Development of a new class of oxygen ion mixed conductors.

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Abstract

Dense ceramic membranes with mixed ionic and electronic conductivity can separate 100 % pure oxygen from air at elevated temperature. They constitute a much cheaper and more environmentally friendly alternative to cryogenic distillation. Today the challenge with dense ceramic membranes is the long term stability. The high operating temperatures (>800°C) of the membranes lead to chemical degradation and mechanical failure. Hexagonal manganites has the ability to reduce the problems associated with high temperatures. Compared to perovskites, which conduct through formation of oxygen vacancies, hexagonal manganites have the ability to form interstitial oxygen due to a less closed packed structure. In this thesis, it is reported on how the structure and oxygen storage ability changes with varying yttrium deficiency in hexagonal YMnO$_3$. It is important to investigate the possibility of cation vacancies in the structure, as little is known about the defect chemistry of these materials. Yttrium deficient samples of YMnO$_3$ made by solid state synthesis was fired in air, and annealed in varying atmospheres. Hexagonal YMnO$_3$ has the ability to lose 15-20 % yttrium without any notable changes in the structure and composition of yttrium deficient YMnO$_3$. Above this limit, reflections from Mn$_3$O$_4$ hausmannite appears in x-ray diffractograms. With a higher degree of yttrium deficiency in the hexagonal structure prepared in air, a slight contraction of the unit cell along both a and c direction is found. After annealing in O$_2$ atmosphere, the structure expands along the a axis and contracts along the c-axis. The opposite behavior is seen in inert (N$_2$) atmosphere, where the structure expands in c direction and contract along the a axis with increased yttrium deficiency in YMnO$_3$. Above 600 °C, during heating of YMnO$_3$ and Y$_{0.80}$MnO$_3$, oxygen goes out of the materials in both N$_2$ and O$_2$ atmospheres. Upon cooling in O$_2$ atmosphere, a reversible oxygen uptake is seen around 400 °C for both YMnO$_3$ and yttrium deficient YMnO$_3$. For stoichiometric YMnO$_3$ the reversible oxygen adsorption is suggested to be attributed to interstitial oxygen, while for stoichiometric yttrium deficient YMnO$_3$ it is believed to be filling of oxygen vacancies. The reversible oxygen uptake of YMnO$_3$ was calculated to be $\delta = 0.034$. Moreover, in this thesis it is also reported on the synthesis of an asymmetric membrane consisting of 15 % titanium substituted YMnO$_3$. The asymmetric membrane did not yield a dense functional layer of nanocrystalline YMn$_{0.85}$Ti$_{0.15}$O$_3$ on top of the porous support after sintering.
Sammendrag

Keramiske membraner for oksygen separasjon basert på en blanding av ionisk og elektronisk ledende oksid, er billigere og mer miljøvennlig enn kryogenisk destillasjon for produksjon av 100 % rent oksygen. En utfordring som er knyttet til dagens keramiske membraner er kjemisk degradering og redusert leveid tid som følge av de høye arbeidstemperaturene over 800 °C. I forhold til perovskitter, som brukes mye som materialet i keramiske membraner i dag, har heksagonale manganitter muligheter for å romme interstitielt oksygen og dermed være ionisk ledende ved mye lavere temperaturer. I denne oppgaven rapporteres det om hvordan strukturen og muligheten for oksygen lagring endres med yttrium underskudd i heksagonal YMnO₃. Det er lite man vet om defekt kjemien i heksagonale manganitter, og det er derfor her forsket på kation vakanser i YMnO₃ for å finne ut mer om dette. Pulver med varierende yttrium underskudd i YMnO₃ har blitt laget med fast stoff syntese, brent i luft, og videre varmebehandlet i forskjellige atmosfærer. Heksagonal YMnO₃ har muligheten for å miste 15-20 % yttrium uten noen spesielle endringer i strukturen. Over denne grensen, kommer refleksjoner av en Mn₃O₄ fase opp i røntgen diffraktogrammene med økende yttrium underskudd i YMnO₃. En liten kontraksjon av strukturen både langs a og c aksen for yttrium underskudd prøver brent i luft. I oksiderende atmosfære (O₂) blir strukturen ekspandert langs a aksen, men kontraherer langs c-aksen, og i inert atmosfære (N₂) skjer den motsatte med økende yttrium underskudd i YMnO₃ ved at c aksen ekspanderer mens a aksen kontraherer. Ved oppvarming av heksagonal YMnO₃ og Y₀.80MnO₃ i både oksiderende og inert atmosfære går oksygen ut av materialet ved omtrent 600 °C. Ved avkjøling i oksiderende atmosfære er det et reversibelt oksygen opptak rundt 400 °C for både YMnO₃ og Y₀.80MnO₃. Denne reversible adsorpsjonen av oksygen er trodd til å ha en sammenheng med yttrium vakanser i Y₀.80MnO₃ og interstitielt oksygen i YMnO₃. Oksygen opptaket under avkjøling er beregnet til å være δ= 0.034 for YMnO₃ i oksiderende atmosfære. I tillegg har det i denne oppgaven blitt laget en asymmetrisk membran av 15 % titan dopet heksagonal YMnO₃. Et tett funksjonelt lag bestående av sintrede nano-partikler av YMn₀.₈₅Ti₀.₁₅O₃ ble ikke oppnådd på en porøs bærer.
<table>
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<th>Abbreviation</th>
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| XRD          | X-Ray Diffra
c                            |
| SEM          | Scanning Electron Microscope       |
| SE           | Secondary Electrons                |
| BSE          | Back Scatter Electrons              |
| MIEC         | Mixed Ionic and Electronic Conduc
tor                                    |
| TGA          | ThermoGravimetric Analysis         |
| HS           | High Spin                           |
| HTXRD        | High Temperature X-ray Diffraction |
| CN           | Coordination Number                 |
| TEM          | Transmission Electron Microscope    |
| VESTA        | Visualizing for Electron and Structural Analysis (software) |
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1. Introduction

Oil is the most valuable commodity in world trade today. Currently, almost 1000 barrels oil a second are used every day, but the proven reserves for fossil fuels are progressively decreasing[1]. The continuing use of fossil fuels produces harmful effects such as pollution and greenhouse gases that threatens the human health and give global warming. The steadily increasing energy needs in nowadays society forces continuous research for alternative environmentally friendly energy production techniques. One approach to meet these global demands is the continuously research on more environmentally production of syn-gas to the reforming of natural gas to liquid fuels. Both the common Fisher-Tropsch- and Methanol synthesis for reforming natural gas requires synthesis gas (H$_2$ +CO) as feedstock [2, 3]. The modern approach for production of syn-gas is the slightly exothermic partial oxidation of methane [4]. In the following schemes (1.1) and (1.2), both steam reforming and partial oxidation are illustrated.

\[
CH_4 + H_2O \leftrightarrow CO + 3H_2 \quad (1.1)
\]

\[
CH_4 + \frac{1}{2}O_2 \leftrightarrow CO + 2H_2 \quad (1.2)
\]

The partial oxidation of methane requires 100 % pure oxygen to improve its performance due to the absence of nitrogen. Conventionally, oxygen separation to produce pure oxygen is a big business with nearly 100 million tons of oxygen produced every year [5]. The currently most established technique for large- scale oxygen separation is carried out by the cryogenic process [6]. This process is very expensive and energy costing, due to requirements such as low temperature and elevated pressure. An alternative is to use mixed ionic and electronic conducting (MIEC) membranes in combination with a reforming catalyst. Because MIEC membranes are 100 % selective for oxygen, the syn-gas production is reduced to just one single step. MIEC materials offer to separate oxygen from air electrochemically, and in addition to the application as oxygen permeable membranes, they are important applications e.g. cathode materials for solid oxide fuel cells and oxygen sensors [2, 3, 7].

MIEC membranes are also important in carbon capture and storage (CSS) especially in cooperation with coal power plants. [8]. “Carbon capture and storage could deliver up to 55% of the mitigation effort needed for the stabilization of atmospheric CO$_2$ levels by 2100”, Rachel Spratt, Environmental Resource Management [9]. Coal is the most abundant and dirtiest fossil fuel, and different fossil fuel power plant concepts for CSS are currently under development, such as oxy fuel power plants. In this process the fossil fuel is combusted using pure oxygen and the result is a gas steam consisting of almost 90-95 % pure CO$_2$ in the dried flue gas. In the combustion process, the capturing of CO$_2$ can be done more easily compared than when air is
used, and stored in a safe geological site. Energy penalties can be reduced by 35 % or more by integrating dense ceramic membranes in coal power stations [6].

1.1 MIEC membranes

The two main types of oxygen separation systems based on dense ceramic membranes are (i) pure oxygen conducting membranes and (ii) mixed ionic-electronic conducting membranes (MIEC) [10]. The main difference between the two types of membranes is the driving mechanism. When oxygen diffuse through the membrane, an opposite but simultaneous flux of electrons have to charge compensate for the oxygen flux. In pure conducting materials, these electrons have to go in an external circuit, and the oxygen yield can be found from the applied current. In MIEC membranes, a gradient in oxygen partial pressure ($\Delta p_{O2}$) is the driving force for oxygen permeation. Figure 1.1, gives a steady state profile of the chemical potential gradient across a thin MIEC membrane on a porous substrate.

**Fig 1.1:** A schematic model of an asymmetric mixed ionic and electronic conducting membrane (MIEC).

Oxygen permeation through a dense MIEC membrane involves three different steps: (i) oxygen adsorption and dissociation into oxide ions, at the high partial pressure of oxygen side of the membrane, (ii) the simultaneous bulk diffusion of holes/electrons and charged oxygen (iii) release of oxide ions into molecular oxygen. The slowest moving specie or slowest process is expected to limit the rate of oxygen permeation. When the flux is governed by bulk diffusion it can generally be described with classic Wagner theory [3, 10, 11]

$$J_{O_2} = -\frac{RT}{4\pi^2p^2} \times \frac{\sigma_{\text{ion}}\sigma_{\text{el}}}{\sigma_{\text{ion}}+\sigma_{\text{el}}} \times \nabla \mu_{O_2}$$ (1.3)
Where $J_{O_2}$ is the oxygen permeation flux in mol m$^{-2}$ s$^{-1}$, $R$ is the gas constant, $F$ is the Faraday constant and $\sigma_{\text{ion}}$ and $\sigma_{\text{el}}$ represents the partial conductivities of oxygen ionic and electronic defects. $L$ represents the membrane thickness and $\nabla \mu_{O_2}$ is the oxygen chemical potential gradient. Moreover, a substitution of the chemical potential with oxygen partial pressure ($p_{O_2}$), and an integration over the membrane thickness $L$ between oxygen partial pressure on the air side ($P_1$) and on the fuel side ($P_2$) gives the integrated Wagner equation for bulk diffusion \[2\],

$$J_{O_2} = -\frac{RT}{4L^2} \times \frac{\sigma_{\text{ion}}\sigma_{\text{el}}}{\sigma_{\text{ion}} + \sigma_{\text{el}}} \times (\ln P_1 - \ln P_2) \tag{1.4}$$

Consequently, reducing membrane thickness for bulk controlled diffusion will increase the $O_2$ permeation rate \[12\]. For thinner membranes, the resistance to oxygen ion conduction will be decreased until a certain limit, known as the critical thickness ($L_c$), which also depends on operating conditions. There are few reports concerning thin films of hexagonal YMnO$_3$, while for the perovskite structure, a critical film thickness should be 25 $\mu$m or less\[12\]. For thin membranes, a porous support is needed for mechanical stability also referred to as an asymmetric membrane. For a thickness below $L_c$, the interphase of the membrane becomes the region of higher resistance, shown in Figure 1.2a. Wagner equation is not applicable when the oxygen becomes determined by the surface exchange reactions. Decreasing partial pressure of oxygen ($P_{O_2}$) gives increased control of oxygen flux by surface kinetics, Figure 2.1b.

**Fig. 1.2:** Oxygen flux through membrane as a function of the reciprocal thickness of the membrane $1/L$. a) As the membrane becomes thinner, the oxygen flux is increasingly dominated by surface exchange kinetics, b) Decreasing $P_{O_2}$ leads to increased control of oxygen transport by surface kinetics. $1/L_c$ is the reciprocal characteristic thickness where surface exchange is dominating.
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The critical thickness can be expressed by the surface exchange coefficient \( k_{\text{chem}} \) and the chemical diffusion coefficient \( D_{\text{chem}} \) as follows [2, 13],

\[
L_C = \frac{D_{\text{chem}}}{k_{\text{chem}}}
\]  

(1.5)

For asymmetric membranes with a thickness value less than \( L_C \), the surface exchange coefficient \( k_{\text{chem}} \) is the limiting factor for oxygen permeation. Thus no further increase in oxygen flux is feasible without increasing \( k_{\text{chem}} \).

1.1.1 Material requirements

A major challenge in processing supporting MIEC membranes is the selection of a suitable fabrication method and materials for the porous substrate, taking into account requirements as chemical compatibility, similar thermal expansion coefficient, good mechanical strength and porosity for adequate diffusion of gas [8, 12]. In addition, stability in reducing atmospheres, membrane design, sealing and costs are important factors in processing MIEC membranes.

Non-stoichiometric oxide materials have been heavily researched because of their reversible oxygen storage/release capacities (OSC), and the most used dense ceramic membranes today are perovskite or fluorite structures [5, 10, 14]. Several perovskite materials have demonstrated high OSC values (>2500 µm·O/g) and low re-oxygenation temperatures (< 200 °C) [14]. However, these materials require either reducing atmosphere or high temperatures (> 700 °C) to form oxygen vacancies. The challenge for MIEC membranes today are the long-term chemical and mechanical stability associated with the high operating temperatures. The chemical driving force for oxygen permeation can cause chemical degradation in the material, and further on, a difference in mobility of the cations in the structure can lead to decomposition [15]. Lot of research are focusing on trying to increase the oxygen flux and to reduce the operating temperatures for the MIEC materials to below 700 °C, since lower operating temperatures reduce energy consumption and thermal degradation of the membrane [10]. A very recent work from Remsen et.al [14] showed that hexagonal DyMnO\(_{3+δ}\) can accommodate a large excess of oxygen up to \( δ=0.35 \). In Figure 1.3, the reversible absorption of large amounts of oxygen around 200-300 °C and sharply desorbed oxygen at elevated temperatures is shown for DyMnO\(_{3+δ}\). Hence, it is suggested that hexagonal manganites have the ability to accommodate oxygen interstitials and thus have ionic conduction at lower temperatures. This is promising for the next generation of MIEC membranes because it may solve the problems with long-term stability due to the high operating temperature used today.
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Fig. 1.3: Oxygen content as a function of temperature for Dy1-xYxMnO3+δ with a heating/cooling rate of 0.1 °C/h. Figure is reprinted from Remsen et.al [14].

As the name “MIEC membrane” states, both ionic and electronic conductivity are important material requirements. Ionic conductivity often relates to defects in the crystal structure, while electronic conductivity often is associated with electronic defects [10]. A defect is an imperfection in the crystal structure that differentiate a perfect crystal (only at 0 K) from a crystal. The concentration of defects in crystalline solids are important because they can modify important properties [16].

