Nanostructured Europium and Palladium Oxide Substituted Lithium Manganese Oxide [LiEu_(x)PdO_(y)MnO₃] Perovskite Material for Li Ion Battery Cathode



By

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June, 2021

https://etd.uwc.ac.za/

DEDICATION

This work is dedicated to the memory of my loving grandparents:

John and Joyce Peter



DECLARATION BY CANDIDATE

I TUMISO EMINENCE MABOKELA declare that NANOSTRUCTURED EUROPIUM AND PALLADIUM OXIDE SUBSTITUTED LITHIUM MANGANESE OXIDE MATERIAL FOR LI ION BATTERY CATHODE APPLICATION is my own work and that all the sources that I have used or quoted have been indicated and acknowledged by means of complete references and that this work has not been submitted before for any other degree at any other institution.

Signature	Add	

Date7 JUNE 2021.....

11+1



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"Trust in the LORD with all your heart and lean not on your own understanding; in all your ways acknowledge Him, and He will make your paths straight." - Proverbs 3:6

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ABSTRACT

The 4th Industrial revolution which is to be mainly powered by cleaner energy technologies has necessitated the scientific community to develop new high-tech energy storage materials. Furthermore, 4IR is associated with increased use of handheld and portable devices which require energy carriers such as supercapacitors and batteries with high power densities and high capacities. Although layered materials such as highly lithiated manganese oxides have paved the way in the development of new high-tech energy storage materials for Li-ion batteries, there is still huge room for improvements in such materials to achieve even greater electrochemical performance. It has been reported that highly lithiated manganese oxides can be modified using a variety of methods, such as surface coating, doping, and acid treatment to improve their stability and general electrochemical performance. Thus, this research focused on integrating these modification strategies to develop electrochemical superior highly lithiated manganese oxides to be applied as aqueous based Li-ion battery cathodes. In this work, europium doped highly lithiated manganese oxide Li₂MnO₃ were synthesized through sol-gel synthesis and electrochemically evaluated. The synthesized europium doped Li₂MnO₃ were then treated with mild HNO3 as acid treatment, which has a synergistic effect on its electrochemical performance. Lastly, A jointly modified highly lithiated manganese oxide Li₂MnO₃ was fabricated by doping with europium and then decorating its surface with PdO via sonochemical methods to form a corrosion protective layer. The produced materials were characterized by Powder X-ray Diffraction (PXRD), Fourier Transform Infrared Spectroscopy (FTIR), Small Angle X-ray Scattering (SAXS), Inductively Coupled Plasma Optical Emission Spectroscopy (ICP-OES), High Resolution Scanning Electron Microscopy (HR-SEM), High Resolution Transmission Electron Microscopy (HR-TEM), and their electrochemical properties were evaluated using Cyclic Voltammetry (CV), Electrochemical Impedance Spectroscopy (EIS) and Galvanostatic Charge Discharge (GCD) in 1 M Li₂SO₄ electrolyte solution. The synthesized materials Li₂Mn_{0.95}Eu_{0.05}O₃, Acid treated Li₂Mn_{0.95}Eu_{0.05}O₃, and PdO@Li₂Mn_{0.95}Eu_{0.05}O₃ exhibited good electrochemical properties and yielded discharge capacities of 4.20, 17.8 and 6.4 mAh.g⁻¹ respectively.

RESEARCH OUTPUT

Publications

- ➤ Mabokela, T.E., Nwanya, A.C., Ndipingwi, M.M., Kaba, S., Ekwere, P., Werry, S.T., Ikpo, C., Kwena, M. and Iwuoha, E., 2021. Recent Advances on High-Capacity Li Ion-Rich Layered Manganese Oxide Cathodes. *Journal of The Electrochemical Society*. (Published 16/07/2021) See appendix 1
- ➤ Somo, T.R., **Mabokela, T.E.**, Teffu, D.M., Sekgobela, T.K., Hato, M.J. and Modibane, K.D., 2021. Review on the effect of metal oxides as surface coatings on hydrogen storage properties of porous and non-porous materials. *Chemical Papers*, pp.1-15. (Published 05/01/2021) *See appendix 2*
- Somo, T.R., Mabokela, T.E., Teffu, D.M., Sekgobela, T.K., Ramogayana, B., Hato, M.J. and Modibane, K.D., 2021. A Comparative Review of Metal Oxide Surface Coatings on Three Families of Cathode Materials for Lithium Ion Batteries. Coatings, 11(7), p.744. (Published 22/07/2021) See appendix 3



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LIST OF ABBREVIATIONS

AAS : Atomic Absorption Spectroscopy

Ag/AgCl : Silver-Silver Chloride Reference Electrode

BET : Brauner-Emmett-Teller

CSIR : Council for Scientific and Industrial Research

CV : Cyclic Voltammetry

EDS/X : Electron Dispersive X-ray Spectroscopy

EDTA : Ethylenediaminetetraacetic acid

EIS : Electrochemical Impedance Spectroscopy

EV : Electric Vehicle

FTIR : Fourier Transform Infrared Spectroscopy

GCD : Galvanostatic Charge-Discharge

HF : Hydrogen Fluoride

HR-SEM : High Resolution Scanning Electron Microscopy

HR-TEM : High Resolution Transmission Electron Microscopy

ICP-OES :Inductively Coupled Plasma Optical Emission

Spectroscopy

Li/Li⁺ : Lithium metal Reference Electrode

Li-ion : Lithium fon : Lithium

LRMO : Lithium Rich Manganese Oxide

MS : Mass Spectroscopy

NiCd : Nickel Cadmium

Ni-MH : Nickel-Metal Hydride

NMP : 1-Methy-2-Pyrollidone

N₂O : Nitrous Oxide

PdO : Palladium Oxide

PVDP : Polyvinylidene fluoride

PXRD : Powder X-ray Diffraction

REE : Rare Earth Elements

rGO : Reduced Graphene Oxide

SAED : Selected Area Electron Diffraction

SAXS : Small Angle X-ray Spectroscopy

TGA : Thermogravimetric Analysis

TM : Transition Metal

USD : United States Dollar

XPS : X-Ray Photoelectron Spectroscopy

XRD : X-ray Diffraction



LIST OF SYMBOLS

Å : Angstrom

 β : Beta

Rct : Charge Transfer Resistance

 Ω : Ohm

γ : Scan Rate

R_{SEI} : Surface Film Resistance

 θ : Theta

λ : Wavelength



CHAPTER 1

INTRODUCTION

1.1. Background of the Study

1.1.1. The global energy demands

The global energy demand is increasing at an alarming rate due to the rapid increase in the global population [1]. This rapid rise in the global population is associated with mounting industrialization and urbanization activities which have greatly increased the global energy demand [2],[3]. Currently, fossil fuel derived oil and gas play a major role in feeding this demand as estimates indicate that over 85% of the global energy demand is met by the burning of fossil fuels [4]. The transportation sector alone consumes over 30% of the global energy in the form of petroleum oil and gas [4], and it was reported that in 2017 approximately 64.5% of all the electricity generated was derived from oil, gas, and coal [6]. With the global population expected to rise even more to 8.5 billion in the year 2030, 9.7 billion in 2050 and 10.9 billion in 2100 [2], it is expected that more fuels will be needed to meet our rapidly increasing demand for energy.

1.1.2. The need for renewable energy

The use of fossil derived fuels to feed the ever-increasing demand for energy is becoming unsustainable and presents a myriad of environmental, economic, and health problems [7]. As depicted in figure 1.1, during the burning of fossil fuels, large amounts of greenhouse gases such as carbon dioxide (CO₂), methane (CH₄) and nitrous oxide (N₂O) are emitted into the atmosphere [7]. Naturally, these gases permit rays of light coming from the sun to reach the surface of the earth but trap the rays that are radiated back from earth surface [8]. As a result of the trapping of the reflected rays, the earth's average temperature increases, maintaining warm climatic conditions and this process is referred to as the greenhouse effect [8]. Scientists have reported that a more aggressive form of this phenomenon has been taking place around the world and is associated with increased greenhouse gas emissions from the burning of fossil fuels [9]. This human enhanced process has had some adverse effects on the

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environment such as melting of sea and continent glaciers, rise in the sea level, severe drought and rising average global temperatures [9].

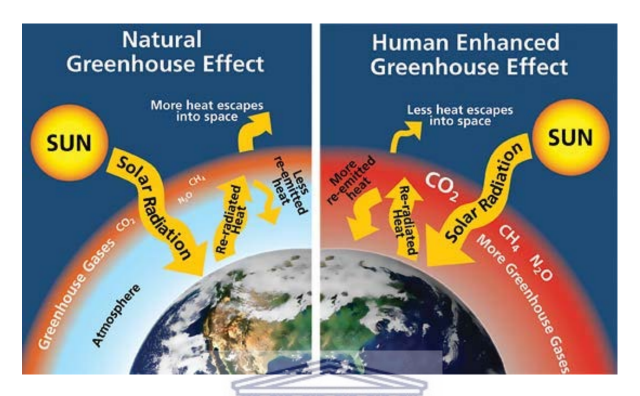


Figure 1.1: A comparison of the natural Greenhouse effect to human induced global warming [7].

