Chapter 7

The importance of nanostructured materials for energy storage/Conversion

M.J. Mochane*1, T.C. Mokhena1, L.Z. Linganiso1, T.E. Motaung1, T.H. Mokhothu2, and A. Mtibe2

1Department of Chemistry, University of Zululand (KwaDlangezwa Campus), Private Bag X1001, 3886, South Africa
2Council for Scientific and Industrial Research (CSIR), Nonwovens and Composites Research Group, 4 Gomery Avenue, Summerstrand, South Africa

Abstract

The finite nature of fossil fuels and concerns over greenhouse effect make the effective utilization of energy and conservation key issues. Using environmentally friendlier materials to either produce or store energy become extremely important to overcome the disparity between the energy supply and demand. Nowadays, the replacement of conventional energy with renewable sources such as hydrogen storage, thermal storage, lithium batteries and supercapacitors received relentless interest. Nanomaterials garnered much interest because of their superior properties such as large surface area, favourable carrier properties, mechanical stability, and both high electrical and thermal conductivity resulting from their nanosized structures. Due to their structural diversity and excellent properties, nanostructured materials and their composites have been extensively studied for energy storage/conversion related applications including photo-catalyst for water splitting, hydrogen storage materials, and thermal energy storage materials, as well as electrode materials for various lithium batteries and supercapacitors. This chapter summarizes the recent developments, limitations and challenges of the nanostructured materials for energy storage/conversion applications. The limitations and challenges of using nanostructured materials to enhance the performance of the energy storage materials have also been addressed.

Keywords: fossil fuels, energy storage, lithium-ion battery, greenhouse gases

Corresponding authors: mochane.jonas@gmail.com; mokhenateboho@gmail.com
7.1 Introduction

The global utilization of fossil fuels grows at a disturbing and unsustainable rate. The release of greenhouse gases and other toxic pollutants are at an unacceptable elevation which is not good for environmental concerns. In most regions of the world, direct solar radiation is thought as a prospective source of energy storage. The future stability of our society depends on the alternative energy sources that are renewable and environmental friendly [1,2]. Researchers globally are investigating on finding a new and renewable energy sources. Energy storage is important in a sense that it reduces the gap between energy supply and demand; it further enhances the reliability and performances of energy systems [3]. This could lead to saving of fuels and making cost effective systems by storing the wastage of energy. To make sense of the whole discussion, it is clear that storage will probably enhances the performance of a power plant for an example, thereby loading level and higher efficiency which would result in energy conservation and reduced generation cost overall [4].

Throughout our existence as human beings, our advances have been based on the discovery, development and employment of new materials. Materials have been considered as the main the core of all technologies as most technological discoveries have been achieved through the development of new materials [5]. The recent technologies are such that the nanomaterials are heavily starting to play a significant in the production of new fields of science and nanotechnologies. In order for us to develop technological discoveries in the production of effective renewable energy storage/conversion, we need nanomaterial to fully develop the potential of renewable energy. As such, the potential of nanomaterials must be discovered at a more fundamental level.

Nanotechnology applications and development in the field of energy storage include (i) nanostructured materials and separators for super capacitors and batteries. This includes lithium ion batteries for automotive drives and stationery mass materials and electrolytic systems, designation of new types of batteries such as lithium air and lithium sulphur. Carbon nanotubes based electrode materials for super capacitors and batteries (ii) nanoporous adsorption storage for mobile gas storage (hydrogen, natural gas). A battery is known as container consisting of one or more cells, in which chemical energy is converted into electricity and used as a source of power. In this chapter different types of energy storage
which include lithium-ion battery, sodium air battery, supercapacitor and metal hydrid storage are discussed.

7.2 Classification and production of nanomaterials

Nanotechnology is a core of multidisciplinary science that has roots in almost all the fields of Science and Technology. It discusses the production of structures at a very small scale and on the level of nanometers. Nanoscience is the research of fundamental principles of structures having sizes between 1 and 100 nm. Generally, nanomaterials are the main core of nanoscience together with nanotechnology. The study of nanomaterials is a broad and exciting area of research and development that has been growing significantly worldwide in the past few years. Nanomaterials are not really new materials, even though they may appear as new materials. Volcano dust has been existing for some time in nature. The release of ash during volcano emission can reach temperatures over 1,400°C and has a very complex structure consisting of solid and liquid particulate matter lifted by the hot gas current [6]. The distribution of ash into the atmosphere resulted in a gas temperature lowering and gas composition changes which led to the accumulation of the particles as either clusters thorough chemical reactions or based on electrostatic forces of attraction.

There are also other man made nanoparticles such as carbon based materials and fumed Titania (TiO$_2$). One of the most common ways of classifying nanomaterials is to differentiate them according to the production method. There are two kinds of methods used for the production of nanomaterials which includes physical and chemical methods. The physical-mechanical method produces nanoparticles in the nanometer sized range by milling or grinding larger particles (Figure 7.1). A common name for this method is the top-down way of making nanomaterials. The mechanical-physical crushing methods for producing nanoparticles involve various milling, whereby milling uses thermal stress and is energy intensive.
The physical method is subdivided into mechanical and phase-change methods as depicted in Figure 7.2. It is noted that both kinds of physical methods doesn’t results in the chemical change for the production of nanoparticles. The physical-phase change method involves the production nanoparticles via phase change process. Typical examples of physical-phase method include direct precipitation, whereby a material in solution is precipitated into a powder of nanoparticles, and as well as laser ablation in which the vaporized material is condensed into a solid nanomaterial.
Figure 7.2  Schematic presentation of the methods used for the production of nanomaterials

Figure 7.3 shows an experimental set-up for colloid preparation by laser ablation in solution. The basic concept of laser ablation occurs as follows: When the laser beam is concentrated on the surface of a target material in liquid phase, the temperature of the irradiated spot quickly increases, vaporizing the target material. Due to the increasing in temperature, the collision of the vaporized species and the neighbouring molecules result in the excitation of the electron state together with light emission and production of electrons as well ions, forming laser-induced plasma spreading out in a shape resembling a feather [7]. The plasma structure depends on a couple of factors such as the target material, ambient media, pressure and laser condition. One could mention that the low-pressure laser ablation gas is preferred for generation of a large plume and as a result small particles are produced. The ambient media happen to be important in laser ablation experiments because laser generated particles as mentioned elsewhere in this document react with the surrounding molecules to form complexes such as oxides and other unwanted materials.
Generally, the chemical methods require that the nanomaterial should be synthesized from initial material which is chemically different from the target material. The chemical method is typical bottom-up approach and different types of chemical methods are shown in Figure 7.2. The bottom-up method is built on physico-chemical principles of molecular or atomic self-organization. The advantage of this method is that it produces selected complex structures from atoms or molecules with better controlling sizes of shapes and size range.

7.3 Different types of energy storage nanomaterials

7.3.1 Lithium-ion battery

Rechargeable battery technology has come a long way since its introduction in the early 1990s. According to the literature [8], there is a moderate increase in the energy density of lithium ion systems (Figure 7.4) in the last two decades since their introduction. There are a lot of rechargeable batteries which includes nickel–metal hydride, lead acid, lithium ion
which has been used for electric vehicle application. Amongst the above mentioned energy storage batteries, lithium ion battery has been the most attractive choice for automobile industries despite the fact that it is expensive [9]. Lithium ion batteries have been widely used since the early 1990s. Lithium-ion battery has shown productive performance in the application of automobile, cellular phone, note book computers because of better power and energy density compared with other energy storage devices. Furthermore, the lithium-ion battery has been significantly used because of attractive characteristics such as high efficiency, long cycle life, low discharge rate and high voltage. Another emerging application for Lithium-ion technology is in battery electrical energy storage systems for smart grids that are powered by traditional energy sources like coal, as well as intermittent renewable energy sources like solar and wind.

![Lithium ion energy density improvements since 1991](http://dx.doi.org/10.5772/57139)

**Figure 7.4** Lithium ion energy density improvements since 1991. ©2014 Rohan JF, Hasan M, Patil S, Casey DP, Clancy T. Published in [8] under CC by 3.0 license. Available from: http://dx.doi.org/10.5772/57139.

Ordinary Lithium-ion battery starts as a 10-50 micron sized particles which are coated onto aluminium or copper current collectors together with conductivity enhancers and binders. **Table 7.1** illustrates the main constituents of lithium-ion battery and their characteristics. The most employed cathode electrodes include lithium cobalt oxide, lithium nickel oxide, Lithium nickel cobalt aluminium oxide, Lithium manganese spinel, and Lithium iron phosphate. Carbon-based materials have been the most favoured choice for anodes, with some version of graphite being utilized in a majority of the commercially available lithium-ion batteries. Natural graphite is the obvious choice as anode material compared to other carbonaceous
materials because it is inexpensive. Moreover, the graphite anode electrode display a high capacity through its high degree of crystallinity as well as less profile compared with lithium [10,11]. Graphite as a dominating anode material consist of two phases i.e. predominantly hexagonal (2H), with a small fraction of a rhombohedral (3R) phase which are responsible for the intercalation of lithium ions [10,11].

**Figure 7.5** shows a schematic presentation of typical lithium-ion battery. The cell consists of five regions, including composite negative electrode (anode), composite positive electrode (cathode), separator, and two electrode current collectors; made of copper and aluminium respectively. The two electrodes (anode and cathode) are divided by an electrolyte separator such as lithium hexafluorophosphate. The following half-reaction takes place at cathode and anode respectively:

\[
LiMO_2 \leftrightarrow Li_{z-2} MO_2 + zLi^+ + ze^- \quad (7.1)
\]

\[
zLi^+ + ze^- + 6C \leftrightarrow Li_2C_6 \quad (7.2)
\]

Equations 7.1 and 7.2 are in units of moles, \(z\) is the coefficient for a complete reaction, whereas M depicts the metal in which lithium oxide is formed.
During the process of charging, the voltage exerted across the electrodes proceeds the half reaction in the forward reaction i.e. from cathode to anode as it is depicted in Figure 7.5. The metal in the lithium metal oxide which is represented in the cathode is reduced to form lithium ions which are liberated in the electrolyte. The liberated ions diffuse through the electrolyte and are received into the carbon/graphite anode. During the formation of ions, electrons are produced to provide necessity for charge neutrality. According to the literature the electrolytes does not necessarily facilitate the conduction of free electrons through the system [12]. The electrons are moved across a wire connecting the two electrodes to provide necessary electrons required for the insertion of Li ions into the anode. During discharge, there is deintercalation between the negative electrode particles and solution. The electrons travel from anode to cathode during discharge process which causes positive current to originate from the cathode, as result the cathode act as positive terminal of the cell. Energy is stored in the electrodes in the form of Li-intercalation compounds.
Table 7.1 Major component of lithium-ion batteries and their characteristics [9]