The perovskite structure of LaMnO3 is known to have a degree of freedom in cation stoichiometry [17]. It is therefore natural to investigate the same phenomenon in hexagonal manganites. YMnO3 is one of the most studied hexagonal manganite [18], and due to the layered structure of hexagonal manganites, an decrease in yttrium content will be interesting. Since yttrium oxide is an expensive oxide, it will be beneficial to investigate the ability of decreasing the yttrium content in the structure.

1.2 Research objectives

This thesis has two primary goals, the first one will be to examine how yttrium deficiency impacts the hexagonal structure of YMnO3. It will be investigated how the lattice parameters change from yttrium deficient samples prepared in air atmosphere in addition to be annealed in O2 and N2 rich atmospheres. This is to get a better understanding of the defect chemistry in YMnO3, since hexagonal manganites have the possibility to accommodate interstitial oxygen. The second goal is to investigate the possibility of using hexagonal YMn0.85 Ti0.15 O3 as an oxygen permeable membrane. Titanium substitution was chosen due to its ability to reduce microcracking in YMnO3, as titanium has proven to reduce the anisotropic thermal expansion and porosity in YMnO3. However, the origin of microcracking will not be investigated in this
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thesis. The aim is to fabricate an asymmetric membrane consisting of an active layer on top of a porous support. The same composition will be used for both the active layer and the support to avoid capability problems, which can lead to degradation or cracking. It is important to understand the conditions necessary for a successful preparation of the membrane, as it has never been done before for hexagonal YMnO$_3$. Known synthesis methods for YMnO$_3$ are going to be used in addition to a successful technique used in production of perovskite asymmetric membranes.
2. Literature survey on YMnO$_3$

Rear-earth manganites with their competing orthorhombic and hexagonal crystal structures, with 6- and 5 folded coordinated Mn$^{3+}$, have been studied for over 50 years [14]. The hexagonal manganites has attracted much interest, especially in recent times, because of the coexistence of ferroelectricity and antiferromagnetic order [19]. However, most multiferroic manganites are insensitive to applied magnetic fields due to their antiferromagnetic order. This is a disadvantage in potential applications.

Among hexagonal manganites, YMnO$_3$ is the most studied metal oxide[20]. Bulk YMnO$_3$ has a high ferroelectric transition temperature $T_c \sim 913$K [21] and a low antiferromagnetic transition temperature $T_N \sim 70$-80K [22, 23]. One can note that literature shows a large spread of $T_N$. Increased attention should be given to off-stoichiometry effects for the understanding of the physical properties of hexagonal YMnO$_3$. Unlike perovskite thin films, YMnO$_3$ does not possess a critical film thickness and is therefore a promising candidate for making multiferroic devices in nano scale. Thin films of YMnO$_3$ have also potential in non-volatile ferroelectric random access memory devices.

2.1 Crystal structure

The isostructural RMnO$_3$, where R is a transition or a rare earth metal, can crystallize in either hexagonal or orthorhombic structures. Large rare-earth elements like Dy, Tb and Gd prefers the orthorhombic structure with room temperature space group $Pbnm$ [17]. For smaller ions like Y, Sc, Ho-Lu and In, the hexagonal structure with space group $P6_3cm$ (No. 185) is the preferred ground-state structure.

Hexagonal YMnO$_3$ with space group $P6_3cm$ consist of Mn$^{3+}$ ions coordinated by two apical and three planar oxygen atoms with corner sharing atoms in the ab-plane. This forms a trigonal bipyramidal environment around the Mn ions. Each consecutive layer of MnO$_5$ is rotated along the c-axis by 180° [10, 11]. The bipyramids are tilted with respect to the ab-plane by $\sim 8°$ and to the polar c-axis by $\sim 5°$ [17]. Situated between the MnO$_5$ layers lies Y$^{3+}$ coordinated by 7 oxygen ions. The displacement of Y atoms along the c-axis for the low temperature phase are 2/3 in negative direction and 1/3 in the positive direction[14]. The small displacement of the trigonal bipyramids compared to the Y$^{3+}$ layer gives rise to a spontaneous polarization of $\sim 5$-6μm/cm$^2$ [24]. The main driving force for polarization is electrostatics although partial covalency is found between empty Y 4d orbitals and O 2p orbitals[24, 25]. Hexagonal YMnO$_3$ are therefore referred to as a “geometric ferroelectrics”.

Above the transition temperature (trimerisation temperature), YMnO$_3$ exhibits a paraelectric structure with space group $P6_3/mmc$ (No. 194) [11]. This is a less ordered structure with higher symmetry than the low temperature phase. In this phase compared to the polar $P6_3cm$ are the MnO$_5$ bipyramids not tilted and the Y$^{3+}$ ions are not displaced. The unit cell of the two different space groups are shown in Figure 2.1, and the space group $P6_3/mmc$ is presented with a $P6_3cm$. 


basis to allow comparison. Green atoms display $Y^{3+}$, red atoms $O^{2-}$ and purple polyhedra show the trigonal bipyramids $\text{MnO}_5$.

![Diagram](image)

**Fig. 2.1:** a) The low temperature space group $P6_3cm$. b) The high temperature phase with space group $P6_3cm$ basis c) The high temperature phase $P6_3/mmc$. Green atoms display $Y^{3+}$, red atoms display $O^{2-}$, the purple trigonal bipyramids indicate $\text{MnO}_5$ with black $\text{Mn}^{3+}$ in the middle. Figure is drawn using VESTA software [26] and atomic positions are from Gibbs et al. [27]

The manganese ions in space group $P6_3cm$ are displaced both in the basal plane and in the c-direction. The difference in bond length between Mn-O in-plane and out-of-plane is reported to be 2.05 Å and 1.86 Å respectively [28]. Onset of the bipyramidal tilting at the trimerisation temperature triples the unit cell volume from space group $P6_3/mmc$ to $P6_3cm$ and is given by the following [4]:

$$\frac{1}{3}a(P6_3/mmc) = a(P6_3cm)$$ (2.1)

The temperature of onset for trimerisation of $\text{MnO}_5$ and polar displacement of $Y^{3+}$, as well as the driving force for polarization have been discussed in several reports. Since the phase transition between the two polymorphs is a second order phase transition, it has been argued from group theory and high temperature x-ray diffraction (HTXRD) that hexagonal manganites must undergo two structural transitions, and that there exist an intermediate phase with space group $P6_3/mcm$ [24, 29]. Recent experiments do not support the notation of an intermediate phase. The ferroelectric to paraelectric phase transition have been reported to be around 1250 K [27], but some structural anomalies have been reported around the Curie temperature, but this is not well understood [24, 29]. Selbach et al [24] found a direct and continuously transition from $P6_3cm$ to $P6_3/mmc$ obtained by HTXRD in both $N_2$ and air atmosphere, with no indication of intermediate phase. In a study performed by Bergum et al [17], a second order transition in $\text{YMnO}_3$ at 1020 °C was found by HTXRD in oxygen atmosphere and by dilatometry. There were no indications of contraction or expansion of $\text{YMnO}_3$ between $N_2$ and $O_2$ atmospheres.
Chapter 2: Literature survey on YMnO$_3$

2.2 Phase stability

The orthorhombic and hexagonal polymorphs are close in energy. The binary oxides forming YMnO$_3$ is given by the following:

$$2Y_2O_3(s) + 2Mn_2O_3(s) \rightarrow YMnO_3(s)$$  \hspace{1cm} (2.2)

Since reactions, involving only solids expects to have an entropy close to zero, the enthalpy of the reaction dominates the Gibbs energy. Yokokawa et.al [30] were the first to report the empirical relationship between the Goldschmidt's tolerance factor $t$ and the enthalpy of formation of perovskite from their constituent binary oxides. This relationship has also been shown to be valid for manganites[17], and thus the relative stability between the hexagonal and the orthorhombic structures depends on the Goldschmidt's tolerance factor [31],

$$t = \frac{r_a+r_o}{\sqrt{2(r_b+r_o)}}$$  \hspace{1cm} (2.3)

As seen from (2.3), the tolerance factor is dependent on the radius of each component in an ABO$_3$ structure. The hexagonal phase is stable for a tolerance factor of $t<0.855$, while the perovskite phase is stabilized for a tolerance factor in the range $0.855 \leq t \leq 1$ (Calculated using Shannon’s values at room temperature[32]) [33]. The calculated Goldschmidt's tolerance factor for YMnO$_3$ is 0.854, which is close to the geometric stability limit of the perovskite structure. The hexagonal structure is stabilized for YMnO$_3$ due to the small yttrium ions that wants to lie in layers instead of in between octahedral. The lower the tolerance factor, the less exothermic the enthalpy of reaction (2.2) and for hexagonal YMnO$_3$ it is reported to be $-29.8\pm2.1$ kJmol$^{-1}$. This value is about 1/3 of the corresponding value for orthorhombic LaMnO$_3$. [17]

According to a study on the thermodynamics of the Y-Mn-O system[34] in air, seen in Figure 2.2, hexagonal YMnO$_3$ is metastable for temperatures less than 789 °C and entropy stabilized with respect to Y$_2$O$_3$ and YMn$_2$O$_5$ above this temperature. However, the hexagonal structure is considered stable at room temperature because the transition from orthorhombic to hexagonal phase is reconstructive, and therefore expected to be slow with a high activation energy barrier. The metastability can be extended to larger temperature intervals [24, 34]. Above 1794 °C, the hexagonal phase melts incongruently to $\alpha$-Y$_2$O$_3$(s) and liquid.
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The metastable and denser orthorhombic perovskite forms easily at high pressures or through epitaxial stabilization. Oxidizing conditions and low temperature chemical synthesis routes also favour the perovskite structure. Oppositely can the hexagonal phase be stabilized with epitaxial growth, single crystal growth or by reducing conditions[24]. Since the phase formations are affected by oxidizing/reduction conditions it is suggested by Selbach et.al[24] that oxidizing/reduction of a fraction of Mn$^{3+}$ to Mn$^{4+}$/Mn$^{2+}$ readily can shift the stability of the two phases. This is also suggested by Bergum et.al[17], since both the enthalpy and entropy of reaction (2.2) can be argued to be small. A possible oxidation of Mn$^{3+}$ to Mn$^{4+}$ may have a large effect of the polymorph stability since there exists a substantial enthalpy involved in the oxidation of Mn$^{3+}$ to Mn$^{4+}$. As Mn$^{4+}$ is smaller than Mn$^{3+}$, the oxidation of a fraction of Mn$^{3+}$ will increase the average Goldschmidt’s tolerance factor of the system. The Shannon radius for the high spin cations Mn$^{3+}$ and Mn$^{4+}$ are 0.645Å and 0.53Å respectively [32]. An illustration showing the different radii of manganese with different valence states is shown in Figure 2.3.

Fig. 2.2: Phase diagram for the Y-Mn-O system. Figure is reprinted from Chen et.al [34]

Fig. 2.3: Shannon ionic radius for different valence states of Mn (HS). Values are obtained from [32]
The orthorhombic perovskite LaMnO$_3$ has been found to display a degree of freedom in oxygen stoichiometry in the range 2.947-3.079 at 1200 °C, while the corresponding range for hexagonal YMnO$_3$ was found to be 2.963-3.009[17]. As the perovskite structure can accommodate a higher degree of excess oxygen due to partial oxidation of Mn$^{3+}$ to Mn$^{4+}$, it is therefore expected that oxidizing conditions favour the formation of the orthorhombic perovskite phase. Oxidation of Mn$^{3+}$ is known to influence the phase stability and crystal structure of the multiferroic perovskites BiFe$_{1-x}$Mn$_x$O$_{3+\delta}$ and BiMnO$_{3+\delta}$.

2.2.1 Crystal field stabilization

A contribution to stabilize the hexagonal structure of YMnO$_3$ is the trigonal bipyramidal crystal field stabilization energy associated with the four electrons in the d-orbital of Mn$^{3+}$ [35]. The trigonal bipyramidal and octahedral crystal field splitting for hence hexagonal and orthorhombic phases is represented in Figure 2.4. The red circles in the figure represents the d$^4$ electron of Mn$^{3+}$.

![Fig. 2.4: a) Octahedral crystal field splitting, b) trigonal bipyramidal crystal field splitting. Marked arrows represents the d$^4$ electron of Mn$^{3+}$.](image)

The lowest excitation limit for a d$^4$ electron in the trigonal bipyramidal crystal field splitting is up to the fairly localized z$^2$ orbital. This energy difference depends sensitively on the Mn-O distance within space group P6$_3$cm [28].

2.3 Thermal and chemical expansion of the structure in varying atmospheres

The lattice parameters a and c of hexagonal YMnO$_3$ experiences anisotropic chemical and thermal expansion upon heating [24, 27, 36]. Thermal expansion is a linear trend, and is a
volume change that the material experience upon a temperature change. The additional expansion, when there does not exist a linear trend is called a chemical expansion. The difference between chemical and thermal expansion can be seen in Figure 2.5.

![Graph showing thermal and chemical expansion](image)

**Fig. 2.5:** The contribution of chemical expansion to the thermal expansion. The onset of thermal reduction is $T_0$. Figure is reprinted from H. Lein[12]

Selbach et al [24] found the anisotropic relationship in air, nitrogen and oxygen atmosphere for hexagonal YMnO$_3$ and HoMnO$_3$ and is seen in Figure 2.6. The lattice parameter c for HoMnO$_3$ decreases non-linearly to the trimerisation temperature ($T_s$) around 1300 K, where it has close to zero thermal expansion for about 100 K before it increases with temperature. The increase in unit cell parameter c is more rapid above $T_s$ in N$_2$ compared to air atmosphere. The same behavior is observed for YMnO$_3$, but it is not seen in Figure 2.5 as the data is collected in the temperature range below 1400 K. However, the unit cell parameter a is smaller in air compared to N$_2$ atmosphere, and the difference increases with increasing temperature. The changes in slopes at temperatures 1200 K and 1300 K for the lattice parameter a in HoMnO$_3$, have also been reported by Nenert et al [37] for YMnO$_3$. 

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Fig. 2.6: a) Changes in unit cell parameter c, (b) lattice parameter a and (c) unit cell volume for hexagonal YMnO\(_3\) HoMnO\(_3\) in O\(_2\), N\(_2\) and air atmosphere. Figure is reprinted from Selbach et.al[24].

The increase of the unit cell parameter a with temperature, is assigned to thermal expansion in the material, while the decrease in unit cell parameter c is related to ferroelectric distortion as the alignment of MnO\(_5\) bipyramids causes Y\(^{3+}\) to contract[15]. From about 650 K, the unit cell volume is larger for N\(_2\) than air. The anisotropic chemical expansion for both YMnO\(_3\) and HoMnO\(_3\) is an indication that the oxygen content is sensitive to partial pressure of oxygen. The symmetry and cell dimensions in hexagonal YMnO\(_3\) depends on oxygen content in the structure and its effect on the crystal structure has received little attention. The chemical expansion seen at higher temperatures is attributed to oxygen vacancies in the material.