Furthermore, the need for fossil fuels as primary sources of energy throughout the three industrial revolutions has triggered global socio-economic instabilities as rich countries scramble to increase their wealth. Consequently, the uneven distribution of global fossil fuel reserves raises concerns on 1) host country instabilities which can affect the oil price and 2) national energy security as a continuous and uninterrupted supply of energy is critical to the nation's economy [13]. Moreover, fossil fuels are non-renewable, and their complete depletion cannot be ruled out in the future [14]. Due to the above-mentioned factors, the scientific community has been tasked with the great responsibility of developing renewable and cleaner means of energy generation, to further mitigate climate change and environmental pollution associated with fossil fuel use.

1.1.3. The need for energy storage

To date, several renewable energy sources and technologies are being widely researched and developed as possible alternatives to the use of fossil derived fuels. For electricity generation, wind and solar energy which harness nature's clean energy have emerged as promising candidates due to their reduced carbon footprint in contrast to their non-renewable counterparts [15]. Despite being cleaner and cheaper per unit energy than fossil fuel derived oil, wind and solar energy generation also have limitations [3],[10]. The limitations lie in that the electricity generation is not constant but varies at different times and according to weather conditions. Thus electrochemical/mechanical energy storage systems are pivotal for wide range adoption of renewable energy sources and integration into the grid [11],[21]. On the other hand, electric and hybrid vehicles have shown great promise as possible alternatives to gasoline powered vehicles to curb and ultimately diminish the use of fossil fuels in the transportation sector [16]. Recently, the cost of owning an electric vehicle (EV) is on parity if not less than owning a gasoline powered vehicle when maintenance costs are factored in [17]. However, the feasibility of this technologies as replacements to conventional vehicles is challenged by the current cost of electric vehicle batteries and the battery life [18]. The realisation of the complete replacement of fossil fuels energy with cleaner energy sources is therefore dependent on the development of highly efficient energy storage technologies. There is ongoing research on implantation of different electrochemical energy systems in EVs and batteries as the favourite technology is maturing at an accelerated rate.

1.1.4. Lithium ion batteries for energy storage

Lithium ion batteries have attracted a great deal of attention in the research and development of large grid and electric vehicle energy storage systems [19],[20],[21],[24]. This is due to their higher specific energy, longer life span, power densities and their wide range of temperature versatility in applications in comparison to other commercial rechargeable batteries like lead acid, nickel cadmium (NiCd), nickel-metal hydride (Ni-MH) and sodium-sulphur batteries [9],[24] as shown in figure 1.2. Furthermore, Li ion batteries have shown a proven track record of reliability in the portable electronics market since their introduction in the 1990's [23] and continue to

enjoy great success with a global market size projected to reach USD (United States Dollar) 105.0 billion by 2025 [22].

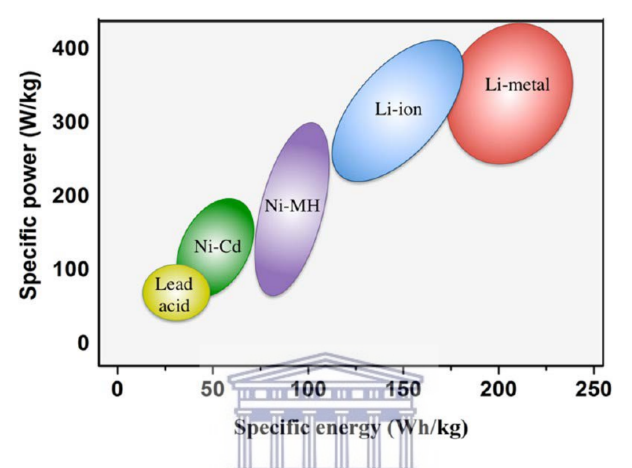


Figure 1.2: Ragone plot of the various battery technologies [9].

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However, currently available Li-ion batteries cannot meet all the requirements for applications in electric vehicles and electrical grid storage systems. These include, extended lifespan, low production cost, environmentally benign, high-energy density, and good safety [25]. For example, the best performing commercial Li-ion battery consists of a cobalt based cathode (LiCoO₂) which is both costly and toxic [26]. The limiting factor to higher capacity and performance in Li-ion batteries is the relatively low capacities of their cathode materials (LiCoO₂, LiNiO₂, LiMn₂O₄, LiFePO₄ and LiMnO₂) with specific capacities ranging from 100–180 mA h g ⁻¹ [27].

This work reports on the use of a novel nanostructured lithium rich manganese oxides (Li₂MnO₃) doped with europium (Eu) and surface coated with Palladium oxide (PdO) composite as high-capacity cathodes for potential grid storage and electric vehicle Liion batteries application. This is because of the low cost of manganese due to its

abundance, high theoretical capacities (459 mAhg⁻¹) and its low toxicity of the materials.

1.2. Problem Statement

The activation of Li₂MnO₃ cathode materials by the loss of lattice oxygen from the structure during the initial cycling results in the irreversible capacity loss and low initial coulombic efficiencies [28]. This occurs because of the detached lattice oxygen reacting with the electrolyte and decomposing it [28]. Furthermore, this oxygen loss destabilises the [MnO₆] octahedral symmetry and induces a phase transformation where transition metals migrate in favour of a more stable tetrahedral geometry (spinel) according to Jahn Teller distortion [29]. It has been suggested that to supress the capacity fading during cycling, the suppression of oxygen dimerization and phase transformation to spinel must be employed [29]. Herein, we report on the strategies that have been employed to suppress oxygen dimerization and phase transformation to spinel [29].

1.3. Motivation

The fabricated composite yields a synergistic relationship which shortens Li⁺ diffusion pathways and improves rate capability. Secondly, surface coating can suppress electrode/electrolyte side reactions and lastly rare earth element (REE) doping which can reduced charge transfer resistance and increase electric conductivity.

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1.4. Research Aims & Objectives

1.4.1. Research Aim

The aim of this study was to prepare, characterize and determine the electrochemical attributes of modified Li₂MnO₃ cathode materials for possible Li-ion battery application.

1.4.2. Objectives

- 1. Prepare Li₂MnO₃, Li₂Mn_{0.95}Eu_{0.05}O₃ and PdO@Li₂Mn_{0.95}Eu_{0.05}O₃.
- 2. Perform structural characterization of Li₂MnO₃, Li₂Mn_{0.95}Eu_{0.05}O₃ and PdO@Li₂Mn_{0.95}Eu_{0.05}O₃.

3. Evaluate the electrochemical performance of Li₂MnO₃, Li₂Mn_{0.95}Eu_{0.05}O₃ and PdO@Li₂Mn_{0.95}Eu_{0.05}O₃.

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CHAPTER 2

LITERATURE REVIEW

2.1. Literature Review

2.1.1. History of Li-ion batterie

Investigations into Li-ion based batteries began as early as the 1970's [1]. This was after the discovery that a Li-ion has the smallest radius in contrast with all the metals and can move reversibly between two interacted materials of different potentials when used as electrodes [1],[2]. This concept was quickly taken up and implemented by M.S Whittingham to produce a battery with lithium metal as an anode and titanium disulphide (TiS₂) as a cathode [2]. Although these types of batteries showed great promise with high energy densities, they had an inherent flaw in that the lithium metal anode exhibited uneven dendritic growth during cycling which ultimately raised safety

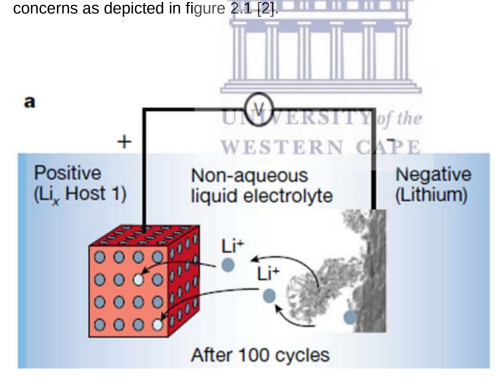


Figure 2.1: Schematic diagram of an early Li-ion battery showing Li dendrite growth on anode [4].

As a result of the mounting safety concerns of lithium metal anodes, a few alternative

anodes where proposed. The newly proposed anodes were fabricated by alloying

lithium with other metals, examples of such anodes included Li/AL and Li/Mn [1],[3].