<table>
<thead>
<tr>
<th>Abbrev</th>
<th>LCO</th>
<th>NLO</th>
<th>NCA</th>
<th>NMC</th>
<th>LMO</th>
<th>LFP</th>
<th>LTO</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>Lithium cobalt oxide</td>
<td>Lithium nickel oxide</td>
<td>Lithium nickel cobalt aluminium oxide</td>
<td>Lithium nickel, manganese cobalt oxide</td>
<td>Lithium manganese spinel</td>
<td>Lithium iron phosphate</td>
<td>Lithium titanate LMO, NCA</td>
</tr>
<tr>
<td>Positive electrode</td>
<td>LiCoO₂</td>
<td>LiNiO₂</td>
<td>Li(Ni₀₈₅Co₀₁₆₃₅Al₀₀₅)O₂</td>
<td>Li(Ni₀₃₃Mn₀₃₃Co₀)O₂</td>
<td>LiMn₂O₄</td>
<td>LiFePO₄</td>
<td>LMO, NCA</td>
</tr>
<tr>
<td>Negative electrode</td>
<td>Graphite</td>
<td>Graphite</td>
<td>Graphite</td>
<td>Graphite</td>
<td>Graphite</td>
<td>Graphite</td>
<td>Li₄Ti₅O₁₂</td>
</tr>
<tr>
<td>Cell voltage (V)</td>
<td>3.7-3.9</td>
<td>3.6</td>
<td>3.65</td>
<td></td>
<td>3.8-4.0</td>
<td>4</td>
<td>3.3</td>
</tr>
<tr>
<td>Energy Density (Wh/kg)</td>
<td>150 mA h/g</td>
<td>150</td>
<td>130</td>
<td>170</td>
<td>120</td>
<td>130</td>
<td>85</td>
</tr>
</tbody>
</table>
The advantages of nanomaterials in lithium-ion battery applications

The designation of a nanomaterial electrode as a Lithium-ion storage material would provide significant improvement in energy, power, and cycle life. The advantage of using nanoparticles as an electrode for lithium-ion battery applications for active energy storage can be seen in two (i) reduced diffusion distance for the lithium-ion particle core to the surface where it transfers to the electrolyte and (ii) nanoparticles provides high surface area because of their size as a result there will be higher electrode-electrolyte contact area [13]. The reductions in the particle size into the nanoscale significantly reduce the mechanical stresses caused by volumetric expansion and contraction during charge and discharge. The majority of lithium-ion cathode materials such as LiCoO₂, Li(Ni₀.₃₃Mn₀.₃₃Co₀.₃₃)O₂ and LiMn₂O₄ consists of lithium and metals more especially the transition ones. The complex compound i.e. lithium and other transition metals may be required to be heated in some cases to achieve the desired oxide or phosphate composition. Heat treatment of large particles in comparison to smaller particles may be heavy and requires large-scale of energy, with a possibility of a different thermal characteristics in comparison to the bulk. If the particle size is large, there will be improper heat treatment which will result into nonhomogeneous composition in the material and hence decline in performance. The use of nanomaterials makes it is easier to sustain a homogenous thermal behavior throughout the material and thereby producing homogenous composition without applying a lot of energy to the complex material.

7.3.1.2 Carbon based nanomaterials for lithium-ion batteries

7.3.1.2.1 Carbon based materials

Carbon is the sixth element in the periodic table of elements, and it became very important in scientific research because it can combine with itself and other chemical elements in different ways [14]. Carbon exists in three forms, known as amorphous carbon, graphite and diamond. In nature, carbon is available in more quantity as coal. There are also man-made carbon materials based on the required application which include: synthetic graphite, synthetic diamonds, adsorbent carbon, cokes, carbon black, and diamond-like carbon [14]. Diamond, graphene, graphite and fullerenes related materials (which include carbon nanotubes and carbon nanofibers) are well allotropes of carbon. Carbon based materials are preferred
choices in many applications including energy storage in lithium-ion batteries for energy storage in comparison to metallic materials because of high thermal conductivity values, less corrosion and cyclic reversibility [15]. Moreover, carbon based materials have the ability to provide active reaction sites and high specific area which allows the addition of stable active materials for high or better electrochemical performances [15].

7.3.1.2.2 Graphene nanoparticles

For the past three decades since the breakthrough of anode materials for lithium secondary battery resulting in the birth and commercialization of lithium ion battery, research on anode materials has been the main focus. There are a lot of anode materials that has been investigated so far including graphitic carbon, amorphous carbon, nitrides, tin oxides and tin-based alloys [16]. However, to our knowledge carbon based materials are still the dominating one available in the market. The reason for the use of carbon materials in this electrode is because (i) they exhibit both higher specific charges and more negative redox potential than most metal oxides (ii) Show better cycling performance than Li alloys because of their good dimensional stability. The insertion of lithium ions into the carbon based materials is known as the intercalation process according to Equation 7.3 below:

\[ \text{Li}_a \text{C}_n \leftrightarrow a\text{Li} + ae + \text{C}_n \]  (7.3)

The carbon host material has a good electrochemical reduction; as a result the lithium ions can easily penetrate into the carbon and form a lithium/carbon intercalation compound Li$_a$C$_n$ and the reaction in equation is reversible. Carbon based materials that are capable of reversible lithium intercalation can be classified into two groups i.e. graphitic and non-graphitic. Graphitic carbons are defined as carbon materials consisting of the element carbon in the allotropic form of graphite. From crystallographic definition, graphite is known as carbons having a layered lattice structure with perfect stacking order of graphene layers. The non-graphitic carbonaceous materials consist of carbon atoms that are mostly arranged in a planar hexagonal network without being close to forming crystallographic order in the c-direction. Graphite has been the main commercially used anode for lithium ion batteries as shown in Table 7.1. According to the literature, graphite experiences a limited theoretical capacity of 372 mA h g$^{-1}$ which is less than what is required of high energy capacity [17]. A
lot of attempt was done in developing other carbonaceous materials in order to obtain better performance. Graphene showed some distinctive advantages compared to graphite in energy storage applications. Graphene is a single-atom thick, two-dimensional sheet of \( \text{sp}^2 \) bonded carbon (Figure 7.6). The specific surface area of graphene is 2620 m\(^2\)g\(^{-1}\) which is much higher than that of graphite with values approximately 1300 and 10-20 m\(^2\)g\(^{-1}\). The large surface area of the graphene nanoparticles in comparison to the graphite particles provides more electrochemical reaction active sites for energy storage. In comparison to graphite, graphene nanosheets are also flexible which is beneficial for constructing flexible energy storage [18].

![Figure 7.6](image)

**Figure 7.6** The three forms of carbon: 0-dimensional, fullerene \(C_{60}\); 1-dimensional, single-walled carbon nanotube; and 2-dimensional, graphene sheet. Reprinted with permission from [18]. Copyright (2009) John Wiley and Sons

A lot of studies investigated the electrochemical properties of graphene as an anode for lithium ion batteries (LIB) and the results are summarized in Table 7.2. Guo *et al.* [19] reported that the graphite nanoparticles (GNS) exhibited a high reversible capacity of 672 mAh g\(^{-1}\) and a good cyclic performance. The authors attributed this observation to the storage of lithium on both sides of the graphene surface. Furthermore, another study [20] showed the storage of lithium ion on both surfaces of the GNS as well as the edges and covalent sites [20]. One of the advantages of graphene nanoparticles is the large surface areas which can provide more electrochemical reaction active sites for energy storage. The lithium storage
properties of graphene nanosheets (GNS) materials as high capacity anode material for rechargeable lithium battery were investigated by Yoo and co-workers [21]. The authors reported a specific capacity of GNS to be approximately 540 mAh g\(^{-1}\) which is much higher than that of graphite as it is revealed in Figure 7.7. The authors from the same study observed higher capacity values (730 and 784 mAh g\(^{-1}\)) by embedding CNTs or fullerene macromolecules into the graphene layers to increase the interlayer distance which probably provided additional sites for the accommodation of lithium ions. The electrochemical properties of graphene paper as anode electrode in lithium batteries was investigated in study [22]. Generally, the graphene papers were found to be mechanically strong and electrically conductive with a very high young modulus value (41.8 GPa). However, the electrochemical properties were found to be poor in comparison to the mechanical properties of the graphene paper. The discharge capacity was initially reported to be 298 mA h g\(^{-1}\) which was less than the theoretical capacity of graphite, decreasing to 240 mA h g\(^{-1}\) after 50 cycles. The results conclude that the graphene paper is not suitable for application as the anode material in a rechargeable lithium ion battery.

![Figure 7.7](image)

**Figure 7.7** (a) Relationship between the d-spacing and charge capacity of GNS families and graphite. (b) Charge/discharge cyclic performance of (a) graphite, (b) GNS+CNT, and (d) GNS + C\(_{60}\). Reprinted with permission from [21]. Copyright (2008) American Chemical Society.

The method of preparation of graphite nanosheets has significant impact on the electrochemical properties of the resultant LIBs. Lian et al. [23] investigated large reversible capacity of high quality graphene sheets (4 layers) as an anode material for lithium-ion batteries. The authors reported the first specific capacity of the graphene sheets as high as
1264 mAh g\(^{-1}\) (Figure 7.8) at a current density of 100 mAh g\(^{-1}\). The reversible capacity of graphene sheets with fewer layers is much higher than the previously reported literature [21] with graphene nanosheets (540 mAh g\(^{-1}\) at low current density of 50 mA g\(^{-1}\)). Comparison of the two studies revealed that the prepared graphene sheets with fewer layers (4 layers) have an intensive potential as a candidate of anode materials with high reversible capacity, good cycle performance and high rate discharge/charge capability. A possible reason for higher electrochemical properties for fewer graphite nanosheets was attributed to a large surface area and curled morphology of graphene sheets with fewer layers provided more lithium insertion active sites, such as edge-type sites and nanopores. Similarly, Wang and co-workers [24] reported that less graphite nanoparticle layers (5 layers) produced at low temperature (300°C) showed remarkable electrochemical performance in comparison to 13 layers obtained at 800°C. This was attributed to a higher specific surface area of lesser GNS layers than more graphene layers. In another study [25], graphene nanosheets were prepared from graphite oxide by two different methods i.e. Thermal exfoliation at different temperatures and wet chemistry using aqueous N\(_2\)H\(_4\) and KBH\(_4\) as the reducing agent. Generally, the two systems produced a high irreversible specific capacity irrespective of the method used for preparation of GNS. However, when comparing the two reducing agents, the GNS synthesized with N\(_2\)H\(_4\) showed better performance in comparison to KBH\(_4\). The authors also reported that the GNS produced with KBH\(_4\) and the thermal temperature of 800°C resulted in poor electrochemical performance of the GNS.

![Figure 7.8](image.png)  
Figure 7.8  Cycle performance of graphene sheets at the current densities from 100 mA g\(^{-1}\) to 1000 mA g\(^{-1}\). Reprinted with permission from [23]. Copyright (2010) Elsevier
Graphene nanoparticles just like any other nanoparticles suffer from agglomeration and restacking due to the van der Waals as a result decreasing the electrochemical performance of graphene nanoparticles in LIBs [26]. The following section discusses recent progress in the surface modification of graphene anodes by metals and/or metal oxide.

