by these authors\(^{160}\) are in close proximity to ours. Different masses were also loaded onto the electrode and the capacitance measured using CD and EIS to obtain an insight into the impact of different mass loadings (20-80 µg range) on capacitance. As seen in figure 4.16, the specific capacitance decreased slightly from the lowest mass loading (∼1320 Fg\(^{-1}\) at 20 µg) to the highest mass (∼907 Fg\(^{-1}\) at 80 µg).
Figure 4.16: Plot of specific capacitance versus mass loading of the MWCNT-NiTAPc determined from the (a) galvanostatic discharge at 1 Ag\(^{-1}\), and (b) electrochemical impedance spectroscopy at 0.3 V vs Ag|AgCl (sat’d KCl). Supporting electrolyte 1M H\(_2\)SO\(_4\).

It is interesting to observe from figure 4.16 that, within the limits of experimental errors, the SC of the MWCNT-NiTAPc nanocomposite is approximately 1000 Fg\(^{-1}\) at all mass loadings studied. The decrease in the SC at higher mass loadings (> 20 µg) may be attributed to the thickness of the film on the small geometric area electrode (0.196 cm\(^2\)) employed in this study. In fact, with the exception of the 20 µg loading, all subsequent loadings had to be carried out at several “drop-dry times” to get all the materials loaded on the electrode. Thick film is disadvantageous to
pseudocapacitive behaviour. As the material loading increases more of the electroactive species are buried within the thick film and become ‘redox-silent’. In addition, there will be problems with distribution of potentials within this thick surface-confined modified CNT material. Thus, as a compromise, we used 40 µg (~ 2 mg cm$^{-2}$) to compare the SC values of the three materials.

The maximum supercapacitive parameters were obtained at 1 A g$^{-1}$ and are summarised in Table 4.1. For the MWCNT-NiTAPc (n = 6, 95% confidence level) the following were obtained: SC = 981 ± 57 Fg$^{-1}$ or 200 ±12 Fcm$^{-2}$; SP = 701 ± 1 W kg$^{-1}$ and SE = 134 ± 8 Wh kg$^{-1}$. These values are more than twice higher than those of the MWCNTs alone.

Table 4.1: Supercapacitive properties of basal plane pyrolytic graphite electrode (BPPGE) modified with NiTAPc, acid-functionalised MWCNTs and MWCNT-NiTAPc.

<table>
<thead>
<tr>
<th>Electrode</th>
<th>Supercapacitive parameters</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>SC (F g$^{-1}$)</td>
</tr>
<tr>
<td>BPPGE-NiTAPc</td>
<td>198 ±13</td>
</tr>
<tr>
<td>BPPGE-MWCNTs</td>
<td>335 ± 18</td>
</tr>
<tr>
<td>BPPGE-MWCNT-NiPc</td>
<td>760 ± 15</td>
</tr>
<tr>
<td>BPPGE-MWCNT-tBuNiPc</td>
<td>440 ± 23</td>
</tr>
<tr>
<td>BPPGE-MWCNT-NiTAPc</td>
<td>981 ± 57</td>
</tr>
</tbody>
</table>

$^1$ Specific capacitance (F cm$^{-2}$) calculated per geometric area of the electrode, not the electroactive area.
From Table 4.1, the values of the specific capacitance are comparable to our estimated CV data. However, the values recorded with the galvanostatic discharging are more reliable given their low errors. In general, the SC value for the BPPGE-MWCNT-NiTAPc is indeed greater than most other half-cell or three electrode counterparts reported in the literature.\textsuperscript{161-163} For example, using the same experimental conditions as ours (i.e., 1 A g\textsuperscript{-1}, 1 M H\textsubscript{2}SO\textsubscript{4} with a 3-electrode system), Hulicova Jurcakova \textit{et al.}\textsuperscript{21} recently reported a value of 1117 µFcm\textsuperscript{-2} for the nitrogen-enriched carbon electrode.