The lithium alloy anodes lowered the rate of lithium dendritic growth but in the process

compromised the electrode potential and capacity of the battery [3]. This prompted

Goodenough in the 1980's to investigate new cathode materials with high voltages

which he found to be layered lithiated transition metal oxide of the formula LiMO₂

(where M= Co, Ni, Cr & V) [1],[2],[3]. Goodenough's discovery together with the

discovery of the highly reversible, low voltage Li intercalation-deintercalation process

in carbonaceous material paved the way for the world's first commercial rechargeable

battery by SONY corporation in 1991 [2],[4],[7].

2.1.2. Working principle of Li-lon batteries

The working principle of rechargeable Li-ion battery is based on the ability of the lithium

ions to move reversibly between the anode and cathode of a cell during charging and

discharging as depicted in figure 2.2 [5],[7]. During the charging process, lithium ions

intercalate into the carbonaceous intercalated anode (graphite) from the lithiated metal

oxide cathode (LiMO₂ where M= Co, Mn, Cr or V) [5],[6]. During the discharging

process, lithium ions intercalate back into the lithiated metal oxide cathode from the

carbonaceous intercalated anode (graphite) and this movement of the lithium is

generating a flow of electrons through an external circuit [5],[7]. This process can be

summarized the equations (2.1 - 2.3) below [6]:

Cathode: $LiCoO_2 \leftrightarrow Li_{1-n}CoO_2 + nLi^+ + ne^-(2.1)$

Anode: $C_6 + nLi^+ + ne^- \leftrightarrow Li_nC_6(2.2)$

Cell reaction: $LiCoO_2 + C_6 \leftrightarrow LinC_6 + Li_{1-n}CoO_2$, $E_0 = ~4 \text{ V}$ (2.3)

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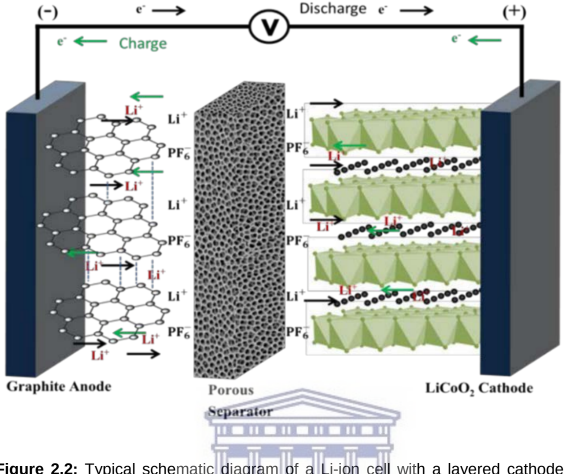


Figure 2.2: Typical schematic diagram of a Li-ion cell with a layered cathode and graphitic anode [6].

2.1.3. Components of Li-lon Batteries WESTERN CAPE

2.1.3.1. Electrolyte

An electrolyte is an important component of a cell that is localized between the electrodes of a cell. It is responsible for the migration of ions between the two electrodes of the cell which is associated with ionic charge transport [11]. Two types of electrolytes for Li-ion batteries exist, liquid electrolytes and solid type electrolytes. The most common type of electrolytes used to date are liquid electrolytes. Examples include lithium hexafluorophosphate (LiPF6), lithium perchlorate (LiClO4) or lithium tetrafluoroborate (LiBF4) [11]. The choice of the electrolyte used in a cell determines the electrochemical window of the cell. The electrochemical window of a cell is a voltage range in which the electrolyte will neither be oxidized or reduced. Properties of an ideal electrolyte are [8],[9]:

- Chemically stable/inert (non-reactive with electrodes and other cell components)
- High ionic conductivity
- Resistant to change in harsh conditions (thermal & mechanical)
- Large electrochemical window ~5 V vs Li⁺/Li.

2.1.3.2. **Anode**

The anode is a negative electrode within the cell at which the process of oxidation takes place, and an electron is released into the external circuit. In Li-ion batteries, lithium metal was found to be the best anode but was quickly replaced by carbonaceous intercalated materials due to safety concerns associated with the uneven dendritic growth during cycling. An ideal electrode must have strong reducing ability, be able to undergo lithium intercalation/deintercalation processes rapidly, have good conductivity and a low cost [12].

2.1.3.3. **Separator**

The separator is put in place within a cell to prevent electronic contact of the two electrodes and to allow free migration of ions from one electrode to another. Typically, separators are fabricated from Polyolefins, natural and/or synthetic fibres (non-woven separators) and ultrafine metal oxide or carbonate particles (inorganic composite separators) [11]. Ideal properties of separators include [11]:

- Chemical stability.
- A relative thickness that will exhibit porosity < 1mm.
- Uniform permeability.

2.1.3.4. **Cathode**

In Li-ion batteries, the physico-chemical properties of the battery are directly dependent on the type of the cathode and anode materials used [14], [17], [18]. These properties subsequently have a greater influence on the performance and the overall cost of the battery [19]. For example, the cathode material used in commercial Li-ion batteries such as LiCoO₂, Li₂MnO₄ and LiFePO₄ have specific capacities of 100~180 mAh.g⁻¹ which are significantly lower as compared to the anode material graphite

which has a specific capacity of ~373 mAh.g⁻¹ [20]. This implies that in order to significantly improve battery performance, the cathode materials become the first component to fine tune because it has a lower specific capacity compared to the anode and the Li-ion transfer when charging and discharging is mainly controlled by the cathode [17],[19].

One family of cathode materials that has generated worldwide interest for Li-ion battery application in electric vehicles, hybrid vehicles and off grid storage is the lithium manganese oxides[21]. This is a result of the low cost of manganese due to its abundance, high theoretical capacities, and the low toxicity of the materials [21]. One such material is the lithium manganese oxide Li₂MnO₃ which possesses a superior theoretical capacity of 459 mAh.g⁻¹[22].

Due to its insulator-like properties, Li₂MnO₃ was thought to be electrochemically inactive [23],[24]. It was in 1991 when Thackeray *et al.* [25] at the CSIR discovered that a new type of electrochemically active layered oxide could be produced by simply leaching out LiO₂ from Li₂MnO₃ by acid treatment at room temperature. When electrochemically cycled, they obtained 0.2Li₂MnO₃0.8LiMnO₂ layered material which ushered the dawn of a new Li₂MnO₃ based layered-layered cathode materials and the notation xLi₂MnO₃(1-x)LiMnO₂ [25].

In another study, it was reported that Li₂MnO₃ could also be activated electrochemically by charging to >4.5 V [25]. These lithium rich manganese oxides (LRMOs) possess specific capacities of up to 280 mAh.g⁻¹ [25]. It has been proposed that the Li₂MnO₃ component stabilizes the structure at lower voltage regions and can also provide addition lithium ions in the high voltage region [25]. Nevertheless, the large-scale application of these LRMOs cathode materials remained limited due to cutoff voltage for activation, irreversible capacity loss in the first cycle and transition metal migration which induces phase transformation during repeated charge-discharge cycles which have been attributed to the Li₂MnO₃ component [25], [26].

As a result of the factors discussed above, it becomes imperative to investigate ways to improve the electrochemical properties of Li₂MnO₃ to fabricate higher capacity Li₂MnO₃ and LMRO cathodes. This review presents research advances in the modification of Li₂MnO₃ to suppress irreversible capacity loss and improve-capacities.

It further presents 1st principle studies of proposed modifications for high capacity Li₂MnO₃ derivatives as well as reported experimental work. Furthermore, it discusses in depth the modification strategies such as Mn-ion substitution, surface coating, surface treatment, combined modifications, and alternative phase embedment. Finally, it also explores the significance of these Li₂MnO₃ modifications, and the effectiveness of these modification techniques are also being compared.

2.1.3.4.1. Li₂MnO₃ structure and electrochemical activity origin

The layered manganese oxide Li_2MnO_3 can be expressed as $Li[Li_{0.33}Mn_{0.66}]O_2$, indicating that it possesses an α -NaFeO₂ type structure with the space group of C2\m monoclinic [20],[27]. In this structure, alternating lithium, close cubic packed oxygen, and transition metal layers are stacked one on the other in an ABCABC stacking order as represented in figure 2.3. From the notation $Li[Li_{1/3}Mn_{2/3}]O_2$, it becomes evident that 1/3 of the inter-slab octahedral sites are occupied by Li^+ and the remaining 2/3 by Mn^{4+} . In other words, 1/3 of the Mn^{4+} in the transition metal (TM) layer are replaced by Li^+ [23],[28]. As a result of this occupancy of Li^+ in the TM layer, a honeycomb configuration consisting of 6 Mn^{4+} surrounding 1 Li^+ exists and can be observed as a superlattice using X-ray diffraction (XRD) as shown in figure 2.4 [20],[27],[29]. This super lattice is characteristic of Li_2MnO_3 materials and can be observed using XRD (figure 2.4) at $20 - 25^{\circ}$ 2θ angles [30],[31].