### 7.3.1.2.3 Surface modification of Graphene nanoparticles anodes

Recent studies have investigated various methods of surface modification of graphene electrodes such deposition of metals, metal oxides, coating with polymers [27-30]. Depending on the modification method, the following improvement may be attained: (i) smoothing the active edge surfaces by removing defects on the graphene surface (ii) forming a dense oxide layer on the graphene surface (iii) increasing the electronic conductivity (iv) Inhibiting structural changes during cycling. According to a reported literature above, deposition of certain metals or metal oxides which are host sites for lithium storage and improvement of the capacity of the anode. Several electrochemical properties (anode electrode) of the graphene with other nanoparticles or metal oxides are shown in Table 2 designed from different literatures. A lot of literatures reported [27,31-34] on the graphene/metal oxides for lithium ion batteries. **Figure 7.9** shows a typical preparation of the graphene/metal oxides composites.

![Figure 7.9](image)

**Figure 7.9** Schematic of the preparation of graphene/metal oxide composites with synergistic effects between graphene and metal oxides [26]. Copyright (2011) Elsevier

The aim of using both graphene and metal oxides for anode electrode is to lower the electrode problems of the individual components being the main active site, however utilizing the advantages of both graphene and metal oxide as active sites to improve the overall electrochemical properties of the system. In the composite both graphene and metal oxides provides different functions i.e. Graphene offer chemical functionality and compatibility whereas metal oxides provide high capacity. The resultant composite similar to a combination of blends provide a new material which is a combination of graphene and metal
oxide with new properties and functionalities. Generally, from the expectation point of view, metal oxide is expected not decrease agglomeration of graphene nanosheet only but also to increase the surface area of graphene nanoparticles leading to better electrochemical properties. Cobalt monoxide (CoO) quantum dot/graphene nanosheet composite was prepared with facile ultrasonic synthesis for LIBs [32]. The resultant CoO/graphene composite showed high performance as anode material for lithium ion battery with a reversible specific capacity of 1592 mAh g\(^{-1}\) after 50 cycles. The improvement in electrochemical properties of the system was ascribed to a well dispersed nanosized CoO quantum dots on conductive graphene matrix (Figure 7.10) which enhanced the active sites for lithium insertion, shorten the diffusion length for lithium ions which is favourable for high capacity and rate capacity.

![Figure 7.10](image)

**Figure 7.10**  (A) XRD pattern, (B) TGA, and (C,D) SEM images of the CQD/GN composites Reprinted with permission from [32]. Copyright (2012) American Chemical Society.

Most recently, stannic oxide (SnO\(_2\))/graphene nanocomposite paper (as both anode for lithium ion battery) were synthesized by the so called facile hydrothermal method [27]. The prepared nanocomposite in this case was chosen with the aim of forming flexible lithium-ion batteries. The flexible SnO\(_2\)/graphene paper showed a discharge capacity of 1195 mAhg\(^{-1}\) after 10 cycles and maintained at 1094 mAhg\(^{-1}\) after 80 cycles. Based on the result above it was concluded that the SnO\(_2\)/graphene paper is one of the candidate for flexible lithium-ion batteries. Similarly, graphite nanoparticles coated with α-Fe\(_2\)O\(_3\) [29] improved the electronic conductivity and diminishing the volume expansion, also proving enough active sites for lithium storage and shorten the lithium ion diffusion pathway. This has suggested that the α-
Fe$_2$O$_3$ coated graphite nanoparticles composite can be used as a potential anode material for high cyclic stability and long life lithium ion battery. The electrochemical properties of recent studies of graphene/metal oxide composite are shown in Table 7.2. Lately, a lot of studies investigated the performance of lithium ion battery in hybrid systems. Different systems (hybrids) were employed such as nanowire-graphene, Bi$_2$Te$_3$-nanoplate/graphene nanosheets [35], and phosphorus-graphene nanosheet hybrids [36] as anodes for lithium-ion batteries. By hybrid materials we are of the understanding that the composites consist of two constituents that are at nanometer or molecular level. A successful phosphorus-graphene nanosheet hybrid was prepared by ball milling method which is regarded as the low-cost method [36]. In this study, the phosphorus particles were grounded into smaller particles and hybridized with graphene nanosheets to form a 3D integrated structure. The authors reported a high specific capacity of 2362 mAh g$^{-1}$ and outstanding capacity retention after approximately 300 cycles at room temperature. The specific capacity of 1623 mAhg$^{-1}$ was obtained at elevated temperatures (60 °C) after 200 cycles. Graphene nanosheet were also mixed with different complexes and doped with different materials for improving the electrochemical properties of LIBs and the results also are summarized in Table 7.2. Guo and co-workers [37] investigated the layer-by-layer assembled composite of graphene nanosheets (GNSs) wrapped in monodisperse cobalt sulfide (CoS$_2$) nanocages for anode performances of lithium ion batteries. The monodisperse nanocages provided additional space to allow the volume changes, shorten lithium diffusion path and reduce agglomerations. As a result, the authors reported a high specific capacity 800 mAhg$^{-1}$ after 150 cycles at 100 mAg$^{-1}$ and 697 mAhg$^{-1}$ after 300 cycles at 500 mAg$^{-1}$. Tang and co-workers [29] investigated the silicon nanoparticles enwrapped with N-doped carbon and combine with N-doped graphene and carbon nanotubes as an anode material for lithium ion battery. The composite delivered magnificent cyclic retention of 892.3 mAh g$^{-1}$ after 100 cycles. The improved electrochemical performance was attributed to the well protected silicon nanoparticles and excellent conductivity provided by N-doping graphene and carbon nanotubes which may release the stress of the silicon due to volume expansion and decrease the transfer resistance of lithium ion in the anode electrode. However, most recently, FeP@reduced graphene oxide (FeP@rGO) prepared at mass ratio of of 1:5 (graphene oxide to FeCl$_3$.6H$_2$O) delivered a reversible discharge of 1000 mAh g$^{-1}$ after. It should be noted that in this case graphene oxide which is a layer of graphene with oxygen atoms was the one attached to the FeCl$_3$.6H$_2$O. Improved electrochemical properties might open options for FeP@reduced graphene oxide as a potential application for advanced LIBs.
Table 7.2 Electrochemical properties of graphene and its synergy with other nanoparticles for lithium ion battery from recent studies

<table>
<thead>
<tr>
<th>Graphene or graphene + other nanoparticles (anode)</th>
<th>Performance improvements</th>
<th>Refs.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cu-supported graphene nanoflakes</td>
<td>Specific capacity of 165 mAh g(^{-1}) over 80 charge-discharge cycle</td>
<td>[30]</td>
</tr>
<tr>
<td>graphene nanosheets (GNS)</td>
<td>Capacity of 540 mAh g(^{-1}) which is higher than that of graphite</td>
<td>[21]</td>
</tr>
<tr>
<td>GNS + CNTs or fullerene</td>
<td>higher capacity values (730 and 784 mAh g(^{-1})) by embedding CNTs or fullerene</td>
<td>[21]</td>
</tr>
<tr>
<td>SnO(_2)-graphene paper</td>
<td>Discharge capacity of 1195 mAh g(^{-1}) after 10 cycles</td>
<td>[27]</td>
</tr>
<tr>
<td>NiCo(_2)O(_4)/graphene</td>
<td>Higher reversible capacity and better cycling stability than NiCo(_2)O(_4)</td>
<td>[34]</td>
</tr>
<tr>
<td>MnO@graphene</td>
<td>Reversible capacity as high as 989.8 mAhg(^{-1})</td>
<td>[33]</td>
</tr>
<tr>
<td>Fe(_2)O(_3)/graphite</td>
<td>Provided a specific capacity of 518.5 mAhg(^{-1}) at current of 1000 mAg(^{-1}) (1000 cycles)</td>
<td>[29]</td>
</tr>
<tr>
<td>Graphene + cobalt sulfide</td>
<td>Delivered a high capacity(\geq)800 mAhg(^{-1})</td>
<td>[37]</td>
</tr>
<tr>
<td>Graphene paper</td>
<td>Discharge capacity dropping from 680 to 84 mAhg(^{-1})</td>
<td>[22]</td>
</tr>
<tr>
<td>Graphene sheet</td>
<td>Specific capacity was as high as 1264 mAhg(^{-1}) at current density of 100 mAg(^{-1})</td>
<td>[23]</td>
</tr>
<tr>
<td>SiliconC@C/graphite</td>
<td>Charge/discharge showed a specific capacity of 919.0 mAhg(^{-1})</td>
<td>[38]</td>
</tr>
<tr>
<td>Anatase TiO(_2) Quantum-Dot/graphene</td>
<td>Great improvements in electrochemical properties</td>
<td>[31]</td>
</tr>
<tr>
<td>CoO Quantum-Dot/graphene</td>
<td>Storage capacity of 1592 mAhg(^{-1}) after 50 cycles</td>
<td>[32]</td>
</tr>
<tr>
<td>Carbon nanofiber/highly branched graphene</td>
<td>Hybrid material showed a reversible capacity of 300 mAhg(^{-1})</td>
<td>[28]</td>
</tr>
<tr>
<td>Bi(_2)Te(_3)</td>
<td>Improved cyclic stability compared with Bi(_2)Te(_3) alone</td>
<td>[35]</td>
</tr>
</tbody>
</table>
### 7.3.2 Sodium Battery

Research has escalated in the exploitation of utilizing sodium as a replacement lithium ion for large-scale energy storage from the economic and ecological view-points. Sodium is cheaper when compared to lithium due to its almost unlimited availability \([40,41]\). Much effort has been dedicated in enhancing the performance of lithium batteries and to overcome some of its challenges such as the accumulation of the charge/discharge products which deteriorate the long-term performance of the battery \([40,41]\). Although much progress has been made on the latter, the possibility of using lithium for future energy generation seems to be overshadowed by its price and toxicity, making its applicability in large-scale energy storage impractical. Sodium ion batteries (SIBs), on the other hand, have a great potential to give a similar or even better performance than lithium-based batteries at a relatively low cost. Although sodium battery seems to be a suitable solution to replace lithium especially considering its low toxicity, the sodium-based batteries still experiencing similar challenges as for lithium. Some of these challenges include low efficiency and poor cycle life, and mostly overheating. Similar to lithium-based batteries, sodium batteries have the same energy storage mechanism in which charging involves the metal ions flows from the cathode (sodium-containing species) to the anode (carbon) through the electrolyte. The discharging process involves the reverse mechanism. Amongst all challenges faced by the sodium-based batteries (SIB), the formation of dendrite (which is predicted to more than lithium based batteries LIB), large volume change and sluggish diffusion kinetics have been the subject of research over the past years \([40,41]\). Different nanoscale materials with various shapes have been studied as the possible solution to overcome these challenges. In this case, the electrodes from various materials in the nanoscale were utilized in order to improve the performance of the battery \([40,41]\).

