SUBSTRATE EFFECT ON THE STRUCTURAL AND ELECTROCHEMICAL PROPERTIES OF ELECTROLYTIC MANGANESE DIOXIDE DEPOSITED FROM SULPHATE SOLUTIONS

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ABSTRACT

We studied the effect of anode substrates such as pure lead (Pb), lead antimony (Pb-Sb), and lead-silver (Pb-Ag) on the structural and electrochemical properties of electrolytic manganese dioxide (EMD). X-ray diffraction (XRD), field emission scanning electron microscopy (FESEM), and chemical analyses were used to determine the structural and chemical characteristics of the EMD samples. The charge–discharge profile was studied in 9 M KOH using a galvanostatic charge-discharge unit. In all the substrates the current efficiencies were more than 99% except with Pb-Sb where it was 90%. Results revealed the nature of the substrate strongly affected the morphology of the deposited material which in turn affected the electrochemical properties of the EMD samples. XRD analyses revealed that the nature of the anode did not affect the crystal structure of the deposited EMD and all the samples were predominantly $\gamma$-MnO$_2$, which is electrochemically active for energy storage applications. The EMD deposited on lead substrate showed superior discharge capacity of 245 mAhg$^{-1}$ when compared with other substrates.

KEYWORDS

Electrolytic manganese dioxide, anode substrate, morphology, discharges capacity.
INTRODUCTION

Use of manganese dioxide/Zn (MnO₂/Zn) batteries in a Laclanche cell is not new but after the improvement of the Laclanche cell for alkaline batteries, this system (MnO₂/Zn) conquered the market due to its low cost, good self life, high temperature performance and ecofriendly nature (Kordesch, 1960; Kordesch, 1978; Kordesch & Weissenbacher, 1994; Marsal, Kordesch, & Urry, 1960). Despite increasing competition from the rechargeable batteries, the primary Zn/Mn batteries, which were manufactured using electrolytic manganese dioxide (EMD) as essential raw material, were expected to remain dominant in the household markets for its relative convenience and low initial cost till today. EMD is prepared by anodic oxidation of manganese sulfate in sulfuric acid solution. The concentration of MnSO₄ and H₂SO₄ are taken in the ratio of 2:1 (Farris & Martin, 2004). Electrodeposition of manganese dioxide is influenced by various parameters such as concentration of electrolyte, current density, pH of the electrolyte and the temperature of the electrolytic bath, apart from this the anode material also play an important role in modifying the structure and properties of EMD. The selection of suitable anode material for the deposition of EMD has been briefly highlighted by Rethinaraj and Visvanathan (1991). It is reported that pure titanium is found to be very attractive anode for the preparation of electro inorganic chemicals, like EMD. Hence in terms of the economical point of view, an attempt is made to investigate the effect of lead based anode substrate on the structure and properties of EMD deposited from synthetic manganese sulphate solution.

EXPERIMENTAL

Materials and Methods for Producing EMD

Electrolytic manganese dioxide (EMD) was prepared from synthetic aqueous sulfate solutions containing 50 g dm⁻³ MnSO₄·7H₂O and 25 g dm⁻³ H₂SO₄ at an anodic current density of 200 A m⁻² in a glass cell. A schematic diagram of the electrolytic cell arrangement is shown in Figure 1. The anodic oxidation of Mn²⁺ to MnO₂ was carried out on various anodes (rolled) like lead (Pb), lead-antimony (Pb-Sb,(Sb-6%), or lead-silver (Pb-Ag)(Ag-1%) placed in parallel to a stainless steel (SS) cathode in their individual experiments. All experiments were carried out at 90-92°C for 6 h. EMD samples deposited on various substrates such as Pb, Pb-Sb and Pb-Ag have been labeled as EMD_Pb, EMD_Pb-Sb and EMD_Pb-Ag, respectively. The electrodeposited MnO₂ was removed from the anode after the deposition period and washed thoroughly with deionized water before drying in an oven. The dried mass was ground and sieved through a 100-μm mesh to obtain EMD powder. Subsequently, the resultant product in powder form was washed repeatedly with deionized water until the sample was sulfate free. The EMD powder was finally dried and cooled in a desiccator and subjected to physical and electrochemical characterization. Morphological characterization was performed on the EMD flakes, those scrapped off the anodes. These were washed with deionized water before analysis.