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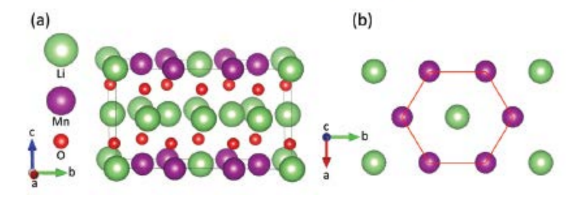


Figure 2.3:(a)Schematic representation of the crystallographic structure of Li₂MnO₃ (b) Honeycomb configuration of 6 Mn⁴⁺ surrounding 1 Li⁺ in the TM layer of Li₂MnO₃ [28].

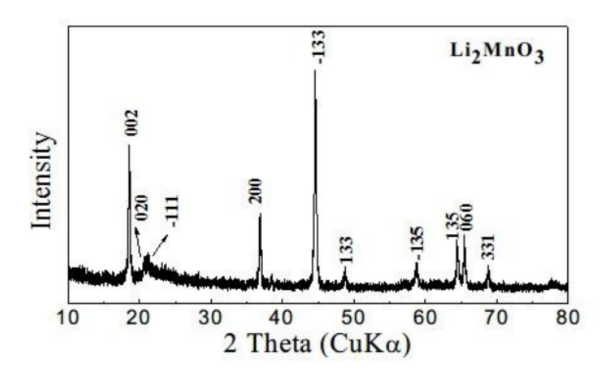


Figure 2.4: XRD pattern of monoclinic Li₂MnO₃ [27].

It is generally accepted that in lithiated manganese oxide cathode materials such as LiMnO₂ and LiMn₂O₄, the electrochemical activity is associated with Li⁺ intercalation and deintercalation which is accompanied by the oxidation of manganese from Mn³⁺ to Mn⁴⁺ [24]. Thus, for a very long time Li₂MnO₃ was thought to be electrochemically inactive due to Mn⁴⁺ being unable to oxidize to higher state of Mn⁵⁺. Today it is well established that Li⁺ ions can successfully be deintercalated and intercalated back into this material, although the origin of the electrochemical activity of Li₂MnO₃ is still a highly debatable subject.

The origin of the electrochemical activity of Li₂MnO₃ is still highly debatable, complex in nature and yet to be fully understood. Initially, the charge compensation in Li₂MnO₃ was attributed to the O losing electrons in order to achieve charge neutrality during the first charge (>4.5 V vs Li/Li⁺). The proposed equation 2.4 was given as:

 $Li_2MnO_3 \rightarrow Li_2O + MnO_2$ (2.4)

Robertson *et al.* [24] did an extensive study on the electrochemical activity of Li₂MnO₃ and suggested that the activity could be attributed more to the ion exchange between the H⁺ proton from the electrolyte and Li⁺ ion from Li₂MnO₃. Similar results were observed even in anhydrous organic electrolytes such as LiPF₆ which have been known to produce hydrogen fluoride (HF) when interacting with even small volumes of water [24]. From this work, it was concluded that the mechanism in Li₂MnO₃ electrochemical activity is the compensation of charge between Li⁺ and H⁺ as opposed to the loss of O²⁻. This conclusion was backed by flame emission spectroscopy (FES), atomic absorption spectroscopy (AAS), X-ray photoelectron spectroscopy (XPS) and thermogravimetric analysis/mass spectroscopy (TGA/MS) results of the sample measured during charge and discharge as shown in table 2.1.

Table 2.1: FES, AAS, XPS and TGA/MS results of the sample measured during charge and discharge [24].

Sample	Calculated Li:Mn	Theoretical	Calculated H	Bulk Composition
	ratio	Li:Mn ratio		
1 st charge 55°C	Li _{0.62} Mn	Li _{0.65} Mn	H _{1.29}	Li _{0.62} H _{1.29} MnO _{2.95}
1 st discharge 55°C	Li _{1.28} Mn	Li _{0.65} Mn	H _{0.66}	Li _{1.29} H _{0.66} MnO _{2.97}

From the first charge, the electrolyte undergoes oxidation to generate H⁺ protons which undergo ion exchange with Li⁺ ions derived from the electrode surface. This implies that H⁺ protons now coexist with Li⁺ ions within the structure of the material. Upon first discharge, the electrolyte reduction process drives the extraction of the H⁺ protons from the structure and Li⁺ ions are intercalated back into the structure. From the table above it can be observed that the loss of oxygen is minimal and thus cannot be the dominant mechanism in the electrochemical activity of Li₂MnO₃.

Although there is significant evidence for the Li⁺/H⁺ ion exchange mechanism during electrochemical cycling, some reports dispute this being the main mechanism for the activity of Li₂MnO₃ [32],[33]. It is suggested that this mechanism is dominant at elevated temperatures where electrode degradation can take place [32]. Yu *et al.* [32] reported that during the de-lithiation process at room temperature, oxygen gas is generated, accounting for ~1/8 moles from Li[Li_{1/3}Mn_{2/3}]O₂ thus further reinforcing the

oxygen loss hypothesis. Chen *et al.* [33] supported the oxygen loss mechanism by suggesting that Li extraction is accompanied by oxygen loss which generates holes on oxygen. These holes are not stable and are stabilized by the formation of molecular oxygen which facilitates the migration of Mn ions onto vacant octahedral sites in the lithium layers and generates a spinel type material [33]. The equation and mechanism supporting the observation are given in equation 2.5 and figure 2.5 respectively.

$$Li_2MnO_3 \rightarrow 2Li^+ + 2e^- + MnO_2 + O_2$$
 (2.5)

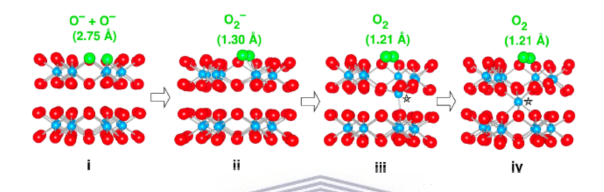


Figure 2.5: Oxygen loss mechanism suggesting that Li extraction is accompanied by oxygen loss which generates holes on oxygen which result in oxygen evolution [33].

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2.1.3.4.2. Challenges facing Li₂MnO₃ as cathode materials

The activation of Li₂MnO₃ cathode materials by the loss of lattice oxygen from the structure during the initial cycling results in the irreversible capacity loss and low initial coulombic efficiencies [27],[28],[29]. This occurs as a result of the decomposition of the detached lattice oxygen through its reaction with the electrolyte. Furthermore, this oxygen loss destabilizes the [MnO₆] octahedral symmetry and induces a phase transformation where transition metals migrate in favour of a more stable tetrahedral geometry (spinel) in accordance with Jahn Teller distortion [28]. It has been suggested that in order to minimize capacity fading during cycling, approaches related to the suppression of oxygen dimerization and phase transformation to spinel must be carried out. Herein, we report on such strategies.

2.1.3.4.3. Li₂MnO₃ Modification Strategies

2.1.3.4.3.1. Particle Size Confinement

Nanoscale confinement has been reported to significantly enhance thermodynamic and kinetic properties of Li-ion cathode materials [34]. Nanostructures have shorter Li⁺ diffusion pathways which can improve the rate capability during Li ion intercalation/de-intercalation processes [35], [36]. Shortened diffusion pathways offer high power density and reduces the time it takes to reach-active sites, resulting in enhanced reaction kinetics [36]. On the other hand, nanostructures (table 2.2) exhibit large specific surface areas which can expose more active sites thus increasing contact between the electrode material and electrolyte leading to greatly improved capacities of the cathode material [37],[38]. Moreover, nanostructures have more defects which can lead to enhanced stability of the structure during cycling and decrease potential barriers [28],[29],[37]. The above-mentioned improvements can be observed in table 2.2, which shows a comparison of the different improvements observed by the nano structuring process using different synthesis approaches.



Table 2.2: Comparison of different Li₂MnO₃ nanostructures and their enhancements.