2.3.1 Microcracking as a result of anisotropic expansion

Application of polycrystalline YMnO\(_3\) is limited by porosity and grain boundary microcracking, which affects the mechanical and physical properties of the material. Tomczyk et.al [36] were the first to address the microcracking observed in YMnO\(_3\). A volume change of 3.7 %, which generated an internal stress of 200 MPa, and a high anisotropic thermal expansion coefficient in the temperature range 600-1000 was found to be the origin of microcracking. The high temperature phase transition from \(P6_3/mmc\) to \(P6_3cm\) which is reflected through the differences in a and c lattice parameters during heating, is associated with the high volume change. It is known that phase transitions that involves ionic rearrangement often is associated with a volume change. In addition can thermal expansion in highly crystalline structures give
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spontaneous microcracking upon cooling[38]. Besides decreasing the grain size below a critical value, suitable doping can give a reduction of both the volumetric change and the anisotropic thermal expansion. Ti substitution, recently shown by Tomcyk et.al[36], have an effective way to reduce the microcracking in YMnO$_3$.

2.3.2 Substitution with titanium in YMnO$_3$

As mention above, increased concentration of Ti in YMnO$_3$, decreases the concentration of microcracks and intergranular porosity[39]. This is because increased Ti substitution has shown to increase the unit cell parameter a, and decrease the unit cell parameter c, up to x= 0.15 [15, 39]. This gives a decreased c/a – ratio for the hexagonal unit cell which in turn reduces the anisotropic thermal expansion in the high temperature phase transition from $P6_3/mmc$ to $P6_3cm$. With X-ray diffraction this phase transition can be seen especially by the disappearance of the reflections at 23° and 36 ° 2θ for x=0.15. The experimental unit cell parameters for Ti substitution in YMnO$_3$ for x = 0.05, 0.10 and 0.15 refined within space group $P6_3cm$ was found by S.Nesdal [15] and are found in Appendix A. Increased titanium substitution (up to x=0.15) in YMnO$_3$ have also in recent preliminary master’s thesis [40] shown a high conductivity compared pure YMnO$_3$. At 400 °C, the conductivity of YMn$_{0.85}$Ti$_{0.15}$O$_3$ was $2\times10^{-3}$ S/cm compared to ranges of $10^{-6}$ S/cm for YMnO$_3$. For increased Ti content above x=0.175, a rhombohedral phase with space group R3c or R3c (bar) is observed by Asaka et.al[41].

2.4 Oxygen non-stoichiometry and defect chemistry of YMnO$_3$

2.4.1 Point defects

The simplest defect imaginable in a crystal is an imperfection at a single atom site. These defects are called point defects and can be divided into intrinsic and extrinsic point defects. Intrinsic point defects involve addition or removal of an atom in the structure, while the extrinsic point defects are defects that result by doping with an impurity atom. The two types of point defects that can occur in a pure material are; (i) vacant atom sites, or vacancies, or (ii) interstitial atoms occupying lattice sites which are normally not occupied in the crystal. A representation of an interstitial atom and a vacancy is seen in Figure 2.7.

![Diagram of interstitial atom and vacancy](image)

**Fig. 2.7:** Point defects in a perfect periodic structure.
Such vacancies and interstitials are always present even in the purest material. For these defects to be stable, the free Gibbs energy ($\Delta G$) of a crystal containing defects must be less than the free Gibbs energy of a crystal not containing defects[16].

$$\Delta G_{\text{crystal with defects}} < \Delta G_{\text{pure crystal}}$$ (2.4)

The concentration of defects will be in a thermodynamic equilibrium at a given temperature. This equilibrium concentration corresponds to the minimum in Gibbs free energy, which is dependent on the enthalpy of defect formation ($\Delta H$) and entropy of configuration ($\Delta S$). Defects can be induced by exposing the material for a certain environment, for example a low partial pressure of oxygen to favor oxygen vacancies or oxygen rich atmosphere to induce interstitial oxygen.

The presence of oxygen vacancies in hexagonal YMnO$_3$ is controversial. It has been reported that oxygen vacancy defects exist[42], whereas the presence of oxygen vacancies within hexagonal YMnO$_3$ was claimed unlikely[37]. Cation diffusion is anyway not pronounced in hexagonal manganites up to a temperature of 1000K, which suggests creation of interstitial oxygen at lower temperatures without formation of oxygen vacancies. The hexagonal phase of YMnO$_3$ is also less dense compared to the known orthorhombic structure, which may give more sites for interstitial oxygen in the structure. Overton.et.al[43] suggests that oxygen vacancies will preferably leave planar site in the MnO$_5$ bipyramids. A chemical expansion for both unit cell parameters $a$ and $c$ was found with decreasing oxygen in the structure of YMnO$_3$. The defect reaction for an oxygen vacancy in YMnO$_{3-\delta}$ is given by the following:

$$YMnO_3 \rightarrow YMnO_{3-\delta} + \frac{\delta}{2}O_2$$ (2.5)

Written in Kröger-Vink notation this becomes

$$2\delta Mn^\ddagger_Mn + O^\circ_O \rightarrow 2\delta Mn^\prime_{Mn} + V^\circ_O + \frac{\delta}{2}O_2$$ (2.6)

$$Mn^\prime_{Mn} \rightarrow e^- + Mn^\ddagger_Mn$$ (2.7)

It is suggested that one oxygen vacancy will affect two Mn atoms. The charge compensated electrons are assumed to be localized on Mn because Mn has several possible oxidation states. The oxidation state for Mn will be reduced from 3+ to 2+ and Mn change from 3d$^4$ to 3d$^5$. 

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Chapter 2: Literature survey on YMnO$_3$

With the same explanation as for oxygen vacancies, it is suggested for interstitial oxygen in YMnO$_{3+\delta}$ that the charge compensated holes are localized around Mn. The oxidation state of the two Mn atoms will be oxidized from 3+ to 4+ and Mn change from 3d$^4$ to 3d$^3$. The defect reaction of interstitial oxygen in YMnO$_{3+\delta}$ is given by the following:

$$\frac{\delta}{2}O_2 + Y MnO_3 \rightarrow Y MnO_{3+\delta} \quad (2.8)$$

In Kröger-Vink notation this becomes

$$\frac{\delta}{2}O_2 + 2\delta Mn^\delta_{Mn} \rightarrow \delta O'^{''}_{i} + 2\delta Mn^\cdot_{Mn} \quad (2.9)$$

$$Mn^\cdot_{Mn} \rightarrow h^\cdot + Mn^\delta_{Mn} \quad (2.10)$$

2.4.2 Electronic defects

The electronic defects in a material are associated with excitation of electrons from valence band (VB) to the conduction band (CB), intrinsic ionization or formation to keep charge balance in the material [10]. In non-stoichiometric oxides, the concentration of electronic defects usually relates with oxygen excess or oxygen deficiency. Thus for perovskites, metal excess or oxygen vacancies are accompanied by formation of electrons, which acts as the major charge carriers in the material and is responsible for the n-type electronic conductivity. In the opposite manner, metal deficiency or oxygen interstitial creates electron holes, which results in a p-type semiconductor, where the electronic conductivity occurs by transport of electron holes in the valence band. To achieve higher conductivity, doping with an aliovalent cation can cause oxygen excess or oxygen deficiency.

YMnO$_3$ is a poor p-type semiconducting material[44] with a believed mechanism for conduction found to be thermally activated hopping of small polarons between Mn$^{3+}$ and Mn$^{4+}$, and migration of oxygen vacancies at higher temperatures [45]. The overlap of Mn orbitals in the basal plane is therefore suggested to be the conductive state for hexagonal manganites [24]. A polaron is a quasiparticle used to describe the interaction between an electron and atom in solid materials[46]. This quasiparticle can be described as an electron with a polarization cloud. As the electron moves around, it will drag the polarization cloud with it, which results in a higher inertial mass and lower mobility compared to a free electron.

The Nernst-Einstein equation are used to relate the activation energy of polaron hopping and the temperature dependence of the mobility for an electron or hole.
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\[
\mu = Ae^{-\frac{E_a}{RT}}
\]  

(2.11)

Where $E_a$ is the activation energy for polaron hopping, $R$ is the gas constant, $A$ is a pre-exponential factor and $T$ is the absolute temperature. The latter relation also holds for ionic conductivity, because the process of polaron hopping is similar to ion hopping between neighbouring sites. Thus, the mobility of an ion increase with increasing temperature. In many cases, the electronic conduction dominates the total conductivity in a material, because the mobility of electrons is about 1000 times higher compared to the mobility of ions in the lattice [15].

2.5 Theory on synthesis methods

2.5.1 Sol gel synthesis

Synthesis of nanocrystalline hexagonal YMnO$_3$ used in this project was adapted from Bergum et al [17], which is a modified Pechini method. This method has successfully synthesized nano-sized crystalline powder of YMnO$_3$. Nitric acid was avoided due to increased stabilization of the metastable orthorhombic phase. Ethylene glycol in equal amounts to the citric acid has shown to have a significant effect on suspending the cations in the solution in addition to helped polymerize and to avoid precipitation. There was shown to be a clear trend in increasing crystallite size with increasing annealing temperature and above approximately 750°C, the crystallite size was dependent on kinetics. An annealing temperature of 825°C of the raw powder gave a crystallite size of 32 nm, while at 900°C a crystallite size of 52 nm was obtained.

2.5.2 Solid state synthesis

Solid state synthesis is a widely used method for preparation of bulk YMnO$_3$. This synthesis involves mixing binary oxides with a mortar and pestle followed up by heating treatment (firing). It is important that the particles are homogeneously mixed in the mortar, and that the particle size is small and uniform. To ensure that the reaction by diffusion occurs at the particle interphase during heating, the powder is pressed into greenbodies, which leads to smaller diffusion distance between the particles. For binary oxides with high coordination numbers, a lot of energy is needed to overcome the energy barrier in the lattice and diffuse over to another site. For such oxides, high temperature is needed. The reaction rate also depends on temperature and by increasing the temperature, the reaction rate increases. The rule of thumb is to use temperatures that are 2/3 of the materials melting temperature. For YMnO$_3$, one firing at temperature 1350°C is often not enough to obtain a pure hexagonal phase[47]. Regrinding, pressing and reheating is necessary to ensure reorientation of the powder to bring more particle surfaces in contact. Different atmospheres during heating can be used to adjust the ion oxidation state in YMnO$_3$. Reducing atmosphere (N$_2$) adjusts the low valence state, while oxidizing atmosphere (pure O$_2$) ensure high valence state[15]
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2.5.3 Solid state sintering

Since most of the samples prepared in this master thesis are polycrystalline ceramics, they need to be solid state sintered. The term “sintering” refers to the pore shrinkage, pore shape change, and grain growth which particles in contact undergo during heat treatment[48]. The major driving force of the sintering process is the achievement of minimum free energy in the material. The possible mass transport mechanisms for solid state sintering are given in Fig 2.5 [49]. The driving force for evaporation-condensation is the difference in vapour pressure for convex and concave surfaces. While for surface diffusion, lattice diffusion from surface, lattice diffusion from grain boundaries and grain boundary diffusion the driving mechanism is vacancy concentration across curves surfaces. Since these mechanisms depends on the activation of material transport, they are strongly temperature dependent. Higher sintering temperatures are needed when the material exhibits stronger bonds in the crystal lattice. Polycrystalline samples of YMnO$_3$ need high sintering temperature, and densities of >90% have been reported when sintered at 1450 °C for 10 hours [24].

![Diagram of mass transport mechanisms in solid state sintering.](image)

**Fig 2.8:** Mass transport mechanisms in solid state sintering.

During the first stage of the sintering process, a very small degree (~5 %) of densification occurs, but neck formation between particles are occurring by surface diffusion (and evaporation-condensation if the vapor pressure is high enough). In this stage, the pores are open with interconnected porosity. Most of the densification (~94 %) occurs in the intermediate stage and grain boundaries are created. The pores are still open, but smaller compared to the first stage. At the end of the intermediate stage, the grain growth starts. During the final stage of the sintering process, the last part of densification occurs. The pores are closed, and are so small that they cannot hinder grain growth. Large agglomerates in the powder is important, as these will act as large particles that might be responsible for exaggerated grain growth.
3. Experimental

In this chapter, the experimental methods used for this project is described. A list of the used binary oxides and other chemicals, their purity and manufacturers are listed in Table 3.1.

Tab. 3.1: List of used binary oxide, manufactory and purity.

<table>
<thead>
<tr>
<th>Binary Oxides</th>
<th>Molecular formula</th>
<th>Purity</th>
<th>Manufactory</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yttrium(III) oxide</td>
<td>Y₂O₃</td>
<td>99.99 wt-%</td>
<td>Sigma-Aldrich</td>
</tr>
<tr>
<td>Manganese(III) oxide</td>
<td>Mn₂O₃</td>
<td>99.0 wt-%</td>
<td>J.T Baker</td>
</tr>
<tr>
<td>Titanium(IV) oxide</td>
<td>TiO₂ (anatase)</td>
<td>99.80 wt-%</td>
<td>Sigma-Aldrich</td>
</tr>
<tr>
<td>Activated charcoal</td>
<td>C</td>
<td>≥ 98 wt-%</td>
<td>Merck KGaA</td>
</tr>
<tr>
<td>Ethylene glycol</td>
<td>C₄H₆O₆</td>
<td>99 %</td>
<td>Merck</td>
</tr>
<tr>
<td>Manganese(II)carbonate hydrate</td>
<td>Mn(CO₃)×H₂O</td>
<td>99.5 wt-%</td>
<td>Sigma-Aldrich</td>
</tr>
<tr>
<td>Yttrium(III) acetate hydrate</td>
<td>Y(O₂C₂H₄)₃ ×H₂O</td>
<td>99.9 wt-%</td>
<td>Sigma-Aldrich</td>
</tr>
<tr>
<td>Titanium(IV) isopropoxide</td>
<td>Ti(OCH(CH₃)₂)₄</td>
<td>97 wt-%</td>
<td>Sigma-Aldrich</td>
</tr>
<tr>
<td>Citric acid</td>
<td>C₆H₈O₇</td>
<td>&gt;99.0 wt-%</td>
<td>Merck</td>
</tr>
<tr>
<td>Reagent grade ethanol</td>
<td>C₂H₅O</td>
<td>≥ 99.8 vol-%</td>
<td>VWR International /</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Sigma Aldrich</td>
</tr>
</tbody>
</table>

3.1 Preparation of powders and sintered samples

3.1.1 Solid State synthesis of crystalline powder

_Prepation of YMnO₃ and yttrium deficient YMnO₃ powder_

All samples of YₓMnO₃ for x=−0.05 to x=−0.30, including YMnO₃ were prepared by conventional solid state synthesis. Prior to use, the binary oxides were dried at 500 °C for 12 hours. Around 200 °C, the oxides were moved from the furnace and kept in a desiccator, where they after short time were weighed out in stoichiometric amounts. The reactants were thoroughly mixed in an agate mortar, before uniaxial pellets of 15 mm were immersed in sacrificial powder on alumina crucibles and fired once in a clean furnace in air at 1450 °C for 12 hours. The used heating and cooling rates was 200 °C/h for every samples.
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Preparation of porous supports for the asymmetric membrane

The stoichiometric powder of YMn$_{0.85}$Ti$_{0.15}$O$_3$ to be used for the porous support was also synthesized by the solid state method. The binary oxides were mixed with carbon black (pore former), and fed into a 250 ml polyethylene bottle. The ratio between carbon black and the total weight of oxides was 1:5. The bottle was filled with yttrium stabilized zirconia (YTZ) grinding media (Ø 5mm), and to achieve a slurry, reagent grade ethanol was added until it reached ¾ of the bottle volume. Table 3.2 gives the composition of the porous powder mixture.