Structural Characterization

X-ray diffractograms were recorded for the EMD powders using PANalytical diffractometer (PW 1830; Philips, Japan) with Mo Kα radiation, λ = 0.71073 Å. The scans were recorded in 2θ range 5–45°. Field emission scanning electron microscope (FESEM) (ZEISS SUPRA 55) was used to examine the surface morphology of the EMD samples.

Electrochemical Characterization

Electrochemical characterization was carried out in a floated cell arrangement. The experimental cell consisted of a zinc strip as anode, and the cathode was in the form of a pellet, made from a uniform mixture of EMD and graphite powder (4:1), with 2–3 drops of 5 % polyvinyl alcohol as binder. The mixture was placed in a stainless steel mesh meant for electrical contact and then subjected to a pressure of 12 t for 3 min by means of a pelletiser (KBr press) in a 20 mm die. The pellet was put into a cell assembly in an electrolyte of 9M potassium hydroxide (KOH) and allowed to equilibrate for 1 h at its open circuit.
RESULTS AND DISCUSSIONS

Electrodeposition of Manganese Dioxide

Electrooxidation of MnO₂ was performed at an anodic current density of 0.02 A cm⁻² for 6 h from the electrolyte containing about 50 g dm⁻³ manganese with 25 g dm⁻³ of H₂SO₄. The electrodeposition of manganese dioxide from MnSO₄–H₂SO₄ solution can be represented by the following steps:

\[
\text{At Anode: } \text{Mn}^{2+} + 2\text{H}_2\text{O} \rightarrow \text{MnO}_2 + 4\text{H}^+ + 2\text{e}^- \quad (1)
\]

\[
\text{At Cathode: } 2\text{H}^+ + 2\text{e}^- \rightarrow \text{H}_2 \quad (2)
\]

\[
\text{Overall: } \text{Mn}^{2+} + \text{H}_2\text{O} \rightarrow \text{MnO}_2 + 2\text{H}^+ + \text{H}_2 \quad (3)
\]

As shown in Figure 2 for both EMD Pb and EMD Pb-Ag the current efficiencies (C.E.) are 99% whereas EMD Pb-Sb is 90%. Likewise, Figure 3 shows that the energy consumption (E.C.) is a minimum for EMD Pb (1.337 kWh/kg) in comparison to that of EMD Pb-Sb (1.788 kWh/kg) and EMD Pb-Ag (1.48 kWh/kg).
X-ray Diffraction Analysis

X-ray diffractograms of the EMD samples deposited on the three lead anode substrates are shown in Figure 4. All the observed diffraction peaks can be indexed to an orthorhombic phase of $\gamma$-MnO$_2$ with lattice constants $a = 8.70$ Å, $b = 2.90$ Å, and $c = 4.41$ Å. These are in good agreement with the standard values (JCPDS card no. 65–1298; $a = 9.27$ Å, $b = 2.87$ Å, $c = 4.53$ Å). The peaks at 2θ values of approximately 10, 16, 19, 24 and 29° correspond to the (120), (131), (300), (160), and (421) planes of $\gamma$-MnO$_2$, respectively. All three EMD samples (EMD$_{Pb}$, EMD$_{Pb-Sb}$, and EMD$_{Pb-Ag}$) were found to contain $\gamma$-phases, with good crystalline behavior. No significant difference is observed among the three types of EMD samples, hence it can be concluded that if electrolytic deposition conditions are constant, the crystal pattern of deposited EMD will be the same regardless the variable anode substrate.
Figure 4 - X-ray diffraction pattern of EMD material deposited on (a) Pb anode (b) Pb-Sb anode and (c) Pb-Ag anode.

Surface Morphology of the EMD

The three types of EMD samples (EMD\textsubscript{Pb}, EMD\textsubscript{Pb-Sb}, and EMD\textsubscript{Pb-Ag}) were investigated by FESEM. It is interesting to note that in all cases, the particle size was in the nano range (~50 to 100 nm), and the particles formed a net-like arrangement during electrodeposition. Although no differences were seen in the XRD for the EMD samples, their morphology appeared to be significantly different. The individual particles of EMD\textsubscript{Pb} (Figure 5a) obtained during electrodeposition consisted of star-shaped particles with four appendages arranged in a netlike fashion. A similar observation of synthesized \( \gamma \)-MnO\textsubscript{2} deposited electrolytically was recorded by Chen, Zhu, Han, Zheng, Yang, & Wang (2009). EMD\textsubscript{Pb-Sb} (Figure 5b) samples show spindle-shaped particles with consecutive occurrence of some agglomerated round shaped particles. However it is interesting to note that EMD\textsubscript{Pb-Ag} (Figure 5c) show spindle-shaped particles which are covered by another layer.