Morphology	Synthesis	Initial	Initial	Retained	Rate	Particle	Measurement Conditions	Ref.
	Method	Discharge	Coulombic	Capacity	Capacity	Size		
		Capacity	efficiency	(mAh.g ⁻¹)	(mAh.g ⁻¹)	(nm)		
		(mAh.g ⁻¹)	(%)					
Nanoparticles	Solid state	335	100	291	245 (46 mA. g ⁻¹)	10	2.1- 4.4 V, 11.5 mA. g ⁻¹ , 3 cycles, RT	[39]
& Nano cubes – Mix	Hydrothermal	323	70	~187.5	-	10-50	2.0 - 4.3 V, 60mA.g ⁻¹ , 20 cycles, RT	[27]
Nanoparticles	Solid state	241.9	7 5.4	182.39	187.23(0.2C)	10-20	2.0-4.6V, 0.1C, 45 cycles, RT	[40]
-	Oxidation reaction	236	78.15	~187.5	100 (914.3mA.g ⁻¹)	~10	2.0-4.9V ,14.3 mA. g ⁻¹ , 30 cycles, RT	[41]
_	Sol gel	173	-	~105		10~40	2.0-4.6 V, 30 cycles, RT	[42]
_	Solid state	124.9	- 2	106.2	80.4 (1/20C)	30-80	2.0-4.6V, 100 cycles, 10mA.g ⁻¹ , RT	[30]
Nanowires	Molten salt	~156	τ	159.8 JN IV E I	~200 (0.04C)	10-25	2.0 - 4.8 V, 20 mA. g ⁻¹ , 27 cycles, RT	[43]
-	Solid State	~80	~50 \	VE 31TE	R 100(5C)	E 50	2.0-4.5V, 100 cycles	[44]
Nanoplate	Precipitation	94	-	82	-	~15	2.0 - 4.8 V, 50 cycles, RT	[45]
Nanoribbon	Ion exchange and oxidation	74	52.11	125	-		2.0 – 4.8 V, 10 mA·g ⁻¹ , 6 Cycles, RT	[46]
Nanobelts	Ion exchange and oxidation	61.2	33.6	~74	-	15	2.0–4.8 V, 200 mA·g ⁻¹ , 30 Cycles, RT	[29]

For instance, Liu *et al.* [27] prepared nanostructured Li₂MnO₃ particles by self-seeding hydrothermal methods and obtained well crystallized nanorods and nano cubes with a size range of 10-50 nm as can be observed in figure 2.6 (a). The nanostructures exhibited an initial discharge capacity of 323 mAh.g⁻¹ and a specific capacity of 243 mAh.g⁻¹ within the voltage window 2.0-4.3 V. In another study, the molten-salt method was used to synthesize Li₂MnO₃ nanowires (figure 2.6 (b)) from Mn₂O₃ nanowire precursor [43]. The nanowires showed an initial discharge capacity of 156 mAh.g⁻¹ compared to 53 mAh.g⁻¹ from the bulk Li₂MnO₃ in the potential window 2.0 to 4.8 V and current density of 20 mAh.g⁻¹ [43]. After 27 cycles, the nanowires retained a discharge capacity of 159.8 mAh.g⁻¹. In both studies, the exceptional capacities are attributed to shorter Li⁺ diffusion pathways and high surface areas which exposed more particles to the electrolyte leading to a lower activation potential.

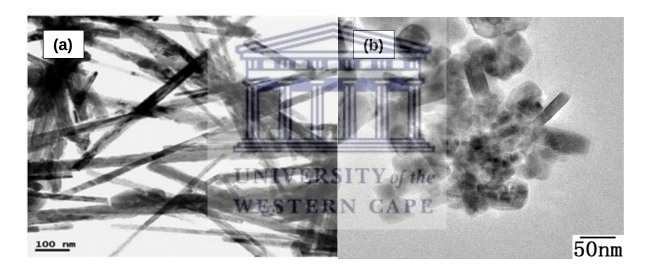


Figure 2.6: TEM images of (a) Li₂MnO₃ Nanowires and (b) Li₂MnO₃ Nanorods and nano cubes [27] [43].

Freire *et al.* [39] reported on nanostructured Li₂MnO₃ obtained by solid state reaction of Li₂O and MnO₂ followed by high energetic grinding at 700 rpm for 20 hours. From TEM studies, the average crystallite size was found to be 10nm in diameter indicating that the material was nanostructured [39]. The initial discharge capacity exhibited was 335 mAh.g⁻¹ with no observed drop in capacity during discharge in the potential window 1.2 - 4.4 V vs Li⁺/Li at C/40. After 3 cycles, the material showed a reversible

capacity of 291 mAh.g⁻¹ and a potential of 3 V at C/40 [39]. More interestingly, even at higher rates, the material still exhibited exceptionally high discharge capacities indicating good cycling stability. The authors attributed the high capacity to the first oxidation process (charging to 4.4 V) which induced disordering in the material.

HRTEM analysis (figure 2.7) of the sample after initial cycling indicated a phase transformation from monoclinic Li₂MnO₃ to a disordered nanostructured material [39]. Most of the sample now contained disordered nano-sized cubic crystals along with traces of pristine monoclinic Li₂MnO₃. It is understood that this transformation to nanosized cubic crystals improved Li ion diffusion paths.

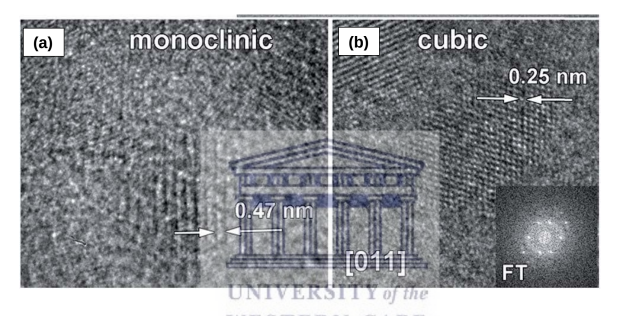


Figure 2.7: HRTEM images of a) single monoclinic and b) cubic nanoparticles [39].

Sayle and co-workers [37] in their 2015 study suggested that the enhancement of electrochemical activity of nanostructured Li₂MnO₃ may also originate from the stabilisation via 'point defect scaffold as shown in figure 2.8. From their Molecular Dynamics simulations, it was determined that the nanostructured Li₂MnO₃ had a high degree of defects and exhibited Li, Mn, and O vacancies with a mix of Li and Mn in all layers (figure 2.8). Unlike in the bulk Li₂MnO₃ which consists of alternating Li ion and transition metal layers, the nanostructured Li₂MnO₃ contained transition metal layers which are held apart by Mn ions (point defects) in the Li ion layer. This Mn ions stabilise the structure of the material during deintercalation thus preventing structural collapse and allowing the smooth diffusion of Li ion in and out of the structure.

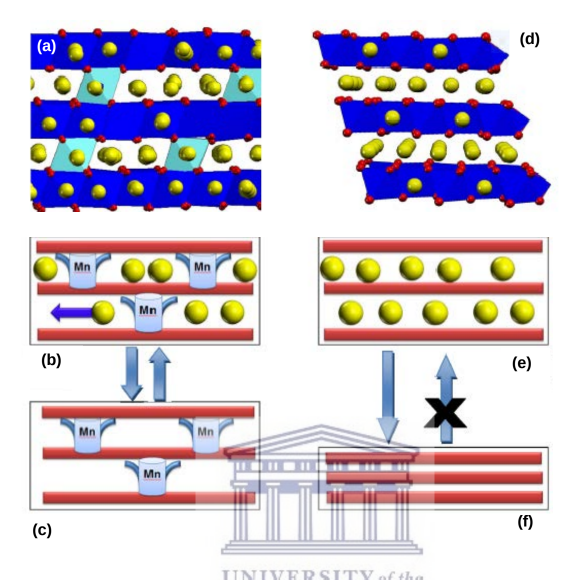


Figure 2.8: Model structures of the proposed stabilization by point defect scaffold [37].

From table 2.2, it is evident that the synthesis method and conditions also play a pivotal role in the electrochemical performances of the nanostructured materials. The synthesis method determines the number of defects (staking faults, Li, Mn or O vacancies and cationic site mixing) which are reported to improve performance of the cathode material [29],[37]. Menon *et al.* [47] reported a similar finding when he compared solid state and sol gel synthesized Li₂MnO₃. Sol gel synthesized Li₂MnO₃ contained more staking faults and exhibited a superior performance compared to the solid-state counterparts [47]. An enhanced performance due to Li and Mn cationic mixing and defects was also observed by Matsunaga *et al.* [48] in their report, where a low synthesis temperature induced Li/Mn cation mixing defects which promoted the smooth diffusion of Li ions.

Table 2.3: Comparison of different coating materials and their enhancements.