<table>
<thead>
<tr>
<th>Chemical</th>
<th>Amount [g]</th>
<th>Amount [wt-%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mix of binary oxides:</td>
<td>20.01</td>
<td>83.33</td>
</tr>
<tr>
<td>Y$_2$O$_3$, Mn$_2$O$_3$, TiO$_2$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carbon black</td>
<td>4.002</td>
<td>16.67</td>
</tr>
<tr>
<td>Total mixture</td>
<td>24.01</td>
<td>100.0</td>
</tr>
</tbody>
</table>

The milling time for the slurry was 16 hours, before the grinding media was removed and the slurry was dried in a rotation evaporator. The dried powder was mortared for a short time, before it was uniaxial pressed into pellets with diameters of 30mm and heights of 4-5mm. The temperature program for calcination and sintering of the pellets, seen in Figure 3.1, was chosen to eliminate the pore former and to gain some structural strength to the support. A slow heating rate and dwell step at 500°C was used to ensure slow burn-off of carbon black, which in turn reduces the probability of destroying the support. The burn-off of carbon was performed in a calcination furnace, while sintering at higher temperatures was performed in a clean furnace. The density of the substrates were measured after sintering by a caliper, followed by a characterization in XRD. The first pellet that was made (P1) was first sintered at 1200 °C for 1 hour in air after the carbon burn-off. This gave a non-phase pure sample, and a second firing at same temperature and time was done in N$_2$ atmosphere. The second pellet (P2) was only sintered in N$_2$ atmosphere, as seen in Figure 3.1.

![Fig 3.1: Temperature program used during sintering of porous support (YMn$_{0.85}$Ti$_{0.15}$O$_3$).](image-url)
3.1.2 Preparation of the active layer – citric acid method

The citric acid synthesis method described in this section is based on Bergum et al. [17], who published a paper giving a synthesis route yielding nanocrystalline YMnO$_3$. Nanocrystalline powders were made to ensure that the dip coating results in a homogeneous layer on top of the porous support.

Nanocrystalline powder was prepared by stoichiometric amounts of Y$^{3+}$ and Mn$^{3+}$ precursors and the Ti$^{4+}$ precursor titanium isopropoxide (TTIP). The preparation of Y$^{3+}$ and Mn$^{3+}$ precursors and the gravimetrically determination of the precursors molality can be seen in Appendix B. 1.4 ml of TTIP was added to a stirred solution containing 0.03 mole of citric acid (CA) and 1 dl distilled water in a small beaker. A 5 ml syringe with a glass tube attached by a tight rubber tube, see Figure 3.2, was used to suck up TTIP liquid and deliver it to the beaker.

![Syringe with a leakage proof attached glass tube.](image)

To avoid hydrolyzation, argon gas was flushed into the syringe before start, and during the delivery argon gas was flushed to the tip of the glass tube all the way to the CA solution. Some of the TTIP hydrolyzed when it came in contact with the solution, but after 1 h of stirring (440 rpm) on a hotplate with a temperature of 60 °C, the solution became blank, and all TTIP was solved in the CA solution. The cation precursors was held into a 2 L beaker and ethylene glycol (ET) was added. The molar ratio used between CA and ET was 1:1. The beaker was placed on a hotplate with a temperature of 150 °C and stirred (500 rpm) with a magnet until the gel had formed. A list of the content in the solution is presented in Table 3.3.

<table>
<thead>
<tr>
<th>$\text{Y}^{3+}$ precursor</th>
<th>$\text{Mn}^{3+}$ precursor</th>
<th>TTIP</th>
<th>CA$_{\text{total}}$</th>
<th>Ethylene glycol</th>
<th>Amount of YMn$<em>{0.85}$Ti$</em>{0.15}$O$_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0315 mol</td>
<td>0.0267 mol</td>
<td>0.00470 mol</td>
<td>0.780 mol</td>
<td>0.780 mol</td>
<td>0.0315 mol</td>
</tr>
<tr>
<td>721 g</td>
<td>621 g</td>
<td>1.40 ml</td>
<td>361 g</td>
<td>105 g</td>
<td>6.04 g</td>
</tr>
</tbody>
</table>
Chapter 3: Experimental

The gel was dried at 140 °C for 12 hours in a drying cabinet, before it was slowly heated to 200 °C for 3 hours to make it possible to extract the gel into a crystallization bowl, without breaking the beaker. The amount of the gel was about a third of the volume of the bowl. The dried gel was calcined in a crystallization bowl with the temperature program described in Figure 3.3. An overview of the whole citric acid method used in this chapter can be viewed in the flowchart in Figure 3.4. Calcined powder was then annealed in air at 800 °C for 2 hours to become nanocrystalline. The powder was characterized by XRD before and after annealing.

**Fig. 3.3:** Temperature program for calcination of the Y$_{0.85}$Ti$_{0.15}$O$_3$ gel.

**Fig 3.4:** Flowchart over the citric acid method used to prepare nanocrystalline Y$_{0.85}$Ti$_{0.15}$O$_3$ powder
3.1.3 Deposition of the active layer by dip coating

To limit the deposition of the dense functional layer to one side, the porous supports were attached to glass plates using liquid latex. A layer of liquid latex was smeared between the glass plate and the support, on the sides and a thin layer was deposited around the edges on the top surface. After solidification of the latex, a solution containing 0.626 g nanocrystalline YMn$_{0.85}$Ti$_{0.15}$O$_3$ powder and 22.5 ml ethanol was made (0.027g/ml). The nanocrystalline powder was sieved (100 µm) prior to limit agglomerates. To ensure randomly distributed particles and to avoid sedimentation during the dip coating, the solution was ultra-sonified for 1 min before each round of coating. A functional layer was deposited by dipping the glass plates with porous supports into the solution for 5 seconds. The membranes were drying for 10 min in room temperature, before the next layer was deposited (double coated). The first membrane that was made (P1) was triple coated, while the second membrane (P2) was double coated. Before sintering, membranes were dried for 1 hour in room temperature and carefully removed from the latex and the glass plates. An overview of sintering temperatures, sintering hours and number of deposited layers on each membranes are given in Table 3.4.

<table>
<thead>
<tr>
<th>Porous substrate sample</th>
<th>Number of dip coatings</th>
<th>Sintering temperature [˚C]</th>
<th>Time [h]</th>
<th>Atmosphere</th>
</tr>
</thead>
<tbody>
<tr>
<td>P1</td>
<td>3</td>
<td>1200/1400</td>
<td>2/1</td>
<td>Air</td>
</tr>
<tr>
<td>P2</td>
<td>2</td>
<td>1400</td>
<td>1</td>
<td>Air</td>
</tr>
</tbody>
</table>

3.2 Characterization of the powder

3.2.1 Structure and composition by x-ray diffraction

X-ray diffraction were used to characterize the powder by using BRUKER D8 ADVANCE DAVINCI with Cu Ka radiation. Most of the diffractograms were scanned for 4 hours with a 2θ range of 14-110˚ and a linear angle of 0.3”. A shorter scan for 30 min with a 2θ range of 14-75˚ where mainly used to have control of the synthesis, while longer scans was used to find structural parameters. DIFFRAC.EVA V 3.0 was used to match the XRD paths to PDF cards.

3.2.2 Structure and composition analysis by Rietveld and Pawley method.

The analysis of XRD paths were done in Bruker AXS Topas-academic software version 4.2. The XRD path of YMnO$_3$, porous support (P2) and nanocrystalline YMn$_{0.85}$Ti$_{0.15}$O$_3$ were refined for unit cell parameters, atomic positions in the unit cell, background and sample displacement. An overview of instrument and correction settings are seen in Table 3.5. Both the low
temperature - and high temperature phase for hexagonal YMnO$_3$ were used as starting values in the Rietveld refinement. The starting values for unit- and atomic positions are obtained from Gibbs et.al[27] and can be seen in Table 3.6 and Table 3.7 for space group $P6_3cm$ and $P6_3/mmc$ respectively. A figure showing the different atom sites in space group $P6_3cm$ is seen in Appendix A.

XRD diffractograms of $Y_{1+x}$MnO$_3$ for $x=-0.05$ to $x=-0.30$, were only refined for unit cell parameters, background and sample displacement (Pawley method), due to the anisotropic tension in Y deficiency samples making it hard to fit all the intensities of the peaks. An hkl phase with space group $P6_3cm$ was used with the same unit cell parameters, instrument – and corrections settings as for the Rietveld analysis.

**Tab. 3.5:** Instrument and correction settings for Bruker AXS Topas-academic software version 4.2.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary radius/Secondary radius</td>
<td>280 mm/280 mm</td>
</tr>
<tr>
<td>2θ angular range of LPSD</td>
<td>3' *</td>
</tr>
<tr>
<td>FDS angle</td>
<td>0.3' *</td>
</tr>
<tr>
<td>Linear PSD Sample length</td>
<td>20 mm'</td>
</tr>
<tr>
<td>Source length</td>
<td>12 mm</td>
</tr>
<tr>
<td>Sample length</td>
<td>15 mm</td>
</tr>
<tr>
<td>Receiving slit length</td>
<td>12 mm</td>
</tr>
<tr>
<td>Primary/secondary soller</td>
<td>2.5'</td>
</tr>
<tr>
<td>N Beta</td>
<td>30</td>
</tr>
<tr>
<td>LP factor</td>
<td>0</td>
</tr>
<tr>
<td>Background</td>
<td>7</td>
</tr>
</tbody>
</table>

*These parameters varies with a short scan of 30 min.

**Tab. 3.6:** Structural parameters within space group $P6_3cm$. The unit cell parameters are $a=6.141\ 51(9)$ Å and $c=11.4013(2)$ Å. Data is taken from Gibbs et al.[27].

<table>
<thead>
<tr>
<th>Atoms</th>
<th>Np</th>
<th>x</th>
<th>y</th>
<th>z</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y1</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0.2728(5)</td>
</tr>
<tr>
<td>Y2</td>
<td>4</td>
<td>1/3</td>
<td>2/3</td>
<td>0.2325(4)</td>
</tr>
<tr>
<td>Mn</td>
<td>6</td>
<td>0.3177(9)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>O1</td>
<td>6</td>
<td>0.3074(4)</td>
<td>0</td>
<td>0.1626(4)</td>
</tr>
<tr>
<td>O2</td>
<td>6</td>
<td>0.6427(3)</td>
<td>0</td>
<td>0.3355(4)</td>
</tr>
<tr>
<td>O3</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0.4744(6)</td>
</tr>
<tr>
<td>O4</td>
<td>4</td>
<td>1/3</td>
<td>2/3</td>
<td>0.0169(5)</td>
</tr>
</tbody>
</table>
Tab. 3.7: Structural parameters within space group $P6_3/mmc$. The unit cell parameters are $a=3.618961(15) \, \text{Å}$ and $c = 11.34090(9) \, \text{Å}$. Data is taken from Gibbs et al. [27]

<table>
<thead>
<tr>
<th>Atoms</th>
<th>Np</th>
<th>x</th>
<th>y</th>
<th>z</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y1</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Mn</td>
<td>2</td>
<td>1/3</td>
<td>2/3</td>
<td>1/4</td>
</tr>
<tr>
<td>O1</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>1/4</td>
</tr>
<tr>
<td>O2</td>
<td>4</td>
<td>1/3</td>
<td>2/3</td>
<td>0.085 57(7)</td>
</tr>
</tbody>
</table>

### 3.2.3 Microstructure

The cross section and the surface of the asymmetric membranes was examined with scanning electron microscope (SEM). The SEM micrographs were made on a Hitachi S-3400N using a Rhodium electron-source with SE and BSE detector. Applied voltage was 20 kV and SE was used to obtain the SEM micrographs. The membranes were attached to the sample holder using a double-sided sticky carbon tape. Prior to SEM, were the membranes divided in two, polished with grade # 800 and grade #1200 silicon carbide grinding paper, before they were washed in ethanol and ultra-sonified for 3 min. The polished cross-section was dried in a drying chamber in 10 min. Since the material is conductive, it was not necessary to coat the samples with gold or carbon.

### 3.3 Affect by atmosphere on unit cell parameters in yttrium deficient YMnO$_3$

To see how the unit cell parameters change in inert (N$_2$) and oxidizing (O$_2$) atmospheres, powders of Y$_{1+x}$MnO$_3$ for $x = -0.05$ to $x = -0.30$, prepared by solid state synthesis were placed in each their alumina crucibles and heated in a tube furnace with N$_2$ atmosphere. A characterization of the heated samples by x-ray diffraction were performed shortly after. The samples were then heated in O$_2$ atmosphere, with a slow cooling rate. Figure 3.5 represents the temperature program in both N$_2$ and O$_2$ atmosphere.

![Temperature program used for heating the Y deficient samples of YMnO$_3$ in N$_2$ (left) and O$_2$ (right) atmosphere.](image)

**Fig. 3.5:** Temperature program used for heating the Y deficient samples of YMnO$_3$ in N$_2$ (left) and O$_2$ (right) atmosphere.
Chapter 3: Experimental

3.4 Mass change in YMnO$_3$ and Y$_{0.80}$MnO$_3$ in varying atmosphere

Thermogravimetric analysis (TGA) of YMnO$_3$ and Y$_{0.80}$MnO$_3$ were performed using a Jupiter Netzsc STA 449C. The same temperature program was used for both samples, as seen in Figure 3.6. A more detailed program of the different stages and dwell times can be seen in Appendix F. Two different backgrounds were obtained by performing the measurement according to the program with an empty alumina crucible in nitrogen and oxygen. To obtain a good result, the background scan was subtracted from the sample scan. Gas flow rate of the aperture was set to 30 ml/min.

Fig. 3.6: Temperature and time program for the TGA analysis on YMnO$_3$ and Y$_{0.80}$MnO$_3$. 