It is of interest to note that the similarities of sodium and lithium could give the scientists an edge regarding many studies conducted on lithium; however due to their difference especially

<table>
<thead>
<tr>
<th>nanoplatelet/graphene</th>
<th>[34]</th>
</tr>
</thead>
<tbody>
<tr>
<td>NiCO(_2)O(_4)/graphene composites showed improved capacity and better</td>
<td></td>
</tr>
<tr>
<td>MoS(_2)-coated vertical graphene nanosheet</td>
<td>[39]</td>
</tr>
<tr>
<td>Compared with MoS(_2)-carbon black, MoS(_2)/VGNS developed on Ni foam showed enhanced electrochemical properties</td>
<td></td>
</tr>
</tbody>
</table>
their size (viz. 1.02 Å for sodium and 0.76 Å for lithium) could cause such discrepancy with regard to the materials used for lithium batteries. For example, the use graphite as anode material for sodium yields extremely low capacity, while remarkable results were reported in the case of lithium-based batteries when graphite is an electrode. In addition, sodium has high reactivity with low melting temperature and has the similar problems as lithium when it comes to formation of dendrite. A wide variety of nanomaterials have been tested to enhance the performance of the SIBs. It is of significant importance that the material used has to bear some valuable properties such as large energy for ion adsorption, high thermal stability, high electronic conductivity, high ion diffusion and good structural stability [42,43].

7.3.2.1 Carbon-based materials

Carbon-based materials received lot of interest as the electrodes for sodium battery with regard to their performance in lithium-based batteries. Despite that, their unique structural properties and abundant availability makes them very interesting materials to enhance the performance of the batteries for large-scale energy storage. For example, Liu et al. [44] developed a binder-free multilayer nanocomposites core-shell microfiber electrode consist of hydrothermal synthesized SnO$_2$ nanocrystal layer on conductive carbon cloth (SnO$_2$/CC) by surface coating. Further coating with ALD Al$_2$O$_3$ (Al$_2$O$_3$/SnO$_2$/CC) was carried out to prevent detachment of the active material. It was reported that SnO$_2$/CC exhibited charge capacity of 529 mAh g$^{-1}$ and retained only 30% of initial capacity after 40 cycles, while Al$_2$O$_3$/SnO$_2$/CC exhibited charge capacity of 470 mAh g$^{-1}$ and maintained a 371 mAh g$^{-1}$ specific capacity at 100$^{th}$ cycle.

Graphene is one of the interesting materials which can be utilized to grow different nanomaterials, especially for high performance electrocatalytic or electrochemical devices [40,45,46]. The unique features of graphene such as explained elsewhere in this document (viz. lightweight, high surface are and conductivity) add more value to the formulation in which the nanoparticles are grown). Yang et al. [46] synthesized graphene oxide using modified Hummers method. The reduction of graphene oxide (GO) was performed by ammonia and hydrazine. GO was mixed with SnSe in a weight ratio of 9:1 using ball milling. It was demonstrated that GO/SnSe composites exhibited enhanced reversible capacity of 590 mAh g$^{-1}$ at a current density of 0.05 Ag$^{-1}$ and rate performance of 260 mAh g$^{-1}$ at a current density of 10 Ag$^{-1}$. The cycle stability with capacity retention of 98% over 120 cycles at a current density of 1 Ag$^{-1}$ was achieved. Phosphorus features extraordinary capacity of 2596
mAh g\(^{-1}\) (i.e. 7 times higher than the commercial graphite); however its low conductivity and rapid structural degradation limit its application as electrode [47]. Recent study by Zhang et al. [47] reported that amorphous phosphorus/nitrogen-doped graphene exhibited remarkable stable cyclic performance and excellent rate capability. The electrode exhibited excellent capacity retention greater than 85% over 350 cycles, remarkable cycle stability (0.002% decay per cycle from 2\(^{nd}\) to 350\(^{th}\) cycle with rate capability of 809 mAh g\(^{-1}\) at a current density of 1500 Ag\(^{-1}\). Other possibility to improve the performance of graphene-based electrodes includes fabricating nanocomposites composed of graphene and transition metal dichalcogenides [48, 49]. This is as a result of improved electrical conductivity and structural stability of the composite electrode. A 2D SnSe\(_2\)/rGO nanocomposite prepared by hydrothermal method was reported by Zhang et al. [49]. A stable SnSe\(_2\) layered structure was prepared from NaHSe solution for the first time. Nanocomposites (SnSe\(_2\)/rGO) exhibited initial discharge capacity of 756 mAh g\(^{-1}\) and a charge capacity of 660 mAh g\(^{-1}\); and after 100 cycles a reversible capacity of 515 mAh g\(^{-1}\) was obtained. On other hand, Xiong et al. [48] used sulphur doped graphene and nanostructured Sb\(_2\)S\(_3\) as an electrode for SIB. This composite exhibited a high specific capacity of 792.8 mAh g\(^{-1}\) at a current density of 0.05 A g\(^{-1}\), good rate capability (591.6 mAh g\(^{-1}\) at 5 A g\(^{-1}\)) and excellent cycle life (83% capacity retention for 900 cycles at 2 A g\(^{-1}\)). It can be concluded that the inclusion of other nanostructured materials into the graphene layers can enhance the performance of the electrodes for sodium batteries. This in turn shows the potential as sodium batteries as another form for future large scale energy storage.

### 7.3.2.2 Non-carbon materials

Non-carbon materials include layered materials with excellent capacity which can potentially replace graphite [50-52]. These materials include transition metals with chalcogen such as S, Se and Te. Their chemical formula is usually expressed as \(MX_2\) where \(M\) represents metal and \(X\) is chalcogen. The layers are bonded together by weak van der Waals forces which can accommodate ions and can further be exfoliated to accommodate more metal ions. Very few studies were conducted based on these materials as electrodes for sodium battery [50,51]. Mortazavi et al. [51] pointed out that Na ions were able to intercalate in MoS\(_2\) which provided adsorption sites, however resulted in phase transformation from hexagonal phase to tetragonal phase giving rise to a maximum theoretical capacity of 146 and a low average electrode potential in the range of 0.75-1.25 V. Through barrier energy calculations of 0.68
and 0.28 eV for Na diffusion in 2H- and 1T-MoS$_2$ predicted moderately fast charge and fast discharge rates, respectively. It was concluded that 1T polytype deliver better electrochemical performance because of its inherited metallic electronic structure, higher capacity for Na ions and their faster mobilities. Lan et al. [50] studied molybdenum dinitride (MoN$_2$) as a potential candidate for Na-ion batteries anode. MoN$_2$ in T-type structure was studied using first principle method. It was found that sodium ions can adsorb onto the monolayer with maximum capacity of 864 mAh g$^{-1}$ and diffusion barrier was 0.12 eV. Sodium ions were also found to adsorb separately without forming clusters which can contribute the overall performance of the battery especially during cycling. These materials (i.e. transition metal dichalcogenides) show potential for their application in stationary energy storage since graphite is not amenable for Na ions intercalation.

7.3.2.2.1 Borophene

Borophene is a metallic and lightweight material having two-dimensional (2D) structured sheets of boron [42,43,53]. It has a similar structure as grapheme hence the name borophene. It is capable of adsorbing sodium ions due to the availability of the adsorption site with strong binding energy. Mortazavi et al. [42] studied the capability of borophene as an anode material for Na storage. It was reported that the binding energy of 2.92 eV for Na was achieved which decreased with increase in Na content. Using Bader charge analysis to evaluate the charge transfer, it was pointed out that borophene with 2D structure have remarkable capacity of about 1380 mAh g$^{-1}$ for Na which shows the capability of using borophene for rechargeable ion batteries. In addition, the thermal stability of the borophene up to 750 K also shows the capability of this material as anode material. Similarly, Liang et al. [53] studied the possibility of borophene as anode from two different structures, namely a closed-packed triangular B layer (B$_\Delta$) and hexagonal voids or vacancies in B layer (B$_\square$). Using the first principle calculations to evaluate the potential of borophene as anode material for sodium ion, they found out that borophene has ad adsorption energy of -1.448 eV for all boron in the plane (B$_\square$) and -1.731 eV for B$_\Delta$. The fully sodiated phase for respective B$_\square$ and B$_\Delta$ was Na0.083B and Na0.1B corresponding to a theoretical specific capacity of 248 mAh g$^{-1}$ (496 mAh g$^{-1}$ for two sides) for B$_\Delta$ and 298 mAh g$^{-1}$ (596 mAh g$^{-1}$ for two sides) for B$_\square$.

7.3.3 Sodium-air battery (SAB)
The operation process of air electrode relies on the reaction of oxygen either alkali metal to yield alkali oxides [41,54,55]. The cathode is usually made up of porous carbon and/or porous metal to deliver current for oxygen reduction and also act as the host for the product. The discharge process involves the filling the voids with the oxide products and stops when all the voids are all filled. Oxygen reduction and oxidation processes rely on the catalyst coated onto the porous matrix. Thus it is of significant importance for the reaction between the oxygen and the alkali ions and electrons from conductive matrix for oxygen reduction at the catalyst sites during discharging. The reactions for sodium and oxygen can be expressed as follows: [54,56,57]

\[
\begin{align*}
Na^+ + O_2 + e^- & \rightarrow NaO_2 \quad E = 2.263 \text{ V (7.4)} \\
2Na^+ + O_2 + 2e^- & \rightarrow Na_2O_2 \quad E = 2.330 \text{ V (7.5)} \\
4Na^+ + O_2 + 4e^- & \rightarrow Na_2O \quad E = 1.946 \text{ V (7.6)}
\end{align*}
\]

The products such as NaO₂, Na₂O₂ and Na₂O were found to accumulate on the cathode which affects the electrochemical cyclability performance of the battery. Therefore, it of significant important for the electrode to have proper porous structure with appropriate pore volume and pore size for adequate oxygen diffusion. Porous structured materials with large surface area were studied as suitable solution to overcome the accumulation of electrochemical products building up on the surface of the electrode as summarized in Table 3. Carbon-based materials have been mostly studied due to their unique properties such excellent electrical conductivity, high surface area and controllable structure and porosity.

### 7.3.3.1 Carbon-based materials

Although the research based on the sodium air battery is still in an infant stage the abundant availability of sodium makes it a potential candidate for large scale application. However, the discharge products (viz. sodium superoxide (NaO₂), Na₂O₂H₂O, Na₂O₂, NaOH and sodium carbonate (Na₂CO₃) have tendency of accumulating on the surface of the cathode electrode during charging/discharging cycle which affect the performance of the battery (Table 7.3) [58-60]. This has called upon new materials having porous structure to accommodate the discharged products without affecting the performance of the battery especially the blockage of the oxygen by the discharged products [61]. Morphological properties such as surface area, pore volume and size are of significant importance to improve the cycling performance by
reducing the over accumulation of the discharge products [58]. Various carbon based materials such as graphene, MWCNT, CNT [54,62] have been considered as promising cathodes for sodium air batteries. For example, Sun et al. [54] demonstrated that diamond-like carbon thin films can be used as electrode at room temperature. Specific discharge capacity of 1884 mAh g\(^{-1}\) (565 µAh cm\(^{-2}\)) at 1/10 C and 3600 mAh g\(^{-1}\) (108 mAh cm\(^{-2}\)) at 1/60 C was achieved using carbonate-based electrolyte. Graphene nanosheets (GNS) received much interest due to their unique valuable properties such as large specific area, high electronic conductivity and high electro-catalytic activity towards the oxygen reduction reaction (OER). GNS were employed as an oxygen electrode for nonaqueous SAB by Liu et al. [63] and showed high discharge capacity of 9268 mAh g\(^{-1}\) at 200 mA g\(^{-1}\). It was demonstrated that the electrochemical properties GNS for SABs was higher than that of the carbon electrode with high catalytic activity for both oxygen evolution reaction (OER) and oxygen reduction reaction (ORR). Graphene nanosheets synthesized by the oxidation of graphite powder were functionalized with platinum nanoparticles [61]. The platinum (Pt) nanoparticles (NPs) were grown by \textit{in situ} on top of the nanosheets via hydrothermal method. It was found that the discharge capacity of platinum-based graphene reached 7574 mAh g\(^{-1}\) as compared to pure graphene with only 5413 mAh g\(^{-1}\) at the current density of 0.1 mA cm\(^{-2}\). This was attributed to presence of the platinum nanoparticles (i) acting as oxygen reservoir because of the presence of large number of effective sites to adsorb oxygen, (ii) well-distributed Pt NPs accommodated discharged products, which alleviated the blockage of oxygen diffusion channels, and (iii) the high conductivity of the Pt NPs favours the conductivity of the electrode despite the insulating discharged products on its surface.