It is interesting to note that our supercapacitance value of 135 mFcm\textsuperscript{-2} (200 Fcm\textsuperscript{-2} using 40 µg) is two orders of magnitude or more than a hundred-fold greater than the value described as ‘extraordinary supercapacitance’.\textsuperscript{21} The excellent supercapacitive behaviour of the MWCNT-NiTAPc may be related to the high surface area and porous nature of the nanocomposite that enhance the penetration of solution ions. Importantly, the thin film nature (small active mass used in this study) of the hybrid is advantageous as it shortens the distance which ions and electrons would have to travel to reach the electrode. In addition, the incorporation of pendant amine groups, capable of protonation at this low pH incorporates more positive charges into the NiTAPc macrocycle structure than simply the Ni(II)/Ni(III) metal centre, thus enhancing the interaction with electrolyte anions at the electrolyte-electrode interface and increasing
the observed double-layer capacitance compared to macrocycles without such additional charged groups.

Also, at all current densities of 1 – 30 A g\(^{-1}\) (i.e., 0.1 – 3 mF cm\(^{-2}\)) studied, the charge processes of the MWCNT-NiTAPc showed mirror images of their corresponding discharging counterparts, with no detectable IR drop even at much higher discharge current densities, suggesting a very small equivalent series resistance (ESR) of the electrode. The energy deliverable efficiency (\(\eta / \%\)) was obtained from equation 17 (from chapter 1). The energy deliverable efficiency of the MWCNT-NiTAPc nanocomposite at the \(~ 2\) mg cm\(^{-2}\) was \(~ 121\) % at 1 A g\(^{-1}\) and up to \(~ 100\) % at 1 A g\(^{-1}\).

\[
\eta (\%) = \frac{t_d}{t_c} \times 100 \quad \text{.................................................. (17)}
\]

To obtain an insight into the impact of phthalocyanine ring substituents on the supercapacitive behaviour, similar CV and CD experiments for NiTAPc-MWCNT were also performed with unsubstituted nickel (II) phthalocyanine (NiPc) and nickel (II) tetra-\textit{tert-butyl} phthalocyanine (\textit{tBuNiPc}). Interestingly, the NiTAPc gave superior capacitive data. As an example, at a current density of 1 Ag\(^{-1}\) (for \(~ 20\) \(\mu\)g loading) the specific capacitance decreases as NiTAPc (\(~ 243\) Fg\(^{-1}\)) > NiPc (\(~ 176\) Fg\(^{-1}\)) > \textit{tBuNiPc} (\(~ 126\) Fg\(^{-1}\)). This trend is accordance with the nitrogen contents of the three phthalocyanine species. In addition, the poorer value of the \textit{tBuNiPc} may be related to the absence of
nitrogens with any doner lone pairs to protonate at this acidic pH used in this work or the bulky tert-butyl group that may deform the phthalocyanine framework, thus reducing the supercapacitance. In addition, Kim et al.\textsuperscript{160} proved that alkyl chains reduce supercapacitance. Also, the NiTAPc-MWCNT is more stable on the BPPGE and highly reproducible galvanostatic data than the other nickel phthalocyanines studied. More importantly, NiPc and tBuNiPc and their MWCNT composites are less soluble in DMF than the NiTAPc based electrodes. This superior performance of the NiTAPc is not fully understood at this moment, but may be related to its nitrogen-containing groups, which have recently been observed to greatly enhance supercapacitive behaviour.\textsuperscript{141,164}

\textbf{Figure 4.17:} Typical charge-discharge cycles obtained at the (a) BPPGE-MWCNT (i) and BPPGE-MWCNT-NiPc (ii), (b) BPPGE-MWCNT (i) and BPPGE-MWCNT-tBuNiPc at 1 Ag\textsuperscript{-1} (0.1 mA cm\textsuperscript{-2}). Supporting electrolyte = 1 M H\textsubscript{2}SO\textsubscript{4}. 
The plots given in Figure 4.18 show the relationship between specific capacitance and current density. For the three materials, the highest specific capacitance was observed at low current densities and decreased significantly with increase in current density. This can be explained by the limited access of the pores for charge storage at high current densities by the ions. The contribution by the smaller pores to the overall capacitance is limited. The three kinds of material are therefore suitable for supercapacitors in lower current densities than $5 \text{ Ag}^{-1}$. A levelling of the curves is observed at high current densities.