![Temperature and time program for TGA analysis](image.png)
4. Results

4.1 Structure and characterization of YMnO$_3$ and yttrium deficient YMnO$_3$ powder

The x-ray diffraction pattern of bulk YMnO$_3$ made from general solid state synthesis was matched PDF card 04-011-9577 for space group $P6_3cm$ using Diffrac Eva, and can be seen in Appendix D. Phase pure bulk YMnO$_3$ is further used in the next XRD diagrams to compare Y deficient YMnO$_3$ samples, including the active layer and porous supports to space group $P6_3cm$. Refined structural parameters of bulk YMnO$_3$ within space group $P6_3cm$ are given in Table 4.1. The high uncertainty of the oxygen positions in Table 4.1 is because of the weak scattering of oxygen. The refinement gave unit cell parameters $a=6.1398(3)$ Å and $c=11.3990(2)$ Å, and a goodness of fit ($R_{wp}$) of 8.162. The Rietveld refinement (difference between experimental and calculated values) of bulk YMnO$_3$ is seen in Appendix D.

<table>
<thead>
<tr>
<th>Atoms</th>
<th>Np</th>
<th>x</th>
<th>y</th>
<th>z</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y1</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0.283(7)</td>
</tr>
<tr>
<td>Y2</td>
<td>4</td>
<td>1/3</td>
<td>2/3</td>
<td>0.242(7)</td>
</tr>
<tr>
<td>Mn</td>
<td>6</td>
<td>0.333(7)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>O1</td>
<td>6</td>
<td>0.301(1)</td>
<td>0</td>
<td>0.159(4)</td>
</tr>
<tr>
<td>O2</td>
<td>6</td>
<td>0.643(9)</td>
<td>0</td>
<td>0.336(2)</td>
</tr>
<tr>
<td>O3</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0.47(5)</td>
</tr>
<tr>
<td>O4</td>
<td>4</td>
<td>1/3</td>
<td>2/3</td>
<td>0.022(5)</td>
</tr>
</tbody>
</table>

4.1.1 Structural analysis of Y$_{1+x}$MnO$_3$ by x-ray diffraction and refined unit cell parameters by Pawley method

X-ray diffractograms of yttrium deficient samples as prepared from solid state synthesis reveals a high degree of freedom of the cation R/Mn ratio. From Y$_{0.85}$MnO$_3$ some weak reflections appear at 18°, 32.5° and 58.6° 2θ that increase in intensity with increased Y deficiency. This new emerging phase was identified as Mn$_3$O$_4$ (hausmannite) with trigonal structure. The most pronounced intensities from this new emerging phase are labelled by a star in Figure 4.1. In the lower end of Figure 4.2, a better view of the shift towards higher 2θ for the Bragg reflections from the (110), (111) and (112) planes is seen with increased yttrium deficiency. The same shift towards right can be seen for the Bragg reflection belonging to the (004) plane. There is not observed any pronounced broadening of peaks with increased yttrium deficiency.
Fig 4.1: X-ray diffractograms of $\gamma$MnO$_3$ samples with varying yttrium deficiency as prepared by solid state synthesis. The lower end of the diagram gives a better overview of the shift towards higher 2θ values for the reflections belonging to the (110), (111), (004) and (112) planes. The vertical black lines in the lower end of the diagram is a guide to the eye to see the shift of intensities better. * is indicating the pronounced intensities of Mn$_3$O$_4$ (hausmannite).
Pawley method of the x-ray diffractograms show that both the a and c unit cell parameters decrease slightly as the yttrium content in the samples decreases, which coincide with the shift of intensities found in Figure 4.1. The refined unit cell parameters in yttrium deficient YMnO₃ in both air (as prepared), N₂ - and O₂ atmospheres are represented in Figure 4.2. There is a trend in higher goodness of fit (R_wp) with increasing yttrium deficiency for as prepared samples, seen in Table 4.2. This is mainly attributed to intensity problems. These intensity problems is more pronounces with Rietveld analysis compared to Pawley fitting, and Pawley method is therefore chosen to give the best fit of the x-ray diffractograms. There is observed a slight degree of broadening with increased yttrium deficiency, but this was not as pronounced as the intensity problems. The abnormal bad fitting seen for Y₀.₉₅MnO₃ and Y₀.₉₀MnO₃ is due to large grain sizes. These samples are not taken into consideration when evaluating trends in the unit cell parameters. This also applies for Y₀.₇₀MnO₃ that experienced peritectic decomposition during sintering.

A higher goodness of fit is obtained for yttrium deficient YMnO₃ heated in N₂ and O₂ atmospheres compared to as prepared yttrium deficient samples. A problem with broadening during Pawley method was more pronounced for these samples compared to as prepared yttrium deficient YMnO₃. This broadening is not easy to see in the x-ray diffractograms for yttrium deficient samples annealed in N₂ and O₂ atmospheres, which can be seen in Appendix E. The x-ray diffractograms of yttrium deficient samples heated in N₂ and O₂ atmospheres also reveals the same high degree of freedom of the cation R/Mn ratio, before reflections appears from Mn₃O₄ (hausmannite) around Y₀.₈₅MnO₃ up to Y₀.₇₀MnO₃. With increased yttrium deficiency in YMnO₃ for samples heated in N₂, a slight shift towards higher 2θ values for the Bragg reflections belonging to the (110), (111) and (112) planes, while the Bragg reflection belonging to the (004) plane does not change. In yttrium deficient YMnO₃ samples heated in oxidizing atmosphere a shift towards higher 2θ is seen for the Bragg reflection related to the (004) plane, while the Bragg reflections related to the (110) and (111) does not change with increased yttrium deficiency.

**Tab. 4.2:** Goodness of fit (R_wp) values from Pawley fitted x-ray diffractograms of yttrium deficient YMnO₃ made from solid state synthesis.

<table>
<thead>
<tr>
<th>Composition</th>
<th>Air (as prepared)</th>
<th>N₂ atmosphere</th>
<th>O₂ atmosphere</th>
</tr>
</thead>
<tbody>
<tr>
<td>YMnO₃</td>
<td>7.301</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Y₀.₉₅MnO₃</td>
<td>11.047</td>
<td>13.336</td>
<td>12.780</td>
</tr>
<tr>
<td>Y₀.₉₀MnO₃</td>
<td>14.072</td>
<td>14.492</td>
<td>16.745</td>
</tr>
<tr>
<td>Y₀.₈₅MnO₃</td>
<td>8.875</td>
<td>11.778</td>
<td>9.645</td>
</tr>
<tr>
<td>Y₀.₈₀MnO₃</td>
<td>10.137</td>
<td>12.285</td>
<td>10.860</td>
</tr>
<tr>
<td>Y₀.₇₅MnO₃</td>
<td>11.070</td>
<td>13.844</td>
<td>12.108</td>
</tr>
<tr>
<td>Y₀.₇₀MnO₃</td>
<td>17.404</td>
<td>15.441</td>
<td>15.739</td>
</tr>
</tbody>
</table>
Chapter 4: Results

After heat treatment in inert atmosphere (N₂), the a unit cell parameter decreases for all the yttrium deficient samples, which the c unit cell parameter increases. The opposite is true for powders annealed in oxidizing atmosphere (O₂), where the unit cell parameter a increase, while unit cell parameter c has a slight decreases with increased yttrium deficiency in YMnO₃. The arrows in Figure 4.2 is a guide to the eye to get a better view of the trends in the unit cell parameters in different atmospheres.

![Graph showing unit cell parameters a and c for varying yttrium deficiency in YMnO₃, in both as prepared in air (black), N₂ (red) and O₂ (blue) atmospheres, obtained by Pawley method.](image)

**Fig. 4.2:** Unit cell parameters a and c for varying yttrium deficiency in YMnO₃, in both as prepared in air (black), N₂ (red) and O₂ (blue) atmospheres, obtained by Pawley method. The arrows are a guide to the eye, to see the trends more clearly.

A decrease in unit cell volume is seen with increased Y deficiency for all powders prepared in both air, N₂ and O₂ atmospheres. The unit cell volume and the c/a ratio is seen in Figure 4.3. In oxidizing atmosphere the c/a ratio decreases with less Y in the structure, while it is opposite in inert atmosphere where the c/a ratio increases.
Chapter 4: Results

4.2 Mass change of YMnO$_3$ and Y$_{1+x}$MnO$_3$ (x < 0) with varying conditions

Thermogravimetric analysis (TGA) show that upon heating in O$_2$ and N$_2$ atmosphere and an anomalously behavior is seen for both YMnO$_3$ and Y$_{0.80}$MnO$_3$. Upon heating in O$_2$ atmosphere, YMnO$_3$ experience an increase followed up by an abnormal and substantial mass loss after 800°C. Y$_{0.80}$MnO$_3$ experience a strange increase in oxygen content around 700°C in oxidizing atmosphere. This together with the choppy behavior seen for Y$_{0.80}$MnO$_3$ in N$_2$ cannot be explained. TGA graphs for both Y$_{0.80}$MnO$_3$ and YMnO$_3$ in O$_2$ and N$_2$ atmosphere is shown in Figure 4.4 and Figure 4.5 respectively. A reversible oxygen adsorption is found during cooling for Y$_{0.80}$MnO$_3$ and YMnO$_3$ in oxidizing atmosphere. The calculated oxygen content ($\delta$) for both powders is left to the discussion, as there is no obvious temperatures for the compounds to be stoichiometric. Oxygen content is not calculated upon heating, because of the strange behaviors as mention above.

Fig. 4.3: Unit cell volume and c/a ratio for varying yttrium deficiency in YMnO$_3$, in both as prepared in air (black), N$_2$ (red) – and O$_2$ (blue) atmospheres, obtained by Pawley method.
Chapter 4: Results

![Thermogravimetric data showing mass loss/gain and temperature as a function of time in O₂ atmosphere upon a) heating and b) cooling.](image1)

**Fig. 4.4:** Thermogravimetric data showing mass loss/gain and temperature as a function of time in O₂ atmosphere upon a) heating and b) cooling.

![Thermogravimetric data showing mass loss/gain and temperature as a function of time in N₂ atmosphere upon a) heating and b) cooling.](image2)

**Fig. 4.5:** Thermogravimetric data showing mass loss/gain and temperature as a function of time in N₂ atmosphere upon a) heating and b) cooling.

### 4.3 Structural analysis of the asymmetric membrane

#### 4.3.1 Structure and composition of porous support of YMn$_{0.85}$Ti$_{0.15}$O$_3$ prepared by solid state synthesis

The addition of carbon black gave successfully porous materials, and the densities and % of crystallographic density compared to YMnO$_3$ are given in Tab. 4.3. The crystallographic density of YMnO$_3$ was found to be 5.132 g/cm$^3$. 

32
Tab. 4.3: Crystallographic density, density of the pellets and sintering atmospheres for the two porous support P1 and P2.

<table>
<thead>
<tr>
<th>Porous substrate sample</th>
<th>Sintering atmosphere</th>
<th>Density [g/cm$^3$]</th>
<th>% of crystallographic density</th>
</tr>
</thead>
<tbody>
<tr>
<td>P1</td>
<td>Air/N$_2$</td>
<td>2.1358</td>
<td>41.62</td>
</tr>
<tr>
<td>P2</td>
<td>N$_2$</td>
<td>2.0220</td>
<td>39.39</td>
</tr>
</tbody>
</table>

Both the substrates were slightly skewed after sintering. This bias originates from the uniaxial pressing as pressure gradients easily can occur, especially with a 30 mm diameter on the pressing tool.

The porous support P1 made by solid state synthesis, was not phase pure after first sintering in air, and some of the reflections belonging to the secondary phase was matched a PDF card belonging to yttrium oxide (Y$_2$O$_3$), while the rest was not identified. It was however not matched with an orthorhombic structure. After a second firing in N$_2$ the porous support became phase pure. The x-ray diffractograms of the porous support P1 fired once in air and reheated in N$_2$ is seen in Appendix C. A displacement of the reflections towards lower 2θ values is observed for the phase pure titanium substituted sample compared to pure YMnO$_3$. This displacement has occurred because of a height difference when preparing the samples prior to XRD scans. A posteriori displacement correction of the data, reveals a displacement of the (004) reflection towards higher 2θ values, while the reflections from the (110), (111) and (112) planes are still displaced towards lower 2θ values.

The same shift of the Bragg reflections is also seen for the porous support P2, in Figure 4.6. This sample was phase pure after one sintering in N$_2$. In both XRD scans of the porous supports a decrease in peak intensity together with an increase in peak broadening is observed compared to YMnO$_3$ sample.
Fig. 4.6: X-ray diffractograms of the porous support (P2) of YMn$_{0.85}$Ti$_{0.15}$O$_3$ sintered in N$_2$ (purple). The x-ray diffractogram is compared to pure YMnO$_3$ within space group $P6_3cm$ made from general solid state synthesis. An excerpt of the (110), (111), (004) and (112) reflections with a smaller range of 2θ values is inserted in the top-right corner of the figure to give a better view of the displacements.

Refined structural parameters and unit cell parameters of the porous support P2 within space group $P6_3cm$, are seen in Table 4.4. Rietveld analysis was not performed on the x-ray diffractogram of the porous support P1 (seen in Appendix C), as both samples comes from the same powder. In Table 4.4, the unit cell parameters and structural parameters of the porous support P2 are compared to bulk YMnO$_3$ made by solid state synthesis. Unit cell parameter a for porous support P2 increases, while the unit cell parameter c decreases compared to bulk YMnO$_3$. The Refined x-ray diffractogram of the porous support P2, can be seen in Appendix D.
**Tab. 4.4:** Structural and unit cell parameters obtained from refined diffractograms of porous support P2 (YMn$_{0.85}$Ti$_{0.15}$O$_3$) and YMnO$_3$ made from solid state synthesis within space group $P6_3cm$.