With its unique features such as large-surface area, high chemical stability, and excellent electrical conductivity, as well as good mechanical properties carbon nanotubes merit special interest as one of promising electrode for sodium air batteries. Jian et al. [64] prepared carbon nanotube paper as binder free air electrode. The paper displayed large discharge capacity of 7530 mAh g\(^{-1}\) at a current density of 500 mA g\(^{-1}\) with 0.5 NaSO\(_3\)CF\(_3\)/DEGDME as electrolyte. A rechargeable sodium-air battery operating at room temperature based on the carbon paper coated with multiwalled carbon nanotubes (MWCNTs) as oxygen electrode was studied recently by Elia, Hasa and Hassoun [65]. It was found that cell provided a discharge capacity of 4260 mAh g\(^{-1}\) leading to a theoretical energy density higher than 8000 Wh kg\(^{-1}\) due to the interconnected structure of MWCNTs suitable for discharge products deposition. The robust network aligned nitrogen-doped CNTs were able to accommodate the discharged products allowing their uniform coverage [66]. This promoted high cycling stability for at least 10 cycles at the current density ranging from 200 to 640 mA g\(^{-1}\), and also sustained high
capacity at 600 and 640 mA g\(^{-1}\) for 44 and 55 cycles before dropping to 100 mA h\(^{-1}\). A binder-free three-dimensional (3D) nanostructured electrode composed of vertically grown nitrogen-doped carbon nanotubes on carbon paper (NCNT-CP) was developed using a spray pyrolysis chemical vapour deposition by Yadegari et al. [62]. CP was covered by randomly oriented fibres with diameter of around 10 µm exhibiting a 3D structure with diverse pore sizes (Figure 7.11a). After nitrogen-doped CNT were grown on carbon paper (CP) a uniform layer of CNTs with the length around 10µm covered the carbon fibres (Figure 7.11b-d). TEM images revealed bamboo-like structure with stacked-cone structure (i.e. characteristic of NCNTs). NCNT-CP composed of micropores and mesopores (as confirmed by BET analyses Figure 7.11g-h) which are essential for oxygen diffusion and sodium ions transportation while accommodating discharged products. The NCNT-CP electrode exhibited a specific capacitance 17 times higher than bare CP (Figure 7.12a) due to its ability to accommodate discharge products while allowing continuous oxygen and sodium ions supply onto the surface of the electrode. It was established that the specific capacity reduces by 1.7 times when the discharge current density increases (Figure 7.12b) corroborating that the electrode had an appropriate structure for air breathing. In comparison with other electrodes based on CNTs from literature, the NCNT-CP performed better as shown in Figure 7.12c.

Figure 7.11 (a-e) SEM and (f) TEM micrographs of (a) pristine CP and (b-f) NCNT-CP air electrode synthesized via SPCVD method for 3 min; (g) nitrogen adsorption/desorption isotherm of the synthesized NCNT-CP air electrode; (h) pore size distribution plot of the NCNT-CP air electrode. Reprinted with permission from [62] Copyright (2015) American Chemical Society.
Figure 7.12  
(a) Discharge curves of pristine CP and NCNT-CP air electrodes (NCNT length of 10 μm) at a current density of 0.1 mA cm\(^{-2}\). (b) Discharge curves of NCNT-CP air electrodes obtained at different current densities of 0.1, 0.2, and 0.5 mA cm\(^{-2}\). (c) The specific capacities are calculated based on the geometric surface area (mAh cm\(^{-2}\)) and also based on the total weights of CP and NCNT-CP electrodes (mAh g\(^{-1}\)); the specific capacity for NCNT-CP and other CNTs containing air electrodes with different architectures including powder-based, CNTs paper, and vertically aligned CNTs from the literature. Reprinted with permission from [62] Copyright (2015) American Chemical Society.
Table 7.3  Recent reports on the electrode for Na-air batteries

<table>
<thead>
<tr>
<th>Electrode</th>
<th>Electrolyte</th>
<th>Air</th>
<th>Specific capacity</th>
<th>Discharge product</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ordered Mesoporous carbon (OMC)</td>
<td>0.5M NaSO₃CF₃/PC</td>
<td>oxygen</td>
<td>500 mAhg⁻¹ (100 mA g⁻¹)</td>
<td>Na₂CO₃</td>
<td>[58]</td>
</tr>
<tr>
<td>Graphene nanosheets (GNS)</td>
<td>0.25M NaPF₆/DME</td>
<td>Dried air</td>
<td>9268 mAhg⁻¹ (200 mA g⁻³)</td>
<td>Na₂O₂</td>
<td>[63]</td>
</tr>
<tr>
<td>MWCNT</td>
<td>0.5M NaSO₃CF₃/TEG-DME</td>
<td>Oxygen</td>
<td>4260 mAhg⁻¹ (100 mA g⁻¹)</td>
<td>NaO₂</td>
<td>[65]</td>
</tr>
<tr>
<td>GNS-Pt</td>
<td></td>
<td></td>
<td>7574 mAhg⁻¹ (0.1 mA cm⁻²)</td>
<td>Na₂CO₃</td>
<td>[61]</td>
</tr>
<tr>
<td>CNT paper</td>
<td>0.5M NaSO₃CF₃/DEGDME</td>
<td></td>
<td></td>
<td>Na₂O 2H₂O</td>
<td>[64]</td>
</tr>
<tr>
<td>Nitrogen-doped CNTs (NCNT)</td>
<td>0.5M NaSO₃CF₃/DEGDME</td>
<td>Dried air</td>
<td>1887 mAhg⁻¹ (25 mA g⁻¹)</td>
<td>Na₂O₂ and NaO₂</td>
<td>[66]</td>
</tr>
<tr>
<td>Nitrogen-doped carbon nanotubes/carbon paper (NCNT-CP)</td>
<td>0.5M NaSO₃CF₃/DEGDME</td>
<td>Oxygen</td>
<td>-</td>
<td>Na₂O₂ and NaO₂</td>
<td>[62]</td>
</tr>
</tbody>
</table>
Supercapacitors

Supercapacitors have low energy density as compared to other energy storage devices (viz. batteries and fuel cells), however they can be used as part of these energy storage materials. It is recognized that the improvements regarding their energy density can revolutionize the future of energy storage devices [67, 68]. In supercapacitors energy is stored in two different mechanisms, i.e. capacitive and pseudocapacitive. The capacitive process (non-Faradaic) relies on the charge separation at the electrode and electrode interface, while pseudocapacitive process (Faradaic) depend on redox reactions that occur on the electrode materials [69]. A supercapacitor cell consists of two electrodes separated by an ionic conductor between them as schematically presented in Figure 7.13. The separator (ionic conductor) has to be permeable for ions to allow the charge transfer while being highly electrical resistance. The electrodes are thin coatings applied to metallic collector (e.g. aluminium). The electrodes can be the same in the case of symmetric cells, while for asymmetric electrodes are different. All these components play a major role on the performance of the supercapacitor, and the most decisive primary parameters are electrodes and electrolytes. For example, the electrolyte conductivity plays major role on the equivalent series resistance (ESR). On the other hand, there has been lot of interest in the developments of advanced electrode to improve the overall performance of the capacitors. A wide variety of nanosized materials, and/or their hybrids were studied as potential candidates to replace the graphitic commercial electrode for advance capacitors with enhanced electrochemical performance as summarized in Table 7.4. The main aim was to reduce the size and increase the contact between the active material and the current collector. Nanomaterials having large surface area can therefore increase contact with current collector to reduce the resistance. In this section we will discuss the application of nanosized materials as either electrode or part of the electrode (Figure 7.13).
7.3.4.1 Carbon materials as electrode materials

Carbon materials with their special features such as high conductivity, relatively low cost and an industrial production processes which makes them easily accessible were most studied materials as electrode for supercapacitors. Due to their large surface area activated carbon has been widely used as one of the potential electrode [67], however other carbon materials such as graphene nanosheets and carbon nanotubes has been also studied in order to enhance the performance of the capacitors [68].

7.3.4.1.1 Graphene

Graphene has received much interest due to their unique properties, namely large surface area, good flexibility, good electrical conductivity, good thermal and chemical stability for various applications especially for large-scale energy storage. Graphene oxide (GO) produced from natural flake graphite with modified Hummers’ method which yield GO sheets with thickness around 1 nm was also studied in [70]. These sheets were used as substrate to grow polyaniline (PANI) nanowires arrays as supercapacitor electrode. Reaching a specific capacitance of 555 F g\(^{-1}\) at a discharge current density of 0.2 A g\(^{-1}\) confirmed the synergistic effect between PANI.
nanowires and GO as potential electrode for supercapacitors. Conductive polymers have a tendency of shrinking and swelling during charging/discharging cycling which affects the performance of the supercapacitors which limit their commercialization [67,70]. It was found that the incorporation of the GO improved the stability of the electrode for more than 2000 cycles with capacitance retention of 92% [70]. Due to the performance brought by graphene and polyaniline (PANI) opens door for their application in flexible electrode with regard to the advancements in technology. These flexible electrodes have a potential as novel energy storage for flexible portable electronic devices, namely roll-up display, electronic paper, stretchable integrated circuits, and wearable systems for personal multimedia, computing, or medical devices. A graphene/polyaniline composite paper prepared by in situ anodic electro-polymerization of aniline monomers into polyaniline film on graphene paper to produce a highly flexible, highly conductive and electrochemical activity free-standing composite electrode as shown in Figure 7.14 was reported in [71]. It was found that the material exhibited the gravimetric capacitance of 233 F g⁻¹ (~135 F cm⁻³) showing their potential as electrode for capacitors. Most recent study based on reduced graphene oxide/polypyrrole/cellulose hybrid papers also justified the possibility of using conductive polymers to induce the flexibility and stability of the supercapacitor electrodes [72]. Introducing natural material, which is cellulose in this case, to serve as supporting frame for RGO-PPy nano-architecture clearly indicate the possibility of introducing third cheaper material to enhance the performance of the electrode, especially mechanical strength to avoid the mechanical failure during service. It was reported that the resulting hybrid was highly flexible and showed high electrochemical activity when tested in a three electrode configuration—(i.e. areal capacitance of 1.20 F cm⁻² at 2 mA cm⁻²). Moreover, excellent cycling stability with capacitance retention of ~90% after 5000 times was obtained.
Yet doping of graphene with electron donors and acceptors can be applied to enhance its performance as electrode [73-75]. GO obtained by Hummer’s method followed by heating in microwave reactor was doped with nitrogen (18 wt%) resulting in a maximum specific capacitance of 461 F g⁻¹ [74]. A facile hydrothermal method with the assistance of amphoteric salt can also be utilized to introduce nitrogen on graphene sheets. However, in this case the specific capacity only reached 242 F g⁻¹ at the current density of 1 A g⁻¹ with cyclability retention of 97.6% after 5000 cycles [73]. Much higher capacity retention (99.7%) was recently reported by Zhu et al. [75] using nitrogen-doped graphene decorated with iron nanorings (Fe₃)₄NRs/NH₂-RGO) after 1000 cycles.