![Graph showing specific capacitance values as a function of current density.](image)

**Figure 4.18:** Specific capacitance values as a function of current density.
4.2.4 Stability studies

The MWCNT-Nickel(ii) phthalocyanine electrodes were tested for stability at high current and were all found to possess remarkable stability over 1000 charge-discharge cycles. The stability of the nanocomposites were tested by continuous cycling at high current density (1mA/cm$^2$). A typical repetitive charge-discharge cycling for 1500 cycles is shown in figure 4.20. The figure clearly shows that the electrode is able to charge and discharge continuously without any significant loss in capacitance.

Figure 4.20: Variation of specific capacitance with the number of cycles at 10 A g$^{-1}$ (1mA cm$^{-2}$). Inset is a portion of the charge-discharge curves. Supporting electrolyte = 1 M H$_2$SO$_4$. 
A small decrease in the specific capacitance is observed for the MWCNT-NiTAPc nanocomposite in the first 100 cycles. A decrease in specific capacitance of 16.7% was observed. In the subsequent 500 cycles a decrease of 6% was observed.
4.3 Supercapacitive properties of electropolymerized film of Nickel (ii) tetra-aminophthalocyanine

Electropolymerized metalloporphyrin and metallophthalocyanine-coated electrodes have been reported expansively as efficient electrocatalysts for chemical, photochemical and analytical applications.\textsuperscript{165-170} Nickel macrocyclic complexes have received a lot of attention as conducting polymer films.\textsuperscript{171,172} Interestingly enough, despite the huge literature on the conductive ability and electrocatalytic properties of electropolymerized nickel phthalocyanine complexes, no report on nickel phthalocyanine polymers as potential supercapacitor electrode materials has been made. Motivated by the results in chapter 3.2 where a composite mixture of nickel (II) tetra-aminophthalocyanine (NiTAPc) and multi-walled carbon nanotubes (MWCNTs) exhibit supercapacitive behaviour, this section explores, for the first time, the capacitive properties of the electropolymeric films of NiTAPc supported on a MWCNT platform.
4.3.1 Cyclic Voltammetry

Figure 4.21 shows typical electropolymerisation process of the NiTAPc on BPPGE, showing growth of scan. As the scan number increases, typical features of electropolymerisation process become evident. The electropolymerisation process has been reported to occur via the oxidation of the amino groups and the N-N coupling resulting in an electroactive polymer on the electrode surface.\textsuperscript{173} Well defined peaks appear at around 0.45 V are observed.

![Cyclic voltammogram](image)

**Figure 4.21:** Cyclic voltammogram of 10 mM NiTAPc in DMF at 100 mVs\textsuperscript{-1} (50 scans increasing from inner to outer)
The same feature is also observed for the BPPGE-MWCNT-polyNiTAPc. Figure 4.22 shows the CV evolutions of the BPPGE-polyNiTAPc on continuous cycling in 0.1 M NaOH. As the cycling continues, the peaks grow steadily until they reach a steady state where no further growth is observed.

**Figure 4.22:** Modified CVs of BPPGE-MWCNT-polyNiTAPc, after 50 scans (increasing from inner to outer) from -0.2 V to 0.8 V in aqueous NaOH 0.1 mol L$^{-1}$ solution at 100 mV s$^{-1}$.