<table>
<thead>
<tr>
<th></th>
<th>YMnO$_3$</th>
<th>YMn$<em>{0.85}$Ti$</em>{0.15}$O$_3$ (P2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$R_{wp}$</td>
<td>8.162</td>
<td>15.876</td>
</tr>
<tr>
<td>$a$ [Å]</td>
<td>6.1398(3)</td>
<td>6.1778(1)</td>
</tr>
<tr>
<td>$c$ [Å]</td>
<td>11.3990(2)</td>
<td>11.367(2)</td>
</tr>
<tr>
<td>$V$ [Å$^3$]</td>
<td>372.14(4)</td>
<td>375.7(1)</td>
</tr>
<tr>
<td>Y1 $z$</td>
<td>0.283(7)</td>
<td>0.27(4)</td>
</tr>
<tr>
<td>Y2 $z$</td>
<td>0.242(7)</td>
<td>0.23(8)</td>
</tr>
<tr>
<td>Mn $x$</td>
<td>0.333(7)</td>
<td>0.33(5)</td>
</tr>
<tr>
<td>O1 $x$</td>
<td>0.30(1)</td>
<td>0.29(5)</td>
</tr>
<tr>
<td>O1 $z$</td>
<td>0.159(4)</td>
<td>0.16(3)</td>
</tr>
<tr>
<td>O2 $x$</td>
<td>0.643(9)</td>
<td>0.65(3)</td>
</tr>
<tr>
<td>O2 $z$</td>
<td>0.336(2)</td>
<td>0.34(0)</td>
</tr>
<tr>
<td>O3 $z$</td>
<td>0.47(5)</td>
<td>0.48(9)</td>
</tr>
<tr>
<td>O4 $z$</td>
<td>0.022(5)</td>
<td>0.02(4)</td>
</tr>
</tbody>
</table>

4.3.2 Structure and composition of nanocrystalline YMn$_{0.85}$Ti$_{0.15}$O$_3$ prepared by citric acid method.

Nanocrystalline powder of YMn$_{0.85}$Ti$_{0.15}$O$_3$ made by citric acid method was made to avoid sedimentation during the deposition of the active layer on top of the porous support, and to obtain a dense functional layer. Pure hexagonal nanocrystalline powder of YMn$_{0.85}$Ti$_{0.15}$O$_3$ was obtained, and is shown in Figure 4.7 together with bulk YMnO$_3$ prepared by solid state synthesis for comparison. Titanium substitution gives a phase transition from $P6_3cm$ to $P6_3/mmc$, especially evidenced by the disappearance of the (012) and (022) super reflections marked by arrows in Figure 4.7. Reflection broadening is also observed for the entire range of reflections in nanocrystalline powder of YMn$_{0.85}$Ti$_{0.15}$O$_3$ compared to bulk YMnO$_3$. The Bragg reflections from the (110) and (111) planes have a pronounced shift to lower 2θ values compared to YMnO$_3$, and the Bragg reflection from the (004) plane shift towards higher 2θ values.
Chapter 4: Results

Fig. 4.7: X-ray diffractogram of hexagonal nanocrystalline YMn$_{0.85}$Ti$_{0.15}$O$_3$ within space group $P6_3/mmc$ compared to bulk hexagonal YMnO$_3$ with space group $P6_3cm$. An excerpt of the (110), (111), (004) and (112) reflections with a smaller range of 2θ values is inserted in the top-right corner of the figure to give a better view of the displacements.

The unit cell parameters $a$ and $c$, as well as structural parameters for nanocrystalline YMn$_{0.85}$Ti$_{0.15}$O$_3$ were found by Rietveld analysis. The XRD pattern was refined within space group $P6_3/mmc$ and gave refined unit cell parameters of $a = 3.565(0)$ Å and $c = 11.26(6)$ Å. Structural parameters from Rietveld refinement of the x-ray diffractogram of the active layer of YMn$_{0.85}$Ti$_{0.15}$O$_3$ is seen in Table 4.5, and the Rietveld refinement can be seen in Appendix D. The only atom position released is the z direction of the O2, because this site is not in high symmetric position. The refinement did not yield an adequately fit to the experimental XRD graph, illustrates by the high goodness of fit of 18.045. One of the main reasons for this is due to restriction in Topas, which does not take into account anisotropic strain the material experience during Ti substitution on Mn site. The crystallite size of nanocrystalline YMn$_{0.85}$Ti$_{0.15}$O$_3$ was found to be 36.1 ± 3 nm.
4.4 Microstructure of porous support and active layer of the membrane

The active layers of nanocrystalline YMn$_{0.85}$Ti$_{0.15}$O$_3$ was not dense after sintering. The porous support also as expected became denser after sintering. An overview of the surface of the porous support and active layer is seen in Figure 4.8 for both the double coated and triple coated membranes. It is observed that the porous supports are denser compared to the active layer. The double coated and the triple coated membranes had also almost the same membrane thickness of 25 μm and 25.5 μm, seen in Figure 4.9, showing the cross sections of the membranes.

<table>
<thead>
<tr>
<th>Atoms</th>
<th>Np</th>
<th>x</th>
<th>y</th>
<th>z</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y1</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Mn</td>
<td>2</td>
<td>1/3</td>
<td>2/3</td>
<td>1/4</td>
</tr>
<tr>
<td>Ti</td>
<td>2</td>
<td>1/3</td>
<td>2/3</td>
<td>1/4</td>
</tr>
<tr>
<td>O1</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>1/4</td>
</tr>
<tr>
<td>O2</td>
<td>4</td>
<td>1/3</td>
<td>2/3</td>
<td>0.08(8)</td>
</tr>
</tbody>
</table>
Fig. 4.8: SEM micrographs from the surface of both active layer and porous support of the sintered asymmetric membranes.
Fig. 4.9: SEM micrographs showing the cross section of the sintered asymmetric membranes. Porous support dip coated a) 2 times and b) 3 times
5. Discussion

5.1 The influence of yttrium deficiency in YMnO$_3$.

Yttrium deficiency in hexagonal YMnO$_3$ can be charge compensated by oxygen vacancies, which gives an oxygen poor structure, seen in (5.1) or they can be charge compensated by oxidation of a fraction of Mn$^{3+}$ to Mn$^{4+}$ as illustrated in (5.2).

\[ Y_{1-x}^3+Mn^{3+}O_{3-\frac{3x}{2}} \]  
\[ (5.1) \]

\[ Y_{1-x}^3+Mn_{1-3x}^{2+}Mn_{3x}^{4+}O_{3} \]  
\[ (5.2) \]

It is important to know that these two possibilities are extremes, and that there might exists both oxygen vacancies and Mn$^{4+}$ at the same time to charge compensate for the yttrium deficiency. The defect chemistry in stoichiometric YMnO$_3$ is not well known, and especially not for yttrium deficient YMnO$_3$. The following sections provide a discussion about how reducing/oxidizing atmospheres influence the crystal structure, and suggest probable defect reactions that may arise in the material during different conditions.

5.1.1 Structure and solubility limit

All yttrium deficient samples were fired at 1723 K (1450 °C), indicating that we are in the β-Mn$_3$O$_4$ solubility region of h-YMnO$_3$ according to the estimated Y$_2$O$_3$-Mn$_2$O$_3$ pseudo-binary phase diagram[34]. Trace amounts of β-Mn$_3$O$_5$ appear in the as prepared x-ray diffraction pattern for Y$_{0.80}$MnO$_3$ and get more pronounced with a less degree of yttrium in the structure. The same trend is also seen for the yttrium deficient samples annealed in both N$_2$ and O$_2$ atmospheres. This clearly shows that the solid solubility region of β-Mn$_3$O$_4$ in yttrium deficient hexagonal YMnO$_3$ is extended more to the left in the phase diagram, seen as the dotted red line in Figure 5.1. Since there has not been sintered any yttrium deficient samples at lower temperatures, it is impossible to say what value the slope of curve will exhibit in the vertical direction (dotted orange line, Figure 5.1).

A degree of freedom in the Y/Mn ratio may explain the different unit cell parameters and phase transition temperatures found in literature for YMnO$_3$. In addition, it is promising in terms of reducing the cost, but on the other hand, mechanical and chemical properties must be further investigated to see how less yttrium in the structure affects the material.
Chapter 5: Discussion

**Fig. 5.1:** A suggestion to a revised new solid solubility limit (dotted red line) of $\beta$-Mn$_3$O$_4$ in yttrium deficient YMnO$_3$ than first expected in the pseudo–binary phase diagram for the Y-Mn-O system. The orange dotted line is just an assumption of the slope of the curve (this has not been experimentally shown in this thesis). The originally figure is reprinted from Chen et.al [34]

The surprisingly high solid solubility region even with 15-20 % yttrium deficiency in YMnO$_3$ may be due to the layered structure of hexagonal YMnO$_3$. The conducting state in hexagonal YMnO$_3$ has been attributed to the overlap between the d-orbitals of Mn in the ab-plane[24]. There is less contact between the MnO$_5$ layers along the c direction, and it is therefore reasonable to believe that a removal of a Y-ion, which is situation between the layers, not will destabilize the structure.

Moreover, the peritectic decomposition observed for Y$_{0.70}$MnO$_3$ after sintering indicates that the peritectic decomposition line left in Figure 5.1 at 1822 K is shifted towards a lower temperature ($\sim$1723 K) for highly yttrium deficient samples.

5.1.2 Distortion of the unit cell with varying partial pressure of oxygen

For a higher degree of yttrium deficiency in YMnO$_3$ prepared in air atmosphere, see Figure 4.1, a shift toward higher 2θ values is observed for Bragg reflections associated with the high intensity (110), (111), (112) and (004) planes. A shift towards higher 2θ values indicates a decrease in the distance $d_{hkl}$ between adjacent lattice planes. It is obvious that a change in $d_{hkl}$ is related to a change in a- and c unit cell parameters. The high intensity Bragg reflections related to the a and c axis are illustrated in space group $P6_3cm$ in Figure 5.2. Decreased distance between adjacent planes for intensities related to the a and c axis indicates a compression of the hexagonal structure in both a and c direction.
Fig 5.2: (a) A visualizing of the (110), (111) and (112) planes in hexagonal YMnO$_3$ with space group $P6_3cm$. All planes are dependent on direction $a$, being either perpendicular or having an angle $\alpha$ not being 0 with the $a$-direction. Decreased distance between adjacent planes will give a compression along a direction. (b) The (004) plane is perpendicular to the c-axis, and parallel to the a-axis. A decreased distance between adjacent planes gives a compression along c direction. The figure is made using VESTA software [26], with structural parameters from Gibbs et al. [27]

The same structural trend was found by Pawley fitted x-ray diffractograms of as prepared yttrium deficient samples, Figure 4.2. The refined unit cell parameters $a$ and $c$, showed a slight decreased with increased yttrium deficiency, especially from Y$_{0.85}$MnO$_3$ to Y$_{0.75}$MnO$_3$. This indicates the same compression of the hexagonal unit cell in both ab plane and c-direction. The compression leads to a volume decrease and a slight increase in the c/a ratio compared to stoichiometric YMnO$_3$. There is no obvious explanation for this behavior. However, yttrium deficiency in YMnO$_3$ can be charge compensated by oxygen vacancies or oxidation of a fraction Mn$^{3+}$ to Mn$^{4+}$. A smaller radii of Mn$^{4+}$ compared to Mn$^{3+}$ may explain the compression of the unit cell when less yttrium is introduced.

Reducing condition for stoichiometric YMnO$_3$ will be attributed to oxygen leaving the material due to the partial pressure difference between the material and the surroundings. It is suggested by Overton et.al [43] that oxygen vacancies in YMnO$_3$ preferably will leave planar atom sites in the MnO$_5$ units. An expansion of the structure was also found in both a and c directions, followed up by a decline in the tilting of MnO$_5$ units with less oxygen in the structure of YMnO$_3$. Less decline of MnO$_5$ bipyramids is also associated with less displacement of yttrium along c direction. Oxygen vacancies can be charge compensated by reduction of Mn$^{3+}$ to Mn$^{2+}$,
written in Kröger-Vink notation and is seen in equation (5.3), and since the ionic radius of Mn\(^{2+}\) is larger than Mn\(^{3+}\), the volume of the unit cell will probably increase. The larger ionic radius of Mn\(^{2+}\) will also increase the average Mn-O bond length and thus the a-axis would be expected to increase.

\[
O_2^\circ + 2Mn_{Mn}^x \rightarrow \frac{1}{2}O_2(g) + V_O^* + 2Mn_{Mn}'
\] (5.3)

It might seem reasonable to assume the same trends in yttrium deficient samples annealed in reducing atmosphere as for oxygen vacancies in stoichiometric YMnO\(_3\). In fact, the opposite happens for unit cell parameter a which decrease, while unit cell parameter c slightly decreases with increased yttrium deficiency in YMnO\(_3\). This behavior is also seen in the x-ray diffractograms of yttrium deficient YMnO\(_3\) annealed in N\(_2\) atmosphere, as a shift toward higher 2\(\theta\) values is observed for intensities related to the (110), (111) and (112), while the intensity related to the (004) plane hardly shifts. A possible explanation can be that oxygen vacancies are preferably removed from the apical sites around MnO\(_5\), instead of the planar sites. A further reasonable discussion is limited on the basis of the obtained data.

For stoichiometric YMnO\(_3\) annealed in oxidizing atmosphere it is expected that oxygen will go into the material from the atmosphere. At low temperatures, this oxygen adsorption is attributed to interstitial oxygen, as the hexagonal structure of YMnO\(_3\) is less dense compared to the related perovskite phase.Interstitial oxygen can be charge compensated by oxidation of a fraction of Mn\(^{3+}\) to Mn\(^{4+}\) and is seen by the following Kröger-Vink notation

\[
\frac{1}{2}O_2 + 2Mn_{Mn}^x \rightarrow 2Mn_{Mn}' + O_i'
\] (5.4)

The position of the oxygen interstitial in the hexagonal structure is discussed in more details in an own section. A decreased c/a ratio for yttrium deficient samples of YMnO\(_3\) annealed in oxidizing atmosphere is due to a compression along the c axis and an slight expansion along the a direction. Since oxygen interstitials is charge compensated by oxidation of Mn\(^{3+}\) to Mn\(^{4+}\), one would probably expect the structure to be compressed in the planer plane because of the smaller radius of Mn\(^{4+}\) compared to Mn\(^{3+}\). However, a reasonable explanation for this behavior cannot be conducted, and more attention has to be given to the position of the interstitial oxygen, and how this influence the structure.

Limitations of The Pawley fit

Not all calculated graphs found by Pawley fitting were able to yield a good fit to the experimental x-ray graphs of yttrium deficient YMnO\(_3\). One of the reasons for this comes from the limiting factor the fitting program exhibits, which is that it assumes stoichiometric
compound when calculating the best fit. A higher degree of yttrium deficiency, gave a higher
strain seen by the limiting fit. The strain is believed to be anisotropic, however the broadening
of the intensity related to the (004) plane was not pronounced and it is therefore hard to say.
The layered nature of the low temperature hexagonal phase is one of the reasons for the strain.
The abnormally high goodness of fit values for Y_{0.95}MnO₃ and Y_{0.90}MnO₃ is attributed to large
grain size which occurred during sintering. It is suggested that less yttrium in the structure will
increase the diffusion distance in the structure, and hence it is believed that increased yttrium
will give smaller grains. A decrease in the sintering temperature will also give smaller grains.

5.1.3 Oxygen storage ability

It was not clear where to assume stoichiometric compounds from the thermogravimetric data
during heating. This applies especially for Y_{0.80}MnO₃ during heating, seen in Figure 4.4 and
Figure 4.5. In O₂ atmosphere, the sudden increase in oxygen content around 600 °C for
Y_{0.80}MnO₃ with a following decrease has no obvious explanation. This does also apply for the
sudden decrease in oxygen content for YMnO₃ around the same temperature region. The
excess oxygen is therefore only calculated during cooling in O₂ and N₂ atmosphere, as this
behavior was more similar to what Remsen et.al[14] found for hexagonal manganites.