One of the major problems that affect the performance of the graphene-based electrodes is restacking of the sheets which reduce the effective surface area. The introduction of the polymers such polypyrrole can intercalate between the sheets and keep them apart which can improve the

Figure 7.14  (top) Illustrative fabrication process and (bottom) digital camera images of (left) two freestanding G-papers (30 mm × 10 mm) and right a flexible G-paper. Reprinted with permission from [71]. Copyright (2009) American Chemical Society.
electrochemical activity of these graphene-based electrodes. In [76] electrochemical polymerization of pyrrole (PPy) on partially exfoliated graphene sheets on top of graphite foil electrode (Ex-GF) was carried out in the presence of sulfonated aromatic compounds (viz. 1,5-naphthalene disulfonate (NDS) and 2-naphthalene sulfonate (NMS) for ‘permanent’ doping anions to afford dual doped PPy which anchors partial exfoliated graphene to construct Ex-GF/PPy-NDS and Ex-GF/PPy-NMS electrode. Ex-GF/PPy-NDS and Ex-GF/PPy-NMS displayed specific conductance of 351 and 322 F g\(^{-1}\), respectively. This was ascribed to the open structure of Ex-GF/PPy-NDS and synergistic effects between the components.

Another suitable solution to overcome restacking of the graphene sheets relies on the utilization of metal oxides as spacers [77-79]. Recent study conducted on manganese dioxide nanorods intercalated between graphene oxide (rGO-MnO\(_2\)) justified the possibility of using such hybrid materials can enhance the overall electrochemical properties [77]. The nanocomposite hybrid displayed superior specific conductance of 468.48 F g\(^{-1}\) when compared to the pure manganese (358.23 Fg\(^{-1}\)) at the same current density. This hybrid nanocomposite had a retention capacity of 78.4% as compared to 46.22% for pure MnO\(_2\). Despite that high capacity retention (93.5% after 10000 cycles) were also reported by Zhong et al. [79] using 3D graphene oxide/MnO\(_2\) nanocomposites, however they managed to reach a maximum capacity of 278.5 F g\(^{-1}\) at current density 1 A g\(^{-1}\). On the other hand, inclusion of silver (Ag) into MnO\(_2\) and GO (Ag/MnO2/rGO) resulted in a specific capacitance of 467.5 F g\(^{-1}\) at a scan rate of 5 mV s\(^{-1}\) [78]. This hybrid did not show any degeneration after 1000 cycles at the scan rate of 80 mVs\(^{-1}\).

Hybrid composites based on graphene was also studied as an alternative to enhance the electrochemical properties [80-82]. Graphene nanosheets-tungsten composite as an alternative electrode for supercapacitors was reported in [80]. It was reported that the composite material performed better than pure tungsten-based electrode (~32.4 F g\(^{-1}\)) by reaching 143.6 F g\(^{-1}\) (343%) at a current density of 0.1 A g\(^{-1}\). A chemical deposition of gold nanoparticles into the pores of three-dimensional nitrogen-doped graphene nanosheets was recently reported in [82]. A maximum specific capacitance of 168.6 F g\(^{-1}\) at a current density of 1 A g\(^{-1}\) with retention of ~97% of the initial capacitance after 500 cycles was achieved. This was attributed to the synergistic effect of gold nanoparticles and hierarchically porous nitrogen-doped graphene. Fan et al. [81] also studied the sandwich composite comprise of carbon nanotubes (CNT)/graphene nanosheets. Three-dimensional (3D) CNT/graphene sandwich (CGS) structures with CNT as
pillars grown in between the graphene layers by chemical vapor deposition (CVD) method. GCS was applied as an electrode in supercapacitor and a maximum capacitance of 385 F g\(^{-1}\) was obtained at a scan rate of 10 mV s\(^{-1}\) in 6 M KOH aqueous solution. This was attributed to the presence of CNT in between the graphene sheets which act as ‘spacer’ to allow the diffusion of electrolyte ions resulting in improvement of the electrochemical properties. Furthermore, the CNT forms an interconnection with graphene providing conductive network within the sheets reducing the resistance of the electrode.

### 7.3.4.1.2 Porous carbon materials

Activated carbons have been the most studied carbon material [67,68]. This is as a result of the cost related to the activated and their large specific area. These materials are usually obtained from carbon-rich natural sources by heat treatment in an inert atmosphere (carbonization) and chemical treatment. The chemical process involves oxidation of the carbon-rich material in hot alkali or acidic medium. As a result, the carbon material become more porous with large surface area ranging between 1000-2000 m\(^2\) g\(^{-1}\). One of the limitations of these materials is control over the pore size distribution which hurdles their fully exploitation. It is noteworthy mentioning that most of the commercially available devices include carbon active based electrodes. Very recent study by Peng et al. [83] showed that nitrogen-doping of microporous carbon from natural plant (pomelo mesocarps) can enhance the electrochemical properties. Pomelo mesocarps was used to produce porous carbon nanosheets by heat treatment at 800\(^\circ\)C under nitrogen flow. It was reported that nitrogen-doping resulted in high specific conductance (~245 F g\(^{-1}\) at 0.5 A g\(^{-1}\)) and stability in Na\(_2\)SO\(_4\) for 10000 cycles and retained 96.5% maximum capacitance. Clearly, natural materials can be applied as a source of carbon to produce low cost supercapacitors electrodes or even in other energy storage applications [83-85]. Mesoporous carbon from lignin was activated by potassium hydroxide (KOH) and carbon dioxide (CO\(_2\)) as supercapacitor electrode [84]. It was reported that the CO\(_2\) and KOH activated carbon had large surface area (~1148 m\(^2\) g\(^{-1}\) and pore volume of 1.0 cm\(^3\)g\(^{-1}\)) and displayed the specific capacitance of 102.3 F g\(^{-1}\) and 91.7 F g\(^{-1}\), respectively. Rice-husks and beet sugar to produce micro- and mesoporous activated carbon was studied by Kumagai et al. [86]. The specific conductance reached ~116 and 106 F g\(^{-1}\) at scanning rate of 5 mV s\(^{-1}\) for activated carbon from both rice husks and beet as well as rice husk,
respectively. Nitrogen-containing activated carbon was reported in [87] using silk fibroins as the carbon source. Application of adequate heat treatment turned the nitrogen atoms from peptides bonds to the silk carbons in the form of nitrogen-containing functional groups. A maximum specific capacitance of 52 F g\(^{-1}\) was achieved.

Electrospinning is one of the techniques that afford to produce fibres with diameters ranging from several micrometers to few nanometers. It was recently recognized that electrospinning can be utilized to produce the mats with nanoporous structure with large surface area as self-standing supercapacitor electrode. This can be achieved by electrospinning suitable polymers followed by carbonization by heating in a furnace [88]. In [88] a self-supported and binder-free microporous polyimide (PI)-based carbon fibers were fabricated by electrospinning from a blend of polyvinyl (PVP) and poly acryl amide (PAA). A maximum specific capacitance of 215 F g\(^{-1}\) based on symmetrical two electrode supercapacitor at a current density of 0.2 A g\(^{-1}\) was achieved.

7.3.4.1.3 Carbon nanotubes

CNT with the unique properties such as large surface area, excellent electronic, mechanical and chemical properties open doors for a wide variety of applications. These materials are often produced by catalytic decomposition of hydrocarbons. Depending on the fabrication process and the conditions applied two carbon nanotubes can be obtained, namely multi-walled nanotubes (MWCNTs) and single-walled nanotubes (SWCNTs). The freestanding CNT were produced by chemical deposition vapor (CVD) as an electrode for electrochemical double layer capacitors (EDLCs) [89]. Cyclic voltammogram (CV) was almost rectangular-shaped indicating good capacitive behaviour with relatively low equivalent series resistance (ESR). The specific conductance from the galavanosatic charge-discharge curve was 51 F g\(^{-1}\). Similar study based on multi-walled CNT achieved a maximum capacitance of 20 F g\(^{-1}\) [90]. Interestingly, the specific power density reached a value of 30 kW kg\(^{-1}\). Moreover, the electrode displayed fairly ideal voltammograms even at high scan rate of 1000 mVs\(^{-1}\) which corroborate the applicability of freestanding CNTs as potential electrodes for supercapacitors.

The combination of CNT with metal oxides was also studied to enhance the electrochemical properties [91-93]. A combination of CNT and MnO\(_2\) was studied by Chou et al. [91] by electrodepositing the MnO\(_2\) nanowires onto CNT paper using a cyclic voltammetric technique.
The specific capacitance of the CNT/MnO\(_2\) composite (~129 F g\(^{-1}\)) was superior to pure CNT (32 F g\(^{-1}\)) at the scan rate of 5 mVs\(^{-1}\). In this case CNT acted as a good conductive layer allowing the MnO\(_2\) fibres to facilitate the movement of the electrolytes active materials. The material displayed a good cyclability with 88% retention of initial capacitance after 3000 cycles. This opens doors for the composite based on CNT and metal oxides in flexible materials as compared to non-flexible powdered based composites materials. A porous flexible capacitor based on CNT/MnO\(_2\) was recently reported by Du et al. [94]. The authors used the vacuum filtering method to produce a hybrid of CNT/MnO\(_2\) nanotubes (NT) which exhibited volumetric capacitance of 5.1 F cm\(^{-3}\) and a high energy density of 0.45 mWh cm\(^{-3}\). Their results clearly corroborate the application of these flexible materials for high performance power supplies in wearable electronic devices. Yet another method to produce flexible energy storage materials include the use of conductive polymers/CNT composites as compared to the powder-based rigid composites [95].