Electrochemical Activity

The suitability of the EMDs as battery materials prepared by varying the anode substrates was assessed by discharge behavior (Figure 6) through imposing a constant current over a period of time. MnO\textsubscript{2} as a half-cell versus metallic Zn was used. The samples were subjected to discharge studies in 9 M potassium hydroxide aqueous solutions. The reactions at cathode and anode are shown in reactions 4-6.

\[
\text{Cathode:} \quad \text{MnO}_2 + H_2O + e^- \rightarrow \text{MnOOH} + OH^- \quad (4)
\]
\[
\text{Anode:} \quad Zn + 2OH^- = ZnO + H_2O + 2e^- \quad (5)
\]
\[
\text{Net Reaction:} \quad Zn + 2\text{MnO}_2 + H_2O = ZnO + 2\text{MnOOH} \quad (6)
\]

The first electron of MnO\textsubscript{2} is allowed to discharge via the homogeneous reversible reaction by the movement of protons and electrons into the lattice, resulting in the reduction of MnO\textsubscript{2} to MnO\textsubscript{1.5} (Kordesch, Gsellmann, Peri, Tomantschger, & Chemelli, 1981; Minakshi, 2008; Minakshi, Singh, Carter, & Prince, 2008). This could be possible due to the conversion of MnO\textsubscript{2} into MnOOH in the solid phase. The second electron discharge of MnO\textsubscript{2}, proceeds either in solid or in solution phase, leads to the formation of Mn(OH)\textsubscript{2} containing soluble Mn (II) species, which is formed during recharging of \( \gamma \)-MnO\textsubscript{2} (Mondolini, Laborde, Rioux, Andoni, & Levy-Clement, 1992; Ruetschi, 1984). The formation of the
discharged products is reported to be nonreversible. This system is, therefore, suitable only as a use and dispose battery. Urfer, Lawrence, & Swinkles (1997) reported that the proton–electron pair formed during the discharge process diffuses from the EMD surface to the interior of the material. They maintained that the diffusion rate of the proton–electron pair is a function of the structural, physicochemical, and electrochemical properties of the EMD (Urfer et al., 1997). The theoretical discharge capacity of EMD has been reported as 308 mAh g\(^{-1}\) with regard to one electron discharge step (Li, Wang, He & Zhou, 2010). The discharge capacities versus cell voltage profile of the EMD samples are shown in Figure 6. EMD\(_{\text{Pb}}\) and EMD\(_{\text{Pb-Sb}}\) show discharge capacities of \(~245\) and \(~227\) mAh g\(^{-1}\), respectively, whereas EMD\(_{\text{Pb-Ag}}\) shows a discharge capacity of \(~215\) mAh g\(^{-1}\).

![Figure 5 - FESEM images of EMD samples (a)EMD\(_{\text{Pb}}\) (b)EMD\(_{\text{Pb-Sb}}\) (c)EMD\(_{\text{Pb-Ag}}\).]
CONCLUSIONS

The effect of the anode substrate on the structural and electrochemical behavior of the EMDs produced from synthetic manganese sulphate solutions were reported, and the following conclusions were drawn:

- All the samples show characteristics of $\gamma$-MnO$_2$ which is essential for their electrochemical activity and variation of the anode substrate did not affect the crystal phase of the deposited material.
- FESEM images show that the EMDs deposited on substrates have different surface morphologies, indicating a significant effect of the substrate interface on the shapes and sizes of the EMD particles.
- Tetra-branched star shaped nano particles were obtained for EMD$_{Pb}$, spindle shaped nano particles with frequent appearance of round shaped particles for EMD$_{Pb-Sb}$, and needle like growth of nano particles covered with another layer for EMD$_{Pb-Ag}$ were also seen.
- EMD$_{Pb}$ shows superior discharge capacity of 245 mAh$^{-1}$ against the discharge capacity of 227 and 215 for EMD$_{Pb-Sb}$ and EMD$_{Pb-Ag}$ respectively.

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