The assumption of stoichiometric compound during cooling was done where there was a high
probability to only have Mn^{3+} in the structure. For Y_{0.80}MnO₃ in O₂ atmosphere this is seen as
the plateau between 400°C – and 500°C, while for YMnO₃ in the same atmosphere is it assumed
to be around 450 °C. For YMnO₃ and Y_{0.80}MnO₃ in N₂ atmosphere, stoichiometric compound is
assumed to be around 400 °C and the plateau between 450-500 °C respectively. The excess oxygen
content as a function of temperature for YMnO_{3+δ} and Y_{0.80}MnO_{2.7+δ} in both O₂ and N₂
atmosphere is shown in Figure 5.3 and Figure 5.4 respectively. For the excess oxygen
calculations in Y_{0.80}MnO₃ it was assumed that less yttrium in the structure was charge
compensated by oxygen vacancies, which gives the stoichiometric compound Y_{0.80}MnO_{2.7+δ}.
There is no excess oxygen in the yttrium deficient structure (δ = - 0.002) during cooling in N₂
atmosphere, and the same observation is seen for YMnO₃. This is reasonable as the partial
pressure of oxygen is small. In oxidizing atmosphere an oxygen adsorption of δ = 0.034 is
calculated for YMnO₃, while for Y_{0.80}MnO₃ a smaller adsorption of δ = 0.013 was found.
Chapter 5: Discussion

**Fig. 5.3:** Termogravimetric data showing excess oxygen as a function of temperature in O\(_2\) atmosphere for both YMnO\(_{3+\delta}\) (black) and Y\(_{0.80}\)MnO\(_{2.7+\delta}\) (orange). The calculated excess oxygen is \(\delta = 0.034\) for YMnO\(_3\) and \(\delta = 0.013\) for Y\(_{0.80}\)MnO\(_3\) at room temperature.

**Fig. 5.4:** Termogravimetric data showing excess oxygen as a function of temperature in N\(_2\) atmosphere for both YMnO\(_{3+\delta}\) (black) and Y\(_{0.80}\)MnO\(_{2.7+\delta}\) (orange). The calculated oxygen content is \(\delta = 0.0001\) for YMnO\(_3\) and \(\delta = -0.002\) for Y\(_{0.80}\)MnO\(_3\) at room temperature.
The substantial increase in oxygen content from higher temperatures to around 800 °C is believed to be attributed to filling of oxygen vacancies in both YMnO$_3$ and Y$_{0.80}$MnO$_3$. This is also supported by the chemical expansion seen at higher temperatures for YMnO$_3$ found by Selbach et al. [24]. The filling of oxygen vacancies during cooling from $T > 800$ °C is seen by the following Kröger-Vink notation. This is the same as equation (5.3), only in opposite direction.

$$\frac{1}{2}O_2(g) + V^-_O + 2Mn'_Mn \rightarrow O^x_O + 2Mn^x_Mn$$

From approximately 400 °C – 800 °C a stabilization of Mn$^{3+}$ is assumed, which gives stoichiometric compounds as mention above. For Y$_{0.80}$MnO$_3$, it is here believed that one yttrium vacancy is charge compensated by two oxygen vacancies. This is shown by the following Kröger-Vink notation

$$Mn_2O_3 \xrightarrow{Y_{1-x}MnO_3} 3O^x_O + 2Mn^x_Mn + 2V^-_O + V''_Y$$

When the temperature is around 400 °C the oxygen content is increasing again with decreasing temperature (besides the anomalously behavior seen for Y$_{0.80}$MnO$_{2.7+\delta}$ in N$_2$). The increasing oxygen content for Y$_{0.80}$MnO$_{2.7+\delta}$ in O$_2$ atmosphere is suggested to be filling of oxygen vacancies, with a following oxidation of Mn$^{3+}$ to Mn$^{4+}$ to charge compensate, shown by

$$\frac{1}{2}O_2(g) + V^-_O + 2Mn^x_Mn \rightarrow O^x_O + 2Mn^+_Mn$$

The probability of oxygen vacancies at such low temperatures is believed to be low for stoichiometric YMnO$_{3+\delta}$. The increase in oxygen content is therefore believed to be interstitial oxygen going into the structure. Even though ion mobility is low at these temperatures, can a decrease in activation energy for ion hopping make this possible. The Kröger-Vink notation for interstitial oxygen with a following oxidation of Mn$^{3+}$ to Mn$^{4+}$ is seen in equation (5.4), with Mn$^{4+}$ described as Mn$^{3+}$ with an electron-hole. According to these results, YMnO$_3$ is still believed to be a semiconducting material with p-type conductivity at low temperatures, and n-type conductivity at temperatures around 1000 °C. The same cannot be assumed for Y$_{0.80}$MnO$_3$, as oxygen vacancies are believed to be more dominated at lower temperatures compared to YMnO$_3$. 
Chapter 5: Discussion

If yttrium deficiency in general are charge compensated by oxygen vacancies, leaving planar sites available in the Mn-plane, this may give rise to higher conductivity. As the ion conductivity is believed to be in the ab plane, a removal of oxygen will give a higher degree of available sites that \( \text{O}^{2-} \) can jump between. However, electrical conductivity measurements performed on \( \text{Y}_{0.90}\text{MnO}_3 \) in a preliminary master’s thesis \([40]\) gave a relatively low conductivity (same range as \( \text{YMnO}_3 \)). This only shows that more research has to be done to get a better understanding of the defect chemistry in yttrium deficient \( \text{YMnO}_3 \).

**The possibility of anti-sites in the hexagonal structure**

Since the ionic radii of \( \text{Y}^{3+} \) (0.96 Å \([32]\)) is not much larger than the ionic radii of \( \text{Mn}^{2+} \) (0.83 Å), there might be a possibility of anti-site between Mn and Y in the structure. The anti-site for Mn on Y site is shown for stoichiometric \( \text{YMnO}_3 \) by the following Kröger-Vink notation

\[
\text{Mn}_2\text{O}_3 \xrightarrow{\text{YMnO}_3} \text{Mn}^{x}_{\text{Mn}} + \text{Mn}^{x}_{\text{Y}} + 3\text{O}^{x}_0 \quad (5.8)
\]

Since \( \text{Mn}^{2+} \) forms at higher temperature, the anti-site is thus more likely to happen at higher temperatures. To investigate this probability, anti-site in \( \text{Y}_{0.80}\text{MnO}_3 \) was calculated in Rietveld. The anti-site gave no better fit than without anti-site. The probability is therefore suggested to be small. An empty “Y vacancy” in the middle of the trigonal bipyramidal environment would not give stabilization to the surrounding oxygen, and the structure could perhaps be destabilized. Although the suggestions is pointing towards a lower possibility for anti-site defects at lower temperature, little is known about the structure and one cannot totally decline the possibility.

**5.2.4 The positions of interstitial oxygen in \( \text{YMnO}_{3+\delta} \)**

There are several sites in the hexagonal structure that has enough space to accommodate an interstitial oxygen. Two of the most probable sites for oxygen interstitial is between three Mn atoms in the MnO\(_5\) layers, and is represented in Figure 5.5.
Chapter 5: Discussion

The main reason for these positions of interstitial oxygen is because manganese can exhibit different oxidation states. One interstitial oxygen in the middle of three Mn atoms, will oxidize two Mn$^{3+}$ to Mn$^{4+}$, while one is left as Mn$^{3+}$. It is believed that anti bonding bonds are formed between electrons from the O 2p orbital and the Mn d orbital. Due to electrostatics it is believed that the manganese atoms will be drawn towards the interstitial oxygen, while nearby oxygen will be repelled. It is not clear how this influence the rest of the unit cell. An interstitial oxygen will create an octahedral surrounding around manganese. The octahedral crystal field splitting for manganese in oxidation state Mn$^{3+}$ and Mn$^{4+}$ is viewed in Figure 5.6. The crystal field stabilization is more stable for Mn$^{4+}$ compared to Mn$^{3+}$, as there is needed more energy to excite an electron up to the $e_g$ orbitals.

![Diagram showing octahedral crystal field splitting for Mn$^{3+}$ and Mn$^{4+}$](image)

**Fig. 5.6:** Octahedral crystal field splitting showing Mn$^{3+}$ and Mn$^{4+}$. 

Fig. 5.5: Position of interstitial oxygen (yellow atom) in the $P\bar{6}_3cm$ structure, seen from the c-axis.
5.2 Structure and evaluation of the asymmetric membrane YMn\textsubscript{0.85}Ti\textsubscript{0.15}O\textsubscript{3}

5.2.1 Decreased c/a ratio with titanium substitution in YMnO\textsubscript{3}

15 \% titanium substitution in nanocrystalline YMnO\textsubscript{3} resulted among others in a disappearance of the (102) and (202) super reflections, seen in Figure 4.7. The x-ray diffraction pattern for nanocrystalline YMn\textsubscript{0.85}Ti\textsubscript{0.15}O\textsubscript{3} was matched to the paraelectric high temperature phase which indicates a stabilization of space group $P6_3/mmc$. The phase transition was expected as it has been observed in preliminary master’s thesis [[40]] and by S.Nesdal and Tomczyk.et.al[[15, 39]].

The disappearance of the same reflections were also expected for the x-ray diffraction patterns of the porous supports, but this was not observed. The Rietveld analysis of the porous support P2 within space group $P6_3/cm$, gave unit cell parameters in almost the same ranges as the unit cell parameters for YMn\textsubscript{0.90}Ti\textsubscript{0.10}O\textsubscript{3} found by S.Nesdal[15], and can be seen in Table 5.1. The unit cell parameter $a$ has a value between unit cell parameters $a$ for YMn\textsubscript{0.90}Ti\textsubscript{0.10}O\textsubscript{3} and YMn\textsubscript{0.85}Ti\textsubscript{0.15}O\textsubscript{3}, while unit cell parameter $c$ lies somewhere between values of YMn\textsubscript{0.95}Ti\textsubscript{0.05}O\textsubscript{3} and YMn\textsubscript{0.90}Ti\textsubscript{0.10}O\textsubscript{3}. It is therefore suggested that the porous supports contain less Ti, which is also supported by the non-disappearance of the Bragg reflections belonging to the (102) and (202) planes. This is a strange result as general solid state synthesis is easy to control as the amount of oxide is weighed out manually in stoichiometric amounts.

With increased titanium substitution a decrease in the c/a ratio is seen compared to pure YMnO\textsubscript{3}. The c/a ratio for YMn\textsubscript{0.85}Ti\textsubscript{0.15}O\textsubscript{3} (P2) is found to be 1.840 compared to 1.856 for YMnO\textsubscript{3}. This reduces the anisotropic thermal expansion in the high temperature phase transition from $P6_3/mmc$ to $P6_3/cm$, which also reduce microcracking in the material. It is hard to decide from the SEM micrographs of the porous support and active layer in Figure 4.8, if microcracking in the material has been reduced.
Tab. 5.1: Unit cell parameters, unit cell volume and c/a ratio from Rietveld analysis of the porous support P2 of YMn$_{0.85}$Ti$_{0.15}$O$_3$ and bulk YMnO$_3$ made by solid state synthesis, within space group P6$_3$cm. To compare values, the unit cell parameters, unit cell volume and c/a ratio is shown for increased titanium substitution in YMnO$_3$ obtained by S.Nesdal [27], and the active layer of nanocrystalline YMn$_{0.85}$Ti$_{0.15}$O$_2$.

A shift towards lower 2θ values of the Bragg reflections associated with the (110), (111) and (112) planes were observed for both porous supports and active layer compared to bulk YMnO$_3$. For the Bragg reflection belonging to the (004) plane a shift was observed towards higher 2θ values. Increased Ti in YMnO$_3$ will expand the unit cell in the basal plane, while the c-axis will be compressed. A more pronounced shift to the left is seen for nanocrystalline powder compared to the porous support for the intensity belonging to the (004) plane. This is in accordance to Vegards law that unit cell parameters will scale with the chemical composition. The increase of the unit cell length a, together with a compression along the c-axis is then attributed to a reduced tilting of the MnO$_5$ and less displacement of the yttrium atoms.

Structure analysis consideration of the active layer consisting of nanocrystalline YMn$_{0.85}$MnO$_3$ prepared by citric acid method

When dealing with Rietveld analysis of nanocrystalline powder it is important to have a tight distribution of crystallite size in the powders, as a wide distribution can cause nonrandomness. Today, it is not possible to correct for nonrandomness in Topas. Line broadening will also be an issue for nanocrystalline powders as seen in Fig. 4.7. This gives uncertainties in the calculation of crystallite size, and is probably the reason for the few nm higher value than expected for the crystallite size compared to the results from Bergum et.al found at 800˚ C. For a better indication of the distribution and correct crystallite size, TEM micrographs should be used. It should also be noted that the fundamental parameters for the D8 focus are still not perfects and correction for instrumental errors could be slightly inaccurate.

Besides the high broadening of the peaks seen in the XRD path of nanocrystalline YMn$_{0.85}$Ti$_{0.15}$O$_3$, everything coincide with a phase transition from the low temperature phase to the high temperature phase. Although Bergum et.al [17] found an increase in the unit cell length a, and a decrease in unit cell parameter c, which gave a decrease in unit cell volume with
decreasing crystallite size, there is not seen any abnormal results of the nanocrystalline YMn\textsubscript{0.85}Ti\textsubscript{0.15}O\textsubscript{3}. There is a possibility that decreased crystallite size give a higher symmetry structure because of the increased randomness caused by a loss of coherence in the structure.

5.2.2 Oxygen non-stoichiometry with titanium substitution in YMnO\textsubscript{3}

It is suggested that titanium substitution will stabilize excess oxygen in the material. The proposed mechanism for this is that Ti\textsuperscript{4+} stabilize O\textsuperscript{2−} by the formula YMn\textsubscript{1−x}Ti\textsubscript{x}O\textsubscript{3+(x/2)}, where x/2 is the same as δ. For x = 10, this value becomes δ = 0.05, that almost corresponds to the value obtained by TGA, δ=0.043, found by S.Nesdal\textsuperscript{[15]}. The stabilization of excess oxygen or in other words stabilization of interstitial oxygen is described by the following Kröger-Vink notation,

\[
2\text{TiO}_2^{\text{YMnO}_3} \rightarrow 2\text{Ti}^{\text{Mn}_x} + 4\text{O}^{\text{x-}}_0 + \text{O}^{\text{ii-}}_i \tag{5.9}
\]

This may explain the high electronic conductivity at 400 °C found for YMn\textsubscript{0.85}Ti\textsubscript{0.15}O\textsubscript{3} compared to YMnO\textsubscript{3} in the preliminary master’s thesis \textsuperscript{[40]}, as there is created more charge carrier holes. However, Ti\textsuperscript{4+} has zero electrons in the d-orbital and one would expect a lower electronic mobility. Another assumption is that Ti\textsuperscript{4+} is charge compensated by reduction of Mn\textsuperscript{3+} to Mn\textsuperscript{2+}, seen as follows

\[
\text{Mn}^{\text{x+}}_{\text{Mn}} + \text{TiO}_2^{\text{YMnO}_3} \rightarrow 2\text{Ti}^{\text{Mn}_x} + 2\text{O}^{\text{x-}}_0 + \text{Mn}^{\text{i+}}_{\text{Mn}} \tag{5.10}
\]

This will change manganese from 3d\textsuperscript{4} to 3d\textsuperscript{5}, and one electron has to go up in the fairly localized dz\textsuperscript{2} orbital as shown in Figure. 5.7. This excitation required more energy than the other oxidation states of Mn within the trigonal bipyramidal crystal field splitting.

\[\text{Mn}^{3+} - d^4 \quad \text{Mn}^{2+} - d^5\]

\[\text{Mn}^{3+} - d^4 \quad \text{Mn}^{2+} - d^5\]

Fig. 5.7: Trigonal bipyramidal crystal field splitting showing Mn\textsuperscript{3+} and Mn\textsuperscript{2+}.
Chapter 5: Discussion

5.2.3 Evaluation of the active layer and the porous supports after sintering.

Even though the synthesis of nanocrystalline powder was successful, a dense ceramic membrane was not obtained after sintering, as seen in Figure 4.8. There can be several reasons for this. One reason can be due to agglomerates in the powder that hinders close packing (smaller distance between particles) of the powder before sintering. Powder that is not sufficient close packed gives a high diffusion distance which gives limited densification during sintering. The density of particles solved in ethanol could also have been too high, which also gives a relatively poor close packed powder upon the porous support. This can also be the reason for why the double and triple dip coated membranes had almost the same thickness of the active layer, seen in Figure 4.9. To solve problems with close enough packing of powder on top of the porous support surfactant and/or ball milling can be used. After crystallization of the nanocrystalline powder a following ball milling could decrease the particle size. This powder could be used in a dispersion containing ethanol and surfactants to prevent agglomeration. The dispersion stability can have a significant effect on the performance, resulting in poor adhesion, insufficient reactivity and film thickness. There are however no reports on how to make a stable dispersion using surfactants and hexagonal YMnO$_3$.