### 7.3.4.2 Metal oxides

Metal features unique valuable properties such as high specific capacitance and conductivity making them suitable candidates for electrode fabrication of high density energy supercapacitors. Various metal oxides such as RuO\(_2\), IrO\(_2\), MnO\(_2\), NiO, CoO\(_2\), SnO\(_2\), V\(_2\)O\(_5\) or MnO\(_x\) have been explored for electrode fabrication [96-101]. Amongst all, ruthenium and manganese oxides are the most studied, thus are the only ones discussed in the following subsections. The mechanisms behind the energy storage in mainly from the fast faradic redox reactions that occur between the oxides and the electrolyte.

#### 7.3.4.2.1 Manganese oxides (MnO\(_2\))

MnO\(_2\) is considered as one of the most promising materials for fabrication of electrodes in capacitors at fairly low cost; and it is environmentally friendly with high theoretical capacitance value (~1110 F g\(^{-1}\)). The charge storage of manganese oxides relies on pseudocapacitive reactions on the surface and in the bulk of the electrode. The reaction involves a reversible redox transitions which include the exchange of protons and/or cations with the electrolyte and
transitions between the Mn(III)/Mn(II), Mn(IV)/Mn(III) and Mn(IV)/Mn(IV). This can be expressed as the following \textbf{Equation (7.7)}: \cite{102, 103}

\[ MnO_2 + C^+ + e^- \leftrightarrow MnOOC \]  

(7.7)

where \( C^+ \) are metal cations/protons from the electrolyte (Li\(^+\), Na\(^+\), K\(^+\)). Since the physical and chemical structure of MnO\(_2\) affect pseudocapacitive much work has been done to improve the overall performance. This is as a result of limited specific capacitance of MnO\(_2\) (100-200 F g\(^{-1}\)), lack of structural stability and long-term cyclability. It was recognized that the cyclability is mainly defined by the microstructure, while the specific capacitance relies on chemical hydrated state. Much efforts have been dedicated to the synthesis of the MnO\(_2\) with desirable morphology, chemistry and crystal structure. For example, for highly crystalline MnO\(_2\) the ion exchange is limited due to loss of surface area, but there is an increase in conductivity. In turn, there is tradeoff between these two parameters (ion exchange and conductivity) which requires special attention to control the synthesis conditions \cite{103}. The effect of the crystal structure on the specific capacitance is well discussed in the review paper published by Wei \textit{et al.} \cite{104}. This well documented review clearly discussed on how the preparation processes which changes the crystal structure of MnO\(_2\) which in turn affect the specific capacitance of the resulting material. The most suitable solution to overcome the shortcomings of MnO\(_2\) is to include the substrate which is highly conductive and highly porous \cite{104}. Various materials such as graphene oxide \cite{79}, carbon nanotube \cite{102} and electrospun carbon nanofibers \cite{105} have been investigated as potential substrate for MnO\(_2\). Moreover, additional nanoparticles were incorporated to enhance the electrochemical performance of MnO\(_2\)-based electrodes \cite{78}. For instance, Ma \textit{et al.} \cite{78} decorated MnO\(_2\)/graphene oxide with silver nanoparticles to produce a ternary nanocomposite system. In this case, the MnO\(_2\) nanoparticles were grown on graphene oxide followed by co-reduction of Ag\(^+\) and graphene oxide (GO). It was reported that the \textit{in situ} formed Ag and MnO\(_2\) nanoparticles were well-distributed on the surface of reduced GO giving rise a maximum capacitance of 467.5 F g\(^{-1}\) at a scan rate of 5 mV s\(^{-1}\) which was superior to that of MnO\(_2\)/rGO (293.2 F g\(^{-1}\)). Moreover, the nanocomposites did not show any degeneration after 1000 cycles at a scan rate of 80 mV s\(^{-1}\). The inclusion of other nanoparticles improves the electrical conductivity which enhances the electron transfer of active materials. It can be concluded that
MnO$_2$ have a potential of being a part of the next generation of energy storage materials with regard to its performance especially when combined with other conductive materials. For example, very recent study based on MnO$_2$ nanoflake/CNT core-shell particles as electrode for supercapacitors in [102] justified the enhancement of electrochemical performance of the hybrid nanocomposites. The MnO$_2$ coating on CNT particles was obtained by the reduction of KMnO$_4$ in the presence of CNT particles. This method involves two stages: (i) fast nucleation forming amorphous particles, and these particles aggregate in a crystal plane and (ii) come together to form a sheet-like structure through Ostwald ripening as shown in Figure 7.15. In comparison with pure CNTs, the MnO$_2$ coated CNT resulted in compact structure with MnO$_2$ flakes deposited on top of CNT and the contact with neighbouring MnO$_2$ was achieved by increasing the time for growth (Figure 7.15). The electrochemical performance was evaluated in an aqueous Na$_2$SO$_4$ at scanning rate of 5, 10, 20, 50 and 100 mV s$^{-1}$. A rectangular shaped cyclic voltammetry (CV) curves were obtained even at high scanning rate corroborating the good electrochemical capacitor behaviour (Figure 7.16). However, at low content of MnO$_2$ (short growth time) the weak redox was depicted as compared to the higher content (longer growth) in which the peaks were more prominent due to more MnO$_2$ covering the CNT. A complete coverage of the MnO$_2$ onto CNT at higher content led to redox reaction on the MnO$_2$ surface which involves manganese ions Mn$^{4+}$ and Mn$^{3+}$. This reaction involves the exchange of alkali cations or protons from electrolyte, which is expressed as $MnO_2 + X^+ \leftrightarrow MnOOX$, where $X^+$ is $Na^+$ or $H^+$. The maintenance of the CV curves even at high scanning rate was attributed to a strong bonding between MnO$_2$ nanoflakes on the CNT surface and low enough interfacial resistance between MnO$_2$ and CNTs. It was reported that the higher MnO$_2$ content led to high current density than the lower MnO$_2$ content as well as CNT particles. The specific conductance from galvanostatic charge-discharge indicated that the higher MnO$_2$ (370 Fg$^{-1}$) was superior to lower MnO$_2$ content (147 Fg$^{-1}$) as well as bare CNTs (26 Fg$^{-1}$) at current density of 0.5 Ag$^{-1}$ (Figure 17). A comparison between bare MnO$_2$, low content MnO$_2$/CNT and high content MnO$_2$/CNT cyclic performance up to 4000 cycles was performed to evaluate their cyclic stability. It found that the composites maintained ~100% of the initial capacitance, whereas the MnO$_2$ particle electrode revealed decay and exhibited only 7% of the initial capacitance as shown in Figure 7.17. The decay of MnO$_2$ was ascribed to the detachment of MnO$_2$ during insertion/desertion and/or the dissolution of MnO$_2$ into aqueous electrolyte solution,
corroborating the fact that the inclusion of CNT enhanced reversible charging/discharging reaction without debonding. The energy density and a power density were found to be 27 Whkg\(^{-1}\) and 225 Wkg\(^{-1}\) for the higher content of MnO\(_2\) composites.

![Diagram](Image)

Figure 7.15  (up) Schematic fabrication of MnO\(_2\)/CNT core-shell particles. (down) SEM images of (a) CNT particles, (b) low-MnO\(_2\)/CNT core-shell particles, and (c) high-MnO\(_2\)/CNT core–shell particles. High-magnitude SEM images of (d) CNT particles, (e) low-MnO\(_2\)/CNT core–shell particles, and (f) high-MnO\(_2\)/CNT core–shell particles. (g) SEM images of the internal space of the MnO\(_2\)/CNT core-shell particle. Reprinted with permission from [102]. Copyright (2009) American Chemical Society.
Figure 7.16 CV curves of (a) the CNT particles, (b) the low-MnO$_2$/CNT core–shell particles, and (c) the high-MnO$_2$/CNT core–shell particles measured at various scan rates from 5 to 100 mV$s^{-1}$ in a 1 M of Na$_2$SO$_4$ electrolyte solution. Reprinted with permission from [102]. Copyright (2009) American Chemical Society.
7.3.4.2.2 Ruthenium oxide

Among other metal oxides, ruthenium oxide is one of the most studied metal oxide. This is as a result of unique properties such as catalytic activities, metallic conductivity, electrochemical reduction-oxidation, field emitting behaviour, high thermal and chemical stability. This pseudocapacitive material has high specific capacitance of about 1000 Fg$^{-1}$ which makes it the

![Graphs showing galvanostatic charge/discharge curves and specific capacitance](image-url)
most interesting material [106]. It has a wide potential window and long life cycle. Its pseudo capacitance behaviour of RuO$_2$ relies on series of oxidation states, namely Ru(IV), Ru(III) and Ru (II). It also features reversible redox processes. The pseudocapacitive process involves the electro-adsorption of protons on the surface of RuO$_2$ electrode. Generally, it can be expressed as: [69, 106-108].

$$RuO_2 + xH + xe^- \leftrightarrow RuO_{2-x}(OH)_x$$  \hspace{1cm} (7.8)

where $0 \leq x \leq 2$. $x$ changes during insertion/de-insertion occurring at 1.2 voltage window resulting to capacitive behaviour due to ion adsorption following a Frumkin-type isotherm. Brumbach et al. [106] studied the nanostructured ruthenium oxide electrode obtained via high-temperature molecular templating for use in electrochemical capacitors. They reported unusual behaviour for thermally prepared RuO$_2$ materials with voltammograms showing that the templated-RuO$_2$ films (viz. fabricated at high temperatures in the presence of alkyl-thiols) established current 4 times greater than untemplated-RuO$_2$, which displayed resolvable redox peaks, to broad ‘squarish’ voltammograms without resolvable peaks, similar to that obtained for hydrous ruthenium oxide and mixed oxides systems. This was attributed to the access granted by the RuO$_2$ materials for the electrolyte as shown in Figure 7.18. The specific capacitance of untemplated RuO$_2$ was around 200 F g$^{-1}$, while templated RuO$_2$ reached values near 400-500 F g$^{-1}$. These clearly demonstrate that it is possible to reach high capacitance values by using templates which can also be explored for other less expensive metal oxides. The utilized methods afford highly crystalline structure (i.e. leading high conductivity) and densification of the film. SEM images of the templated-films showed a regular array of circular feature, 300 ± 50 nm in diameter, at the surface of the templated-RuO2 (Figure 7.19A-C). At lower concentrations of RuCl$_3$ in the precursor solution resulted in a higher density of the circular features as shown in Figure 7.19A&B which are more on the subsurface as pores (Figure 7.19B). All these features were not observed in the case of untemplated RuO$_2$; hence they were related to the bubble generated and gas evolved from pyrolysis of the templating material. Moreover, ill-defined pores ($\sim$ 4nm in diameter) were observed using TEM (Figure 7.19D&E). It was surmised that the chain length of hexane thiol was too short to support formation of a mesophase but influenced the growth of rutile RuO$_2$ grains and led to a hierarchically porous, nanostructures film which
improved electrolyte access, while protonic and electronic conduction are preserved through the network of hydroxylated RuO$_2$ nanograins.