This is typical of cyclic voltammograms recorded for electroformed nickel macrocyclic-based films in alkaline solution and shows that the film has been formed as a result of the anodic polymerization of the
As the cycling continues, the peaks grow steadily until they reach a steady state where no further growth is observed as shown in figure 4.22. The CV shows one redox couple with formal potential \( (E_{1/2}) \) at \( \sim 0.42 \text{ V} \) vs Ag|AgCl, which is assigned to the Ni(II)/Ni(III) redox process of the nickel centre of the NiTAPc. The appearance of this signature redox couple confirms the successful integration of the polyNiTAPc onto the electrode. The same feature is observed for the MWCNT-polyNiTAPc electrode, except for the higher current response observed for the BPPGE-MWCNT-polyNiTAPc. It has been postulated that the polymerization of nickel macrocyclic complexes in alkaline media is very similar to that of the nickel hydroxide electrode.\(^{173}\) The nickel-nitrogen tetra coordination of the complex is thought to be lost in the polymerization process. Another postulate by Roslonek and Taraszewka proposes that upon electropolymerisation, the complexes are attached via oxo-bridges.\(^{174}\) Cyclic voltammetry study was carried out for all the electrodes in both acid (1 M H\(_2\)SO\(_4\)) and alkaline (1 M NaOH) conditions. Figure 4.23 presents typical cyclic voltammetric evolutions obtained.
**Figure 4.23:** (a) Comparative CV’s for bare BPPGE (i), BPPGE-MWCNT (ii), BPPGE-polyNiTAPc (iii) and BPPGE-MWCNT-polyNiTAPc (iv) in 1 M H$_2$SO$_4$. (b) Effect of electrolyte on capacitance: BPPGE-MWCNT-polyNi(OH)TAPc in acid (i) and in alkali (ii).

The capacitance of the polymeric film per unit area of the electrode ($C_{polym}/F\text{ cm}^{-2}$) was estimated in both the faradaic (ca 0.35 V) and the non-Faradaic regimes (-0.1 V versus Ag/AgCl) using equation (25)

$$C_{polym} (Fcm^{-2}) = \frac{I_{ch}}{\nu A} \quad \text{.......................... (25)}$$

where $I_{ch}$ is the charging current, $\nu$ the scan rate and $A$ the geometric area of the electrode. The capacitance was calculated for both the anodic and cathodic charging current and the average values reported. From the
comparative CVs in acidic medium, the capacitive behaviour of the electrodes follows as MWCNT-polyNiTAPc (97 mF cm\(^{-1}\)) > MWCNT-polyNi(OH)TAPc (50 mF cm\(^{-1}\)) > MWCNT (35 mF cm\(^{-1}\)) > polyNiTAPc (21.9 mF cm\(^{-1}\)). Figure 4.23 (b) shows the impact of electrolyte on the capacitance, where BPPGE-MWCNT-polyNi(OH)TAPc in acid medium is better than obtained in alkaline. Clearly, the CV recorded in acid has a larger current separation as compared to that recorded in alkaline medium.

The overall capacitance in acidic media is therefore higher than that observed in alkaline medium. Recently, Hulicova et al.\(^{141}\) observed a similar phenomenon for nitrogen-doped carbon materials studied for supercapacitor and found that these materials exhibit enhanced capacitive behaviour in 1 M \(\text{H}_2\text{SO}_4\) than in 1 M \(\text{KOH}\) solution. The authors interpreted the situation in terms of the nitrogen functionalities exhibiting stronger interactions with the \(\text{H}_3\text{O}^+\) than with the \(\text{K}^+\) resulting in higher capacitance. The same explanation may also hold in our case, possibly due to the interaction between the nitrogen groups of the phthalocyanine ring and the \(\text{H}_3\text{O}^+\) of the \(\text{H}_2\text{SO}_4\) electrolyte. Based on this observation, further studies were carried out with the more reliable galvanostatic charge-discharge method to unequivocally determine the capacitance values. Scan rate studies were also performed for the MWCNT-polyNiTAPc electrode
4.3.2 Galvanostatic charge-discharge experiments

It is well established that galvanostatic discharge represents the most accurate and reliable strategy for probing the supercapacitance of materials. Figure 4.24 shows the galvanostatic charge-discharge curves obtained. The specific capacitance (SC) was evaluated from the discharge curve using equation 14 from chapter 1 with the mass of the electrode being replaced by electrode area in cm$^2$:

$$SC \ (Fcm^{-2}) = \frac{I \times \Delta t}{\Delta E \times A} \ ........................................ (14)$$