Class of Coating	Coating	Initial	Initial	Retained Capacity	Rate	Measurement	Ref.
		Discharge	Coulombic	after cycling	Capacity	Conditions	
		Capacity	Efficiency	(mAh.g ⁻¹)	(mAh.g ⁻¹)		
		(mAh.g ⁻¹)					
			(%)				
Metal Oxide	TiO ₂	Down from	Maintained at	Up from 81.4 to	Up from 56.54 to	2.1- 4.4 V, 20mA.g ⁻¹ ,	[39]
		212.3 to	100	99.5	99.56	100 cycles, RT	
		206.8			(100 mA. g ⁻¹)		
	Co ₃ O ₄	Up from 85	Up from 60.28	Up from ~40 to 135	-	2.0 – 4.8 V, 10mA.g ⁻¹ ,	[52]
		to 178	to 97.27			50 cycles, RT	
Phosphate	FePO ₄	Up from 94	Up from 45.19	Up from 82	-	2.0 and 4.8 V, 10mA.g ⁻¹ ,	[45]
Coating		to 180	to 91	to 175		50 cycles, RT	
Carbonaceous	Sucrose derived	Up from		·t	Jp from 51.8 to 117.5	2.0 - 4.8 V, 10 mA. g ⁻¹ ,	[51]
material	Carbon	229.1 to	UNIVE	ERSITY of the	(400 mA. g ⁻)	27 cycles, RT	
		280.5	WEST	ERN CAPE			
	PVP derived	Down from	Up from 59.3 to	Up from 126.5 to	Up from 0.3 to 111	2.0 – 4.8 V, 20 mA. g ⁻¹ ,	[53]
	Graphitic	239.3 to 206	61.9	196.3	(100 mA. g ⁻)	10 cycles, RT	
	Carbon						
Active Cathode	LiCoO ₂	Up from 74	Up from 52.11	Up from ~ 110 to	-	2.0 - 4.8 V, 10 mA. g ⁻¹ ,	[46]
Material		to 180	to 72	180		30 cycles, RT	

2.1.3.4.3.2. Surface Coating

Surface coating is one of the most common methods of improving the electrochemical performance of layered manganese oxide cathode materials [49],[50],[51]. Table 2.3 summarizes some of the reported improvement in the electrochemical performance of layered manganese oxides using different coating materials. Oxygen loss and transition metal dissolution occur at the surface of the electrode and thus a suitable coating improves the performance by providing a protective layer which suppresses side reactions between the electrode and the electrolytes as well as prevents migration and dissolution of transition metals [45],[49],[50]. Furthermore, some coatings can react with the surface of the cathode to provide an active layer which improves capacity and stability [45]. When considering surface coatings, it is important to also note that the thickness of the coating plays a crucial role. In a coating such as TiO₂, the increase in TiO2 thickness correlates with the formation of a LixTiO2 species, which exhibits poor electrical conductivity and leads to decreasing discharge capacities [39]. On the other hand, FePO₄ is a direct opposite of this observation. Li₂MnO₃ coated with increased amounts of FePO₄ shows an increase in discharge capacities since it can produce Li_xFePO₄ which is electrochemically active [45]. In the case of carbon coatings, a thin coating is desirable as carbon coating induced the formation of Mn3+ which helped promote electron transport [50]. Thick coating led to high Mn3+ concentrations, which compromise the stability of the material [51].

Metal Oxide Coating

Kim *et al.* [52] synthesized pristine Li₂MnO₃ via sol gel method and proceeded to coat the resulting Li₂MnO₃ with anatase (TiO₂) to form a shell of controlled thickness as shown in figure 2.9. In order to obtain the optimum TiO₂ ratios, oxide coating variations from 4.8 -13.3 wt.% were investigated. The rationale behind TiO₂ as a coating material was based on the difficulty of TiO₂ in reacting with the electrolyte at room temperature according to equation 2.5:

$$TiO_2 + 6HF + 2e^- \rightarrow H_2TiF_6 + 2H_2O$$
 (2.5)

From their results, it was found that the coating had no enhancement effect on the initial cycling capacity but rather the initial discharge capacity of the material diminished with increasing TiO₂ content and was attributed to the thickness of the coating. Nonetheless, the coated samples showed an improved cycling stability after 100 cycles. The best performing sample showed a discharge capacity of 125.2 mAh.g⁻¹ with a capacity retention of 66.3% as compared to 81.4 mAh g⁻¹ at 38.3% shown by uncoated Li₂MnO₃ [52]. In this case, the coating prevented side reactions at the electrode/electrolyte interface as well as Mn dissolution into the electrolyte.

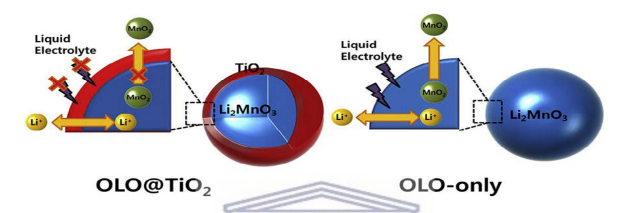


Figure 2.9: An illustration of how a surface coating such TiO₂ affects Li₂MnO₃ [54].

Phosphate Coating

In certain instances, some surface coating materials can improve both the cycling stability and the initial cycle discharge capacity. Such a behaviour was observed when Wang *et al.* [45] produced a core-shell type nanocomposite which consisted of a Li₂MnO₃ nanoplate coated with FePO₄. In their report, it was observed that the coating improved the cycling stability as no significant capacity fading was observed in the coated Li₂MnO₃ nanoplate with cycling. The FePO₄ coating thus prevented electrode corrosion via side reactions at the electrode/electrolyte interface. As opposed to the TiO₂ coating reported by Kim *et al.* [45] which is an "inert" coating, the FePO₄ coating also acted as an active layer which diminished 1st cycle capacity loss and had a specific capacity of 180 mAh.g-1 (figure 2.10).

Moreover, the FePO₄ coating doubled as a Li-ion host for Li ions that cannot be reinserted into the Li₂MnO₃ and thus exhibited a superior initial cycle discharge capacity of 208 mAh.g⁻¹ as shown in figure 2.10. The intercalation of Li ions into the

FePO₄ was evidenced by the presence of a reversible redox peak at 2.7/3.0 V vs. Li⁺/Li which did not occur in the uncoated counterpart. It is also suggested that small amounts of Fe³⁺ from FePO₄ might have been incorporated into the Li₂MnO₃ structure further improving its capacity. Although the suppression of electrode corrosion and the lowered charge transfer resistance can be attributed to surface coating, the nanoscale confinement effect must not be ignored.

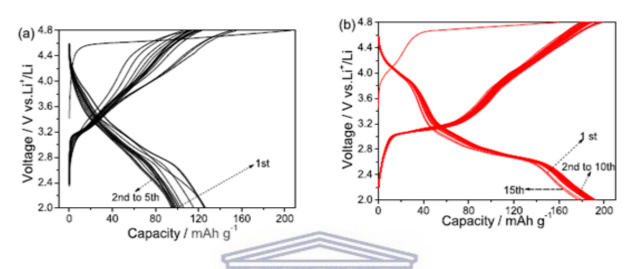


Figure 2.10: Charge/discharge curves of Li₂MnO₃ nanoplates (a) and Li₂MnO₃@FePO₄ nanoplates [45].

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Carbon coating

Surface coating with carbon & carbonaceous materials has its merits as a low-cost method of improving cathode material performance as they possess excellent conductivity [40],[55],[56]. In one publication by Xiong *et al.* [51], Li₂MnO₃ was produced by solid state reaction of Li₂CO₃ and MnCO₃ followed by calcination in air at 500 °C for 4 hours. The obtained powders were mixed with sucrose in ethanol and water while stirring, dried and then calcined at 350 °C for 20, 40 and 60 minutes, respectively [55]. When electrochemical studies were carried out, the carbon coating had a significant effect on the initial cycle discharge capacity of the Li₂MnO₃ and on the rate capacity. The best performing sample delivered a discharge capacity of 280.5 mAh.g⁻¹ and 117.5 mAh.g⁻¹ at 10 mA.g⁻¹ and 400 mA.g⁻¹ current densities respectively as shown in figure 2.11 [55]. The pristine sample on the other hand delivered lower capacities of 229.1 mAh.g⁻¹ and 51.8 mAh.g⁻¹ at 10 mA.g⁻¹ and 400 mA.g⁻¹ current densities respectively [55]. The high conductivity of the carbon

infrastructure on the surface of Li₂MnO₃ enhanced the electron pathway thus reducing charge transfer resistance.

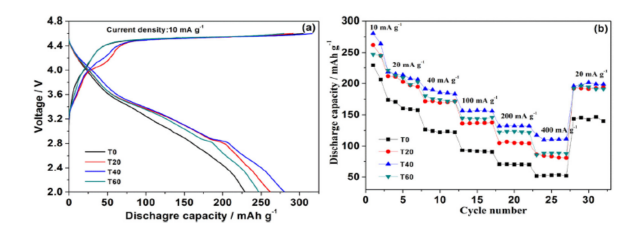


Figure 2.11: (a) Charge/discharge curves of Li₂MnO₃ and various Li₂MnO₃@C samples (b)Cycling stability testing of Li₂MnO₃ and various Li₂MnO₃@C samples [55].

Active cathode material coating

Wang *et al.* [46] produced coaxial LiCoO₂@Li₂MnO₃ nanoribbons which exhibited initial charge and discharge capacities of 270 mAh.g⁻¹ and 180 mAh.g⁻¹ respectively. Even after 30 cycles, the material was able to retain a 180 mAh.g⁻¹, discharge capacity in the potential window 2.0 - 4.8 V vs Li⁺/Li as shown in figure 2.12 (b). The exceptional performance was attributed to the LiCoO₂ coating which provided additional Li⁺ during insertion, suppressed active material corrosion and to the redox couple Co³⁺/Co⁴⁺shown in figure 2.12 (a) which can increase the energy density.