Figure 7.18  Cyclic voltammetry of ruthenium oxide thin films on titanium are shown for (A) thermally prepared RuO$_2$, and RuO$_2$ formed with (B) 1:1 and (C) 1:2 molar additions of thiol relative to Ru in the precursor solution. The electrolyte solution was 1 M H$_2$SO$_4$ and potentials are referenced to Hg/HgSO$_4$. Three voltammograms are shown for each scan rate. Scan rates were 5 (black), 10 (red), 20 (blue), and 50 (green) mV/s. Reprinted with permission from [106]. Copyright (2009) American Chemical Society.
One of the major concerns is the high cost associated with the manufacturing and/or designing an electrode that can exploit the highest capacitance from the least amount of this metal oxide. Therefore, much effort has been dedicated in developing a combine structure of RuO$_2$ and other electron-conductive electro-chemically stable materials as substrates. Various substrates were studied to come up with best substrate and conducive synthesis process to coat RuO$_2$ onto a porous substrate especially carbon-based materials since they comprise most of the required properties (e.g. high conductivity and easily prepared into porous large surface area substrate). These include the likes of graphene [109-111]; carbon nanotubes [112,113]; and carbon nanofibers. It was found that it is possible to overcome the shortcomings of RuO$_2$ especially with regard to the fact that only a part of it participate in the charge-storage mechanism while the
underlying active material remain unreacted. Thus a combination of large surface area materials such as carbon materials with ruthenium oxides can lead to a hybrid nanocomposite with high specific conductance ranging between 150 to 1540 F g⁻¹ depending on the thickness of the films. Recent studies showed that the inclusion of the ruthenium into the electrode enhances the electrochemical performance of the overall nanocomposite materials [113,114]. This is attributed to its unique features such as high capacitance (>700 F g⁻¹), low electrochemical resistance (ESR) and long life cycle [113]. One of the principal reasons that hampers the progress of RuO₂ especially towards commercialization is its limited availability and high cost. Despite that the inclusion of these substrates can reduce the overall price of the electrode which facilitates its commercialization in the mere future.

Hydrous ruthenium oxides (RuO₂ₓH₂O) features high specific capacitance (~900 F g⁻¹) (theoretical 1358 F g⁻¹), and high electrical conductivity (3 × 10² S cm⁻¹) resulting from the interparticle and interlayer hydrous regions which allows protonic conduction [67,109]. This results in high power and high density supercapacitor. Ternary nanocomposites based on hydrous oxides were also studied aiming to enhance electrochemical properties of the electrode. Using a system comprise of multi-walled carbon nanotubes/RuO₂ₓH₂O/ titanium (Ti) as electrode in 1.0 M H₂SO₄ aqueous solution reaches a maximum specific capacitance of 1652 F g⁻¹ at a scan rate of 10 mV s⁻¹ [113]. Such behaviour can be related to the nanopore structure of the material which allows fast and reversible faradaic processes on the electrode surfaces. To reduce the price and restacking of the RuO₂ other cheap nanoparticles or sheets can be incorporated between the RuO₂ layers. However, there is some sort of dependency on the ratio between the incorporated particles and RuO₂ in the hybrid material. Thus optimization of the ratio can lead to a high specific conductance [115]. For instance, Deng et al. [115] found that 40wt% of RuO₂ was the maximum in the hybrid composite of RuO₂/graphene to achieve a specific conductance of 998 F g⁻¹ with excellent cycle stability, and high power density of 20.28 Wh kg⁻¹. Elsewhere, a high specific capacitance of 1017 F g⁻¹ was reported using a cheaper vapour grown carbon fibres as a matrix for the dispersion of RuO₂ₓH₂O [114]. A core-shell structure of carbon/carbon nanotube supported Ag-Ru nanoparticles can be prepared by using Ag to form a hollow structure which later is replaced by ruthenium. This kind of structures offers more enhancements regarding capacitive behaviour by promoting the proton transportation in RuO₂ due to the shortened diffusion distance [116].

Bendable/flexible capacitors based on RuO₂ were also studied to evaluate their applicability especially for wearable devices [117]. In this case, a binder-free bendable RuO₂ thin film can be
fabricated on top of flexible carbon-based materials such as graphene. Cho et al. [117] recently published an article based on RuO$_2$ fully flexible supercapacitor electrode. The authors prepared a RuO$_2$ film by coating it onto graphene-coated copper foil using cathodic electroplating technique. It was stated that the electrode exhibited a specific conductance of 1561 F g$^{-1}$ at a scan rate of 5 mV s$^{-1}$ and significantly improved initial capacitance retention of 98% under bent conditions. This electrode exhibited a high energy density of ~13 Wh kg$^{-1}$ at a power density of ~21 kW kg$^{-1}$. All these improvements were attributed to the improved mechanical adhesion between the RuO$_2$ and the current collector. There was no structural damage after 500 CV cycles.

Other suitable solution to overcome the high cost ruthenium oxides is to utilize synthesizing route that are cheaper and easily accessible. Furthermore, the route has to be user-friendlier and green. Ismail et al. [69] studied the use of Aspalathus linearis natural extract as a reducing/oxidizing agent and stabilizing agent for synthesizing RuO$_2$ nanoparticles. In this case they used RuCl$_3$.H$_2$O, although is extremely destructive to respiratory system, as precursor and mixed with Aspalathus linearis natural plant extract and the synthesis was carried out at room temperature. The RuO$_2$ nanoparticles were deposited on the nickel foam (NiF), 450 g m$^{-2}$ as 3D template. They observed Faradaic redox peaks around 0.28 and 0.38 V for the NiF/RuO$_2$ as the result of adsorption/desorption of electrolyte K$^+$ at the surface of RuO$_2$ which follows Equation 7.8. The CV curves confirmed a reversible electron-transfer process demonstrating that the capacitance was associated with charge storage of RuO$_2$ in KOH aqueous electrolyte due to rapid intercalation of alkali metal cations K$^+$ in the electrode during reduction and oxidation processes. The calculated specific capacitance for RuO$_2$/NiF reached 750, 685, 630, 600 and 480 F g$^{-1}$ at the current density of 10, 25, 50, 75, and 100 A g$^{-1}$, respectively. The cycle efficiency reached 97% retention of initial capacitance over 500 recharge/discharge cycles justify the applicability of this material for long-term. The excellent pseudocapacitive properties and high cycling stability was attributed to the high electrical conductivity of NiF and RuO$_2$ as well as 3D porous structure of the nickel foam which provided easy access to ions from the electrolyte at the electrode/electrolyte interface.
Table 7.4  Summary of recent results from selected studies of the capacitors based on nanomaterials

<table>
<thead>
<tr>
<th>Active material</th>
<th>BET surface area (m²g⁻¹)</th>
<th>Csp (Fg⁻¹)</th>
<th>Scan rate (mVs⁻¹)</th>
<th>C retention (%)</th>
<th>cycles</th>
<th>Refs.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Micro- and mesoporous activated carbon (from Rice Husks and Beet)</td>
<td>1357</td>
<td>116 ± 2</td>
<td>10.5 ± 0.2</td>
<td>-</td>
<td></td>
<td>[86]</td>
</tr>
<tr>
<td>Micro- and mesoporous activated carbon (rice husk)</td>
<td>1442</td>
<td>106 ± 6</td>
<td>7.4 ± 0.4</td>
<td>-</td>
<td></td>
<td>[86]</td>
</tr>
<tr>
<td>Fe₃O₄ NRs/NH₂-RGO</td>
<td>145</td>
<td>99.7</td>
<td>1000</td>
<td></td>
<td></td>
<td>[75]</td>
</tr>
<tr>
<td>Porous graphitic biomass carbon (PGMC)</td>
<td>1732</td>
<td>222</td>
<td>-</td>
<td>84</td>
<td>5000</td>
<td>[85]</td>
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<tr>
<td>MnO₂ nanoflakes/CNT</td>
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<td>-</td>
<td>~100</td>
<td>4000</td>
<td></td>
<td>[102]</td>
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<tr>
<td>RuO₂/graphene</td>
<td>998</td>
<td>94%</td>
<td>1000</td>
<td></td>
<td></td>
<td>[115]</td>
</tr>
<tr>
<td>RuO₂/Carbon fibre</td>
<td>13</td>
<td>824</td>
<td>1000</td>
<td>97</td>
<td>300</td>
<td>[114]</td>
</tr>
<tr>
<td>Core-shell Ag-Ru nanoparticles/Carbon/CNT</td>
<td></td>
<td>819.9</td>
<td></td>
<td></td>
<td></td>
<td>[116]</td>
</tr>
<tr>
<td>RuO₂/activated carbon</td>
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<td>180.2</td>
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<td></td>
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<td>[96]</td>
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<td>Material</td>
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<tr>
<td>Bendable RuO$_2$/Graphene/Cu</td>
<td>1494</td>
<td>5</td>
<td></td>
<td></td>
<td>[117]</td>
<td></td>
</tr>
<tr>
<td>RuO$_2$/Cu</td>
<td>1891</td>
<td>5</td>
<td></td>
<td></td>
<td>[117]</td>
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<tr>
<td>RuO$_2$/graphene/CNT foam</td>
<td>743</td>
<td>507</td>
<td>-</td>
<td>106</td>
<td>8100</td>
<td>[109]</td>
</tr>
</tbody>
</table>
7.4 Concluding remarks

The desire for a portable electronic device has pushed the technological enhancements in rechargeable solid-state batteries. It has been explained in these chapter that a battery is composed of various electrochemical cells that are linked to provide the required voltage and capacity. Generally, a cell consists of a positive and negative electrode separated by electrolyte solution containing dissociated salts. One of the most well-known rechargeable batteries is the lithium ion (LIB) and sodium-ion batteries whereby the ions move from the positive electrode to the negative electrode during charging and back during discharging. We have reviewed recent progress in electrochemical applications of lithium ion, supercapacitors and sodium ion batteries. Different nanostructured materials (viz borophene, graphene, carbon nanotubes, porous carbon materials and metal oxides) have been used as electrode material for lithium ion, supercapacitors and sodium ion batteries. Amongst the above mentioned nanostructured materials, graphene have been mostly used in the three systems discussed in this chapter either as an anode or cathode electrode. The advantages of using graphene as one of the electrode are due to its high surface area, ultra-thin thickness, excellent thermal conductivity and good chemical functionality. Furthermore, graphene structure has controllable surface chemical groups which are advantageous for variety of applications in comparison to well-known nanoparticles such carbon nanotubes and graphite. Graphene, as a negative electrode displayed an improved electrochemical performances i.e. reversible capacities up to 540 mAh g\(^{-1}\) have been achieved, which is more than the practical value of graphite (350 mAh g\(^{-1}\)). The synergistic effect of graphene with other nanoparticles enhances the electrochemical properties further due to (i) graphene nucleating; growing on the surface of other nanoparticles especially metal oxides (ii) other nanoparticles between the layers of the graphene sheet can minimize re-stacking (iii) Graphene can also act as conductive substrate to improve the electrical conductivity as a result the charge transfer of the overall system. Generally, the quality and structure of the nanoparticles depending on the method of preparation used differ from system to system which makes the device behave inconsistently. The above sentence gives a short description generally about the performance of nanostructured materials in energy storage/conversion up to date. For future purposes, attention should be paid to environmental concerns of batteries especially with the addition of nanostructured materials. It is possible that they may come in contact with humans or end up in the environment during their life cycle.
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