The specific capacitance follows the same trend as the CV data: BPPGE-MWCNT-polyNiTAPc (112 mFcm$^{-2}$) > BPPGE-MWCNT-polyNi(OH)TAPc (84.9 mFcm$^{-2}$) > BPPGE-MWCNT (38.7 mFcm$^{-2}$) > BPPGE-polyNiTAPc (27.2 mFcm$^{-2}$). Also, as in the CV, the capacitance of the BPPGE-MWCNT-polyNiTAPc obtained in acid is higher than obtained in the alkaline medium.
The enhanced capacitance of the MWCNT-polyNiTAPc in acid medium may be interpreted in terms of the synergism between MWCNT and polyNiTAPc species (also observed in the CV profiles). It indicates that this electrode possesses features (mainly, conductivity and porosity) that permit free movement of the solution ions to and from the electrode/solution interface. The porous nature of the MWCNT-polyNiTAPc electrode is also observed in figure 4.25. Upon introduction of the MWCNTs, the globular structure of the polyNiTAPc is still evident in figure 4.25 (c) and the deposit shows a network between the MWCNTs and the polyNiTAPc in 4.25 (d).
**Figure 4.25:** SEM image of (a) bare BPPGE, (b) BPPGE-polyNiTAPc, (c) BPPGE-MWCNT-polyNi(OH)TAPc and (d) BPPGE-MWCNT-polyNiTAPc electrode surfaces.

The energy deliverable efficiency (\(\eta / \%\)) was obtained from equation 17 in chapter 1.

\[
\eta (\%) = \frac{t_d}{t_c} \times 100 \hspace{1cm} (17)
\]


The energy deliverable efficiency for BPPGE-MWCNT-polyNiTAPc is 105 ± 11%. The long-term cycling stability of the same electrode as potential supercapacitor material was also investigated. A typical repetitive charge-discharge cycling for 1000 cycles is shown in figure 4.26. It is clear from this figure that MWCNT-polyNiTAPc is capable of undergoing continuous charge and discharge with little or no significant loss in capacitance. This observation also suggests that MWCNT-polyNiTAPc material does not show significant structural changes during the course of charge-discharge processes.

**Figure 4.26:** Typical portion of the charge-discharge curves obtained during a repetitive cycling of the BPPGE-MWCNT-polyNiTAPc electrode at 10 mA cm\(^{-2}\) in 1 M H\(_2\)SO\(_4\) solution, showing excellent stability.
4.3.3 *Electrochemical Impedance Spectroscopy experiments*

To obtain further insights into the impact of the MWCNTs on the capacitive behaviour of the polyNiTAPc, a comparative EIS experiments were carried out for the BPPGE-polyNiTAPc, BPPGE-MWCNT and BPPGE-MWCNT-polyNiTAPc electrodes in 1 M H₂SO₄ conditions. The low-frequency differential capacitance \(C_d = 1/2\pi f Z''\) for each of the electrodes was obtained by running a single potential EIS experiment (biased at 0.25 V) MWCNT, polyNiTAPc and MWCNT-polyNiTAPc at a frequency range of 10 mHz – 100 kHz in 1 M sulphuric acid (Figure 4.27). In all cases, the maximum specific capacitance was obtained at ~ 0.25 V vs Ag|AgCl, with values decreasing as BPPGE-MWCNT-polyNiTAPc (~ 13.2 mF cm\(^{-2}\)) > BPPGE-MWCNT (~ 10 mF cm\(^{-2}\)) > BPPGE-polyNiTAPc (~ 3.17 mF cm\(^{-2}\)). This trend again agrees with CV and CD data. Clearly, the EIS data strongly deviate from the data obtained from the galvanostatic method, and are characteristic of conducting polymeric substances.\(^{147}\) The deviation in specific capacitances calculated from CD and EIS have been explained in section 4.2.2 of this chapter.
**Figure 4.27:** Comparative Nyquist plots for BPPGE-polyNiTAPc (i) and BPPGE-MWCNT-polyNiTAPc (ii) and MWCNT (iii) electrodes at 0.25 V. Inset: The expanded portion of the high frequency region.