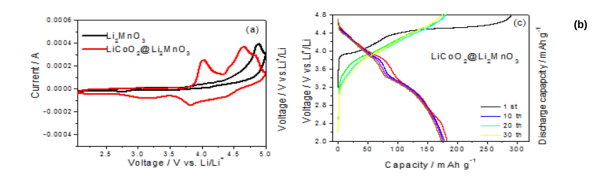


Figure 2.12: (a) CV voltammogram of Li₂MnO₃ and LiCoO₂@Li₂MnO₃, with the latter showing additional 3.8/4.0 V plateau attributed to the redox couple Co³⁺/Co⁴⁺. (b) Charge/discharge curve of LiCoO₂@Li₂MnO₃ [46]

2.1.3.4.3.3. Li/Mn/O site doping

Elemental doping has proved to be a good strategy to improve electrochemical performance of Li₂MnO₃ cathodes and table 2.4 summarizes some of the reports on the strategy. Doping can suppress oxygen loss and prevent structural transformation in Li₂MnO₃ which can lead to enhanced structural stability, electronic and ionic conductivity. In most cases, the least amount of dopant used gave the best results. When excess amounts of dopants are used, the crystal structure of the materials becomes distorted thus leading to instability and ultimately capacity loss. Therefore, doping with minimal amounts of dopants can suppress oxygen release and structural transformations which are responsible for the irreversible capacity loss during 1st cycling.

Transition metal doping

One of the first cases of transition metal doping was when Mori *et al.*[58] doped Li₂MnO₃ with varying amounts of Ruthenium (Ru) and observed a decrease in the electrical resistivity. The Li₂Mn_{0.4}Ru_{0.6}O₃ exhibited a discharge capacity of 192 mAh.g⁻¹ in the 1st cycle and 169 mAh.g⁻¹ in the 10th cycle at 1/10 C [58]. Since Mori *et al.* reported on Li₂Mn_{0.4}Ru_{0.6}O₃, several publications report on the doping of Li₂MnO₃ with transition metals (e.g., Fe, Co, Ti, Ni, Zr etc.) for enhanced performance [59],[62],[63],[64], [71].

In one first principle study, Gao *et al.* [65] showed that doping Li₂MnO₃ with molybdenum (Mo) greatly enhanced its properties as it resulted in a reduced band gap while at the same time inducing an increased number of electronic states in proximity of the Fermi level. As a result of this, Li-ion diffusion would increase, and the stability of oxygen would be enhanced leading to improved stability of the cathode material [65]. Ma *et al.* [66] also reported on the doping of Li₂MnO₃ with Molybdenum and obtained initial charge and discharge capacities of ~300 mAh.g⁻¹ and ~175 mAh.g⁻¹, respectively. The performance was attributed to the Mo ion expanding the lattice thus allowing increased Li⁺ diffusion and lowering of de-lithiation potential which inhibits lattice oxygen loss.

 Table 2.4: Comparison of different dopants materials and their enhancements.

Class of dopant	Dopant	Initial	Initial Coulombic	Retained	Rate	Measurement Conditions	Ref.
		Discharge	Efficiency	Capacity	Capacity		
		Capacity	(%)	(mAh.g ⁻¹)	(mAh.g ⁻¹)		
		(mAh.g ⁻¹)					
Transition Metal	Al	Up from 84 to 99.4	Up from 50 to 58.9	Up from	Up from 84.8 to 97.1	2.0 and 4.8 V, 20mA.g ⁻¹ , 40 cycles,	[57]
				71.23 to		RT	
				96.52			
	Ru	Up from ~25 to 192	Down from ~71.43 to	Improved	Improved	2.0 - 4.8 V, 20 mA. g ⁻¹ , 10 cycles,	[58]
			~70.33			RT	
	Ni	Up from ~28 to ~175	Up from ~56 to ~58.3		-	2.0 - 4.8 V, 10mA.g ⁻¹ , RT	[59]
	Мо	Up from ~37.5 to 175	Down from ~72.12 to		- 11	2.0 - 4.8 V, 10mA.g ⁻¹ , RT	[60]
			56				
Halogen	F	Up from 108 to 200	Down from 118.38 to	Up from	Up from 66 to 125	2.0 - 4.6 V, 10mA.g ⁻¹ ,100 cycles, RT	[61]
			90.09	~94.7 to	(16C)		
			UNIVER	S 184 Y	of the		
Alkali/Alkali Earth	Na	Up from 124.9 to 161	Up from 100 to	Up from	Up from 80.4 to 140.6	2.0-4.6V, 10mA.g ⁻¹ , 100 cycles, RT	[30]
Metals -			268.33	106.2 to	(1C)		
				158.8			
	Mg	Up from 241.9 to	Up from ~74.20 to	Up from	Up from 102 to 123.1	2.0-4.6V, 0.1C, 30 cycles, RT	[16]
		307.5	~87.85	187.96 to	(1C)		
				259.84			

Interestingly, aluminium (AI) doping at the manganese site was also found to greatly enhance electrochemical performance of Li₂MnO₃ [57]. In one study by Xiang *et al.* [57], it was revealed that due to the strong AI-O bond in the doped sample, the doping resulted in a weaker initial cycle capacity for the AI-doped sample. After the initial cycle, the cycling stability of the AI-doped sample is enhanced as depicted in figure 2.13 and the discharge capacity exceeded that of the pristine material. The EIS result of the AI-doped sample shows a reduction in the charge transfer resistance (Rct) from 10037 to 5185 Ω indicating an increase in the charge transfer within the material. These results are in agreement with Kuganathan *et al.*'s findings which indicate that AI is a supreme dopant due to its small ionic radius, solution enthalpy and its trivalent oxidation state which can assist in creating an extra interstitial Li⁺ion in Li₂MnO₃ [19].

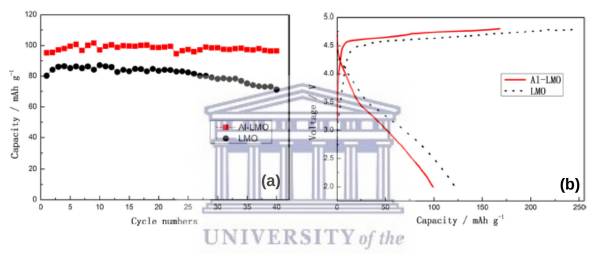


Figure 2.13: Cycling stability testing curves of Li₂MnO₃ and various Al-Li₂MnO₃ samples [57].

Matsunaga and co-workers also studied the effect of nickel- doping in Li₂MnO₃ compounds with varying Ni concentrations [59]. In their report, it was established that the charge and discharge capacities increased with the concentration of the Ni dopant (figure 2.14) [59]. This increase in performance was attributed to the ability of the nickel ion to lower the charge plateau potential therefore leading to higher first cycle charge capacities of up to ~300 mAh.g⁻¹ and discharge capacities of up to ~ 175 mAh.g⁻¹ [59].

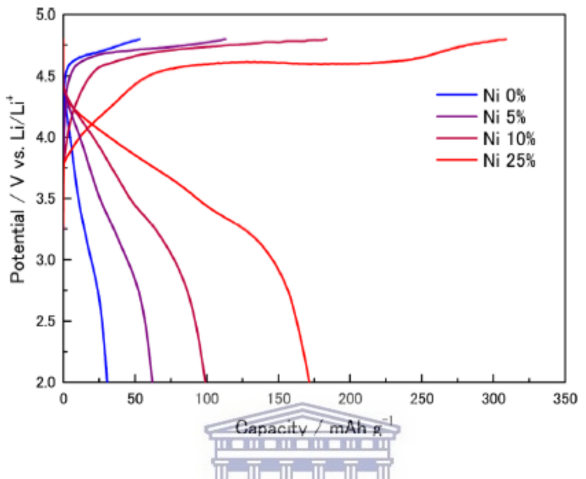


Figure 2.14: Initial Charge/discharge curves of Ni-Li₂MnO₃ [59].

Alkali/Alkaline-earth metal doping VERSITY of the

Alkali/alkaline earth metal doping has been reported in several publications and focuses on the metal substitution at the lithium site as opposed to the manganese site [16], [67]. For example, figure 2.15 shows that substitution of some lithium ions with magnesium ions was performed to yield a material with the formula Li_{1.98}Mg_{0.01}MnO₃ which exhibited a higher initial discharge capacity of 307.5 mAh g⁻¹ which exceeded the 241.9 mAh g⁻¹ of the pristine Li₂MnO₃ material [16]. Improvement in the cycling stability was observed whereby an 84.5% capacity retention was recorded for the doped material [16]. The authors attributed the superior performance to the reduction in charge transfer resistance and the increase in the lithium ion diffusion coefficient [16].