The prospect of using titanium substitution in YMnO$_3$ as membrane material has a great advantage as the electrical conductivity is higher for titanium doped YMnO$_3$ compared to YMnO$_3$. Furthermore, a decrease in the c/a ratio have proven to give reduced microcracking which is a great advantage in processing membranes. In addition, a suggestion of ionic conductivity at lower temperatures due to interstitial oxygen is a great advantage with regards of reducing the operational temperatures for MIEC membranes. However, more attention has to be given to the defect chemistry in YMnO$_3$, and a more systematic synthesis procedure has to be found in order to obtain a dense and functional layer on top of a porous support. The two components of the membrane also have to be in the same structural composition to avoid thermal and chemical expansion problems which will lead to cracks or degradation of the membrane. In this thesis, the same structure was not obtained for both the porous support and active layer due to artificial wrongs.

5.5 Further work

The limiting understanding of point defects in hexagonal manganites, must be further investigated. Thermogravimetric analysis in both oxidizing and reducing atmospheres on 10 % and up to 30 % yttrium deficiency in YMnO$_3$ can give an indication of how the excess oxygen varies for different amount of yttrium in the structure. Dilatometry on yttrium deficient samples will also give an indication of how yttrium deficiency in YMnO$_3$ influence the phase transition between the low and high temperature phase. The understanding of defect chemistry in hexagonal manganites are important as the concentration of defects can tune different properties of the material. Thus for instance optimize the electrical conductivity in YMnO$_3$. The positions of interstitial oxygen and oxygen vacancies are crucial in the understanding of how defects in YMnO$_3$ influence the structure. These positions should be investigated by DFT calculations, to find out which position gives the lowest energy. The next stage when it comes to processing a MIEC membrane is to find a synthesis method that gives a dense and functional
layer on top of a porous support. Addition of surfactants in a slurry with nanocrystalline powder and ethanol will prevent agglomeration and give a denser packed layer before sintering. The performance (oxygen permeation rate) of such a membrane will give a good indication of how the membrane will work in practice and how the properties influence the stability.
7. Conclusion

This work was performed to investigate how the structure and oxygen storage ability changes with different yttrium content in YMnO$_3$. Hexagonal YMnO$_3$ has the ability to lose 15-20% yttrium without any notable changes in the structure and composition for yttrium deficient YMnO$_3$. Above this limit, reflections from Mn$_3$O$_4$ hausmannite appears in x-ray diffractograms. The same solubility limit was also found for yttrium deficient samples of YMnO$_3$ heated in N$_2$ and O$_2$ atmospheres. With a higher degree of yttrium deficiency in the hexagonal structure prepared in air, a slight contraction of the unit cell along both the a and c axis is found. For yttrium deficient samples annealed in oxidizing (O$_2$) atmosphere, the structure expands in the along the a axis and contracts along the c-axis. The opposite behavior is seen in reducing (N$_2$) atmosphere, where the structure expands in c direction and contract along the a axis with increased yttrium deficiency in YMnO$_3$. Above 600 °C, during heating of YMnO$_3$ and Y$_{0.80}$MnO$_3$, oxygen goes out of the materials in both N$_2$ and O$_2$ atmospheres. Upon cooling in O$_2$ atmosphere, a reversible oxygen uptake is seen around 400 °C for both YMnO$_3$ and yttrium deficient YMnO$_3$. For stoichiometric YMnO$_3$ the reversible oxygen adsorption is suggested to be attributed to interstitial oxygen, while for stoichiometric yttrium deficient YMnO$_3$ there is believed to be filling of oxygen vacancies. The asymmetric membrane of YMn$_{0.85}$Ti$_{0.15}$O$_3$ did not yield a dense functional layer on top of the porous support after sintering, and the membrane thickness was the same after double and triple dip coatings. The results from this thesis highlights the huge possibilities hexagonal manganites have for use in MIEC membranes. More attention has to be given to hexagonal manganites to get a better understanding of the properties, and to get one step further to a more environmentally friendly world.
Bibliography

12. Lein, H.L., Mechanical Properties and Phase Stability of Oxygen Permeable Membranes La0.5Sr0.5Fe1−xCoxO3−δ. 2005, Norwegian University of Science and Technology.
13. Kaus, I., et al., Oxygen transport properties in La1−xSrxFex1−yMyO3−δ (M= Cr, Ti), 0.2 <sub>bxb0</sub> 0.8, 0.2 byTi b0.5, 0.1 byCr b0.3. Solid State Ionics, 2007. 178: p. 817-826.


Appendix A: Structural parameters with titanium substitution in YMnO$_3$ within space group $P6_3cm$

The structural parameters of YMn$_{1-x}$Ti$_x$O$_3$ for $x = 0.05, 0.10$ and $0.15$ obtained by S. Nesdal[15] are found in Table A1. The structural parameters were refined within space group $P6_3cm$. A supporting illustration of the structural position is seen for the hexagonal space group $P6_3cm$ in Figure A1. In the same figure, a presentation of the tilting of MnO$_5$ bipyramids and displacement of yttrium ions associated with the low temperature phase is shown.

### Tab. A1: Refined atomic positions, lattice parameters and unit cell volume with increased Ti substitution in hexagonal YMnO$_3$ within space group $P6_3cm$ obtained by S. Nesdal [15].

<table>
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<tr>
<th></th>
<th>YMn$<em>{0.95}$Ti$</em>{0.05}$O$_3$</th>
<th>YMn$<em>{0.90}$Ti$</em>{0.10}$O$_3$</th>
<th>YMn$<em>{0.85}$Ti$</em>{0.15}$O$_3$</th>
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Fig. A1: A schematic representation of the atom positions showing the displacements of Y ions and tilting of MnO₅ bipyramids within space group $P\overline{6}_3cm$ in hexagonal YMnO₃. Figure is reprinted from Katsufuji et.al [50].
Appendix B: Preparation of Y$^{3+}$ and Mn$^{3+}$ precursors for the citric acid method and gravimetrically determination of the precursors molality.

*Preparation of Y$^{3+}$ precursor*

The yttrium precursor was made by adding $\text{Y(O}_2\text{C}_2\text{H}_3)_3\times\text{H}_2\text{O}$ (0.05 mole) to a 1 L conical flask together with citric acid (CA) to a molar ratio of 1:15. Shortly after, 1 L of distilled water was added to the conical flask. The solution was placed on a hot plate holding a temperature of 150 °C, and continuously stirred (480 rpm) with a magnet. After 17 hours (overnight), all powders were dissolved and the solution was completely blank.

*Preparation of Mn$^{3+}$ precursor*

The manganese precursor was prepared by adding $\text{Mn(CO}_3)\times\text{H}_2\text{O}$ (0.05 mole) in a conical flask together with CA to a molar ratio of 1:22. The conical flask was shortly after added 1 L of distilled water. The same procedure for temperature and stirring was followed for the Mn$^{3+}$ precursor as for the Y$^{3+}$ precursor. The yellow solution became clear after about 17 hours.

*Standardization of precursors*

The molality of all solutions has been determined gravimetrically by heating four parallels of porcelain bowls each containing 2 ml of solution. A list of the two precursors prepared in this synthesis, their molality, their annealing temperature as well as the relative amount of citric acid to the cations is given in Table B1. Prior to heating, the porcelain bowls with lao wool lid, were heated at 1000 °C for 1 hour to avoid contamination and measurement wrongs. Figure B1 gives the temperature program for the precursors. The final products were $\text{Y}_2\text{O}_3$ and $\text{Mn}_2\text{O}_3$, as can be seen in the XRD diffractograms in Figure B2.

**Table B1:** A list over the content in the precursors, their molality as well as their annealing temperature, made for the citric acid synthesis route.

<table>
<thead>
<tr>
<th>Precursor</th>
<th>Weight of cation starting material [g]</th>
<th>Citric acid [g]</th>
<th>Relative amount [CA/ cations]</th>
<th>Molality [mmol/g]</th>
<th>Annealing temperature [°C]</th>
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**Fig. B1:** Temperature program used in determining the molality of the precursors gravimetrically.

**Fig. B2:** XRD diffractograms of Mn$_2$O$_3$ (black) and Y$_2$O$_3$. These oxides were formed when gravimetrically deciding the molality of the precursors used for the citric acid method. The vertical lines represent the matching PFD card (00-041-1442) for Mn$_2$O$_3$ (black), and PDF card (00-043-1036) for Y$_2$O$_3$ (red).
Appendix C: X-ray diffraction pattern of pure phase YMnO$_3$ and porous support P1 made from solid state synthesis.

The x-ray diffractogram of phase pure YMnO$_3$ within space group $P6_3cm$. The matching PDF card was 04-011-9577, as is shown as dotted red lines in Figure C1. The x-ray pattern for porous support P1 of YMn$_{0.85}$Ti$_{0.15}$O$_3$ is seen in Figure C2. A posteriori displacement correction of the data, reveals a displacement of the (004) reflection towards higher 2θ values, while the reflections from the (110), (111) and (112) planes are still displaced towards lower 2θ values.

![X-ray diffractogram of phase pure YMnO$_3$ prepared by solid state synthesis within space group $P6_3cm$.](image)

**Fig. D1**: X-ray diffractogram of phase pure YMnO$_3$ prepared by solid state synthesis within space group $P6_3cm$. The x-ray path was matched to the PDF card: 04-011-9577 belongs to hexagonal YMnO$_3$ and can be seen as the dotted red lines in the figure.
Fig. C2: X-ray diffractograms of the porous support (P1) of YMn$_{0.85}$Ti$_{0.15}$O$_3$ made by solid state synthesis that first was heated in air (green), before it was heated in N$_2$ (purple). The (-) indicates a secondary phase of Y$_2$O$_3$. The x-ray diffractograms is compared to pure YMnO$_3$ within space group $P6_3cm$ from general solid state sintering. An excerpt of the (110), (111), (004) and (112) reflections with a smaller range of 2θ values is inserted in the top-right corner of the figure to give a better view of the displacements with titanium substitution in YMnO$_3$ compared to pure YMnO$_3$. A posteriori displacement correction of the data, reveals a displacement of the (004) reflection towards higher 2θ values, while the reflections from the (110), (111) and (112) planes are still displaced towards lower 2θ values.
Appendix D: Rietveld refinement of YMnO$_3$ and YMn$_{0.85}$Ti$_{0.15}$O$_3$ (porous support P2) prepared by solid state synthesis, and nanocrystalline YMn$_{0.85}$Ti$_{0.15}$O$_3$ made by citric acid method.

The Rietveld refinement of YMnO$_3$ prepared by solid state synthesis within space group $P6_3cm$ is seen in Figure D1. In Figure D2 and D3, the Rietveld refinements of the porous support P2 (YMn$_{0.85}$Ti$_{0.15}$O$_3$) and nanocrystalline powder of YMn$_{0.85}$Ti$_{0.15}$O$_3$ are seen respectively. The porous support made by solid state synthesis was refined within space group $P6_3cm$, while nanocrystalline YMn$_{0.85}$Ti$_{0.15}$O$_3$ made by citric acid method is refined within space group $P6_3/mmc$.

![Figure D1: Rietveld refinement of the x-ray pattern for YMnO$_3$ made from solid state synthesis within space group $P6_3cm$.](image)
Fig. D2: Rietveld refinement of the x-ray pattern for the porous support P2 of YMn$_{0.85}$Ti$_{0.15}$O$_3$ made from solid state synthesis within space group $P6_3cm$.

Fig. D3: Rietveld refinement of x-ray pattern from nanocrystalline YMn$_{0.85}$Ti$_{0.15}$O$_3$ made by citric acid method within space group $P6_3/mmc$. 
Appendix E: X-ray diffractograms of yttrium deficient Y\text{MnO}_3 samples annealed in N\textsubscript{2} and O\textsubscript{2} atmospheres

X-ray diffractograms of yttrium deficient samples of Y\text{MnO}_3 heated in N\textsubscript{2} atmosphere, seen in Figure E1, and O\textsubscript{2} atmosphere seen in Figure E2.

**Fig. E1:** X-ray diffractograms of Y\text{MnO}_3 samples with varying yttrium deficiency as prepared by solid state synthesis and heated in N\textsubscript{2} atmosphere. The lower end of the diagram gives a better overview of the shift towards higher 2θ values for the reflections belonging to the (110), (111) and (112) planes. The vertical black lines in the lower end of the diagram is a guide to the eye to see the shift of intensities better. * is indicating the pronounced intensities of Mn\textsubscript{3}O\textsubscript{4} (hausmannite).
Fig. E2: X-ray diffractograms of YMnO$_3$ samples with varying yttrium deficiency as prepared by solid state synthesis and heated in O$_2$ atmosphere. The lower end of the diagram gives a better overview of the slight shift towards higher 2θ values for the reflection belonging to (112) plane. The vertical black lines in the lower end of the diagram is a guide to the eye to see the shift of intensities better. * is indicating the pronounced intensities of Mn$_3$O$_4$ (hausmannite).
Appendix F: Temperature program of the thermogravimetric analysis of YMnO$_3$ and Y$_{0.80}$MnO$_3$.

The temperature program used during thermogravimetric analysis of YMnO$_3$ and Y$_{0.80}$MnO$_3$ made by solid state synthesis is shown in Table F1.

**Tab. F1:** Temperature program of the thermogravimetric analysis of YMnO$_3$ and Y$_{0.80}$MnO$_3$ made by solid state synthesis.

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