The impedance spectrum was satisfactorily modeled using the electrical equivalent circuit involving the modified Randles circuit shown in figure 4.28. The circuit (R (Q [RQ])) used to fit the EIS data involved a solution resistance (R_s = -17.6 Ω), a constant phase element (CPE_1 = 2.2 μF with n_1 = 1), an electron transfer resistance of (R_2 = 23.78 Ω) and constant phase element (CPE_2 = 4.8 mF with n_2 = 0.9). The CPE_1 that replaced the diffusion impedance (Z_w) in the ideal Randles circuit is associated with a high n_1 value (n_1 = 1), describing the high porous nature of the electrode.
The $n_2$ value (0.90) is approximately equal to the ideal capacitive behaviour of $n = 1$.

![Electrical equivalent circuit used in fitting the experimental EIS data obtained for the BPPGE-MWCNT-polyNiTAPc.](image)

**Figure 4.28:** Electrical equivalent circuit used in fitting the experimental EIS data obtained for the BPPGE-MWCNT-polyNiTAPc.

As seen in Figure 4.27, the MWCNT-polyNiTAPc shows near-vertical line as expected for pseudocapacitor. The “knee” or “onset” frequency ($f_o$), which is a measure of the power capability of a supercapacitor, decreases as MWCNT-polyNiTAPc (1.604 kHz). The response time of the capacitor is about 0.6 ms, suggesting that most of the stored energy of MWCNT-polyNiTAPc as a capacitor material is accessible at frequencies as high as 1.604 kHz. This is very interesting considering that most commercially available supercapacitors, including those specifically designed for higher power applications, operate at frequencies less than 1 Hz.\textsuperscript{156}
CHAPTER FIVE

CONCLUSIONS
The capacitive behaviour of three different carbon nanotube-nickel (ii) phthalocyanines nanocomposites have been investigated for the first time in 1M H$_2$SO$_4$. CNT-MPc modified electrodes showed excellent supercapacitive behaviour and remarkable stability. When compared to unsubstituted NiPc/MWCNT or nickel (II) tetra-tert-butylphthalocyanine (tBuNiPc)/MWCNT, NiTAPc/MWCNT exhibited superior supercapacitive behaviour.

The effect of substituent groups on the phthalocyanine macromolecule was also observed. The MWCNT-NiTAPc electrode had the highest capacitance possibly due to the contribution by the nitrogen groups located at the periphery of the phthalocyanine molecule. The contribution by pseudocapacitance effects to the overall capacitance values cannot be overlooked. Alkyl groups on the tBuNiPc macromolecule however have been shown to block the pores in the nanocomposite essential for charge storage. The results also show the contribution by the MWCNTs to the overall capacitance.

Electropolymeric film of a phthalocyanine polymer, supported on MWCNTs platforms, has also been interrogated for the first time. Cyclic voltammetry, galvanostatic charge-discharge cycling and impedimetric techniques confirmed that the MWCNT-polyNiTAPc based electrode exhibit excellent supercapacitance, which may be ascribed to the positive synergism between the MWCNT and polyNiTAPc. The finding reported in
this work is crucial as it could open doors for the investigation of many other electropolymers of metal tetraaminophthalocyanine (MTAPc) complexes and related metal N4 macrocyclic organometallic complexes for application in supercapacitors.

Given the high number of existing MPc complexes and new ones constantly reported in the literature, and the importance of supercapacitors in energy development, we envisage that this novel study is likely to spark research interests in the supercapacitive properties of CNT-MPc hybrid electrodes. Although we have managed to reach a few important conclusions in this work, several aspects still need to be further investigated. Listed below, are some of the suggestions for future work:

1. The effect of different substituent groups on the phthalocyanine molecule and contribution to overall capacitance. e.g. –SO$_3$, phenyl groups etc.

2. The effect of different electrolytes on the capacitive properties of the MWCNT-MPc nanocomposites such as ionic liquids and alkaline medium need also be investigated.

3. The growth of carbon nanotubes on the electrode surface by CVD can produce uniform surface coverage and a desired mesoporous surface by controlling the pore size and pore distribution thus providing a high conducting and high surface area support.
4. Since nitrogen groups have been reported to increase capacitance in carbon material, functionalisation of the MWCNTs with an aim to increase the number of nitrogen groups on the CNT walls may enhance the supercapacitance.

5. An increased electrode surface area and the employment of the more practical two electrode system should be explored.
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