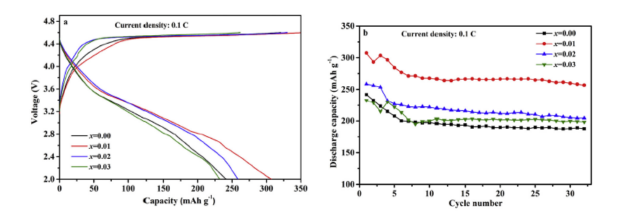


Figure 2.15: (a) Initial charge/discharge curves of Mg-Li₂MnO₃ and (b)Cycling stability testing of Mg-Li₂MnO₃ [16].

A similar type of substitution was performed by Dong *et al.* [30] when he synthesized the material Li_{1.90}Na_{0.10}MnO₃ by the partial substitution of lithium with sodium. The cathode material delivered a discharge capacity of 181 mAh.g⁻¹ in the 1st cycle and exhibited a high-capacity retention of 99.3% after cycling it 45 times at 1/10 C [30]. After 100 cycles at 1/2 C, a discharge capacity of 161 mAh.g⁻¹ was obtained indicating a 98.6% capacity retention [30]. The charge transfer resistance also decreased from 50.8 to 12.7 Ω .

Halogen Doping

Partial oxygen substitution with fluorine (F) has been investigated and found to be effective in enhancing the capacity of Li₂MnO₃ (figure 2.16) [61]. Dong *et al.* [16] reported that F doping weakened the Li-O bond and reduced the particle size of Li₂MnO₃ which shortens the pathway of Li⁺ diffusion (figure 2.16). Moreover, F substitution increased Mn³⁺ species and oxygen vacancies leading to higher capacities and cycle stability as observed in Table 2.4. EIS Nyquist plots also reveal that the charge transfer resistance has decreased in the doped sample.

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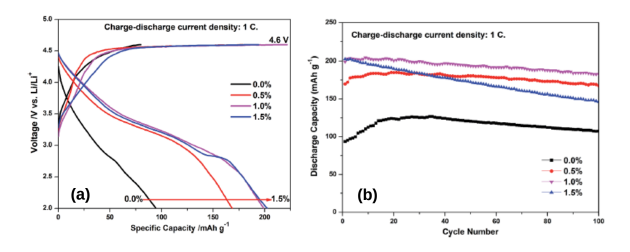


Figure 2.16: (a) Initial Charge/discharge curves of Li₂MnO_{3-x}F_x and (b)Cycling stability testing of Li₂MnO_{3-x}F_x [16].

Rare earth element doping

The use of rare earth elements such as Nd, Ce, Yb, Sc etc. to dope lithium ion cathode materials has been reported in several cathode materials, including LiCoO₂, LiMn₂O₄, LiFePO₄ and xLi₂MnO₃(1-x)LiMO₂ [68],[69],[70],[71]. As it stands, very few experimental reports have been published on rare earth element doping of pristine Li₂MnO₃ electrodes. From first principle studies, rare earth elements as dopants are favourable due to effects such as the ability to expand the lattice which helps to clear the pathway for diffusion thus facilitating ion transport [68],[69],[70],[71], [72], [73]. Furthermore, the expansion of the lattice slightly distorts the structure due to the large ionic radius which leads to smaller crystallites [72]. This offers the material a reduced charge transfer resistance and increased electric conductivity [68], [72].

A first principle study of the effects of rare earth doping was conducted by Zheng *et al.* [68], where they explored the effect of La, Ce, Pr and Sm on Li₂MnO₃. It was observed that La doping changed the properties of Li₂MnO₃ from semiconductor to metallic, while Ce, Pr and Sm doping retained the semiconductor nature [68]. In the case of the latter doped Li₂MnO₃, the band gap was significantly reduced compared to undoped samples which could be important in enhancing the electronic conductivity [68]. Similar results in improvement of electronic structure were observed by Yuzer *et al.* [74] when they experimentally doped Li₂MnO₃ with Nd, Ce and Yb. In their findings,

80 % Ce-doped Li₂MnO₃ showed a strong overlap of electronic levels that resulted in larger molecular bands [74].

It is worth noting that rare earth elements such as Pr, Sm, Nd, Ce, and Yb are favourable for doping as they possess redox couples i.e., RE³⁺/RE²⁺ or RE⁴⁺/RE³⁺ [72]. These redox couples can be electrochemically activated at appropriate potentials and can play a role as redox-active species thus increasing the energy density in cathode materials [72].

2.1.3.4.3.4. Other Modifications

Li₂MnO₃ dispersion

Li₂MnO₃ dispersion on rGO

Wei Zhao *et al.* [40] reported on the dispersion of pristine Li₂MnO₃ over reduced graphene oxide (rGO) to form a Li₂MnO₃/rGO composite as shown in Figure 2.17. Unlike merely coating the Li₂MnO₃ with rGO, their dispersion of Li₂MnO₃ particles over the rGO prevented agglomeration of Li₂MnO₃ particles thus improving the diffusion of lithium ions by shortening the path of diffusion [40]. The composite also showed improvement in the charge transfer resistance and exceptional discharge capacity in comparison with the pristine material [40]. The improved performance of the composite was attributed to the smaller particle size, reduced agglomeration, high surface area and electronic conductivity of rGO [40].

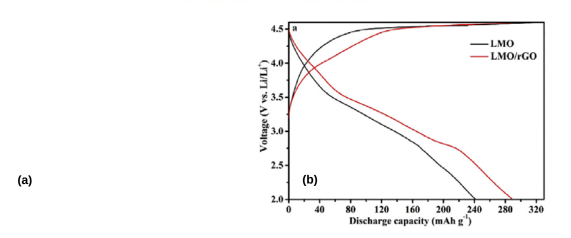


Figure 2.17: (a)Cycling stability testing of Li₂MnO₃ vs Li₂MnO₃/rGO (b) Initial charge/discharge curves of Li₂MnO₃ vs Li₂MnO₃/rGO [40].

Li₂MnO₃ dispersion on CNTs

In an effort to circumvent the agglomeration associated with nano structuring Li₂MnO₃ and improve its conductivity, Choi *et al.* [75] employed microwave assisted hydrothermal synthesis to form a composite of Li₂MnO₃ dispersed on Carbon Nanotubes (CNTs). The composite showed exceptional discharge capacities with electrochemical performance as observed in Table 2.5.

Table 2.5: Comparison of different dispersion substrates and their enhancements.

Substrate	Initial	Initial	Retained	Capacity	Measurement	Ref.		
	Discharge	Coulombic	Capacity	retention	Conditions			
	Capacity	Efficiency	(mAh.g ⁻¹)	(%)				
	(mAh.g ⁻¹)	(%)						
Reduced	289.4	96.7	250.62	86.6	2.0 and 4.6V, 0.1C,	[39]		
Graphene					45 cycles, RT			
Oxide								
Carbon	273	70.54	196.19	91.14	2.8 - 5 V, 0.1C, 10	[75]		
Nanotubes			(0.05C)	(0.05C)	cycles, RT			

2.1.3.4.4. Conclusion WESTERN CAPE

This review has introduced Li₂MnO₃ based materials as possible high energy and high-capacity lithium ion battery cathode materials for electric vehicle and grid storage technologies. Furthermore, it highlighted the challenges associated with such materials including low initial coulombic efficiencies and irreversible capacity loss during cycling. From reviewing all the studies mentioned in this dissertation, it becomes increasingly clear that the performance of Li₂MnO₃ is strongly dependant on its surface properties. It can also be noted that various modifications such as surface coating, ionic doping, nano-structuring, and dispersion on conductive surfaces have the capability of greatly improving the low initial coulombic efficiencies, voltage fade and low-rate capacity associated with Li₂MnO₃ materials.

Furthermore, it has been demonstrated that in some instances the modified Li₂MnO₃ can attain an initial discharge capacity as high as 335 mAh g⁻¹ which represents over 72 % of the theoretical capacity of Li₂MnO₃ (459 mAh g⁻¹) and 100% initial coulombic efficiencies. In other cases, Li₂MnO₃ was dispersed onto rGO and a capacity of >250 mAh g⁻¹was retained even after significant cycling representing >86% capacity retention. It is our belief that although each modification method has its merits and downfalls, Li₂MnO₃ materials would greatly benefit if these methods were exploited jointly. In conclusion, there is still a long way to go before the application of these materials become feasible in electric vehicle and grid storage technologies even with such exceptional results reported in this review. Firstly, we must better understand the origin of the electrochemical activity of such materials as this is still a highly debated matter. Secondly, the best optimum conditions must be established such as the synthesis methods and conditions, electrode thickness and coating/doping concentrations as they play a crucial role in determining the electrochemical performance.

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