Basic Medium Heterogeneous Solution Synthesis of \( \alpha \)-MnO\(_2\) Nanoflakes as an Anode or Cathode in Half Cell Configuration (vs. Lithium) of Li-Ion Batteries

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Abstract: Nano \( \alpha \)-MnO\(_2\) is usually synthesized under hydrothermal conditions in acidic medium, which results in materials easily undergoing thermal reduction and offers single crystals often over 100 nm in size. In this study, \( \alpha \)-MnO\(_2\) built up of inter-grown ultra-small nanoflakes with 10 nm thickness was produced in a rapid two-step procedure starting via partial reduction in solution in basic medium subsequently followed by co-proportionation in thermal treatment. This approach offers phase-pure \( \alpha \)-MnO\(_2\) doped with potassium (cryptomelane type \( K_{0.25}Mn_8O_{16} \) structure) demonstrating considerable chemical and thermal stability. The reaction pathways leading to this new morphology and structure have been discussed. The MnO\(_2\) electrodes produced from obtained nanostructures were tested as electrodes of lithium ion batteries delivering initial discharge capacities of 968 mAh g\(^{-1}\) for anode (0 to 2.0 V) and 317 mAh g\(^{-1}\) for cathode (1.5 to 3.5 V) at 20 mA g\(^{-1}\) current density. At constant current of 100 mA g\(^{-1}\), stable cycling of anode achieving 660 mAh g\(^{-1}\) and 145 mAh g\(^{-1}\) for cathode after 200 cycles is recorded. Post diagnostic analysis of cycled electrodes confirmed the electrode materials stability and structural properties.

Keywords: \( \alpha \)-MnO\(_2\); nanostructured matrix; base medium fabrication; electrode materials; alkali metal ion battery

1. Introduction

Lithium ion batteries (LIB) have attracted huge attention in the battery industry since their introduction in the early 1990s by Sony. The growing attention to large intermittent devices and electric vehicles requires high energy and high power density batteries. Thereby, developing LIB with good performance and excellent stability becomes essential. Metal oxides are studied as electrode materials for LIB due to their high theoretical capacity compared to graphite [1–4]. Among various transition metal oxides, manganese-based oxides (MnO, Mn\(_2\)O\(_3\), Mn\(_3\)O\(_4\), and MnO\(_2\)) have attracted huge interest as electrode materials for their high theoretical capacity, eco-friendliness, cost-effectiveness, and natural abundance [5]. Based on the conversion reaction of each manganese oxide, the theoretical capacity is 750, 936, 1018, and 1230 mAh g\(^{-1}\) for MnO, Mn\(_3\)O\(_4\), Mn\(_2\)O\(_3\), and MnO\(_2\), respectively. Traditionally, MnO\(_2\) is applied as a material for primary lithium batteries and capacitors [6,7]. Recently, the interest in lithium-ion batteries has increased as well for their high theoretical capacity among four different types of manganese oxides [8,9].
However, MnO\(_2\) as an electrode material in secondary ion batteries has been hindered by the large volume expansion, poor conductivity, and poor reversibility [5,8]. To address such an issue, several strategies have been suggested including MnO\(_2\) composite material and fabricating nano-structured MnO\(_2\). Li et al. wrapped MnO\(_2\) with graphene to resolve the conductivity issue [10]. The initial capacity was 1200 mAh g\(^{-1}\) and a stable capacity of 600 mAh g\(^{-1}\) at 400 mA g\(^{-1}\) current density was maintained. MnO\(_2\) composite with improved conductivity has been realized by using graphene [11], carbon nanotube [12], and other metal oxides [13–15]. The size, crystallinity, and morphology control of MnO\(_2\) are important factors that influence the performance. Thereby, it is essential to choose the proper synthesis method. Numerous reports use the acidic medium fabrication method to prepare MnO\(_2\). Chen et al. have prepared MnO\(_2\) nano-rods using KMnO\(_4\) and HCl via the hydrothermal method [8]. The prepared nano-rods have shown an initial capacity of 1200 mAh g\(^{-1}\). However, the cycle performance was unstable. Another example of using acidic medium in hydrothermal process was reported by Feng and coworkers [16]. Despite the high initial capacity (1400 mAh g\(^{-1}\)), the capacity drops to 600 mAh g\(^{-1}\) at second cycle. Li and coworkers prepared hollow urchin-like MnO\(_2\) via mixture of KMnO\(_4\) and sulfuric acid [17]. The hollow urchin morphology is obtained by the Ostwald ripening and delivered an initial capacity of 746 mAh g\(^{-1}\). Still, the cycle retention was 64% after 40 cycles. In addition to anode, there have also been few works on MnO\(_2\) as cathode material reported. 

The majority of research used acidic medium for fabricating MnO\(_2\). Especially, hydrothermal approaches in acidic medium for preparation of the MnO\(_2\) are well reported [18,19]. In realization of this approach, a broad variety of precursors and conditions have been evaluated. In particular, acidic leaching of Mn(II) with H\(_2\)SO\(_4\) from Mn\(_2\)O\(_3\) has been reported [20,21]. Even decomposition of KMnO\(_4\) solutions in acidic medium at controlled low pH has been investigated [22]. The most thoroughly investigated tracks have been, however, the reduction of KMnO\(_4\) by HCl or by MnSO\(_4\) in acidic medium [23,24]. In addition, the use of organic polymers for permanganate reduction has been investigated [25,26]. Other approaches involved oxidation of MnSO\(_4\) solutions by potassium persulfate, K\(_2\)S\(_2\)O\(_8\), or by ozone [16]. Oxidation by atmospheric oxygen has also been used in the co-precipitation approach in basic medium that was using MnSO\(_4\) and MnC\(_2\)O\(_4\) as Mn-sources. Despite acidic medium based MnO\(_2\) showed promising results, acidic medium results in partial reduction of Mn(IV) in the produced MnO\(_2\) nanostructures, leading potentially to lower capacity and instability in cycling of the produced electrode materials.

In this work, we report a successful fabrication of \(\alpha\)-MnO\(_2\) nano flakes by reduction approaches in basic medium, securing preservation of the oxidation state +4. An unusual approach based on seeding of \(\alpha\)-MnO\(_2\) through thermal treatment-induced interaction of amorphous MnO\(_2\), precipitated from permanganate by ammonia, with fine powder of stoichiometric amounts of Mn(II) carbonate resulted in interpenetrated network of phase-pure nanocrystalline material. Table 1 shows the comparison of current work to other’s work on MnO\(_2\) as electrode material. The structure analysis showed successful fabrication of nano-MnO\(_2\) particles. In addition, the preservation of MnO\(_2\) has been observed by Fourier Transformed Infrared Spectroscopy (FTIR) and Thermogravimetric Analysis (TGA). Basic medium based \(\alpha\)-MnO\(_2\) showed successful results as electrode material for LIB and revealed a stable capacity of 690 mAh g\(^{-1}\) and 150 mAh g\(^{-1}\) at 100 mA g\(^{-1}\), respectively. The behavior of \(\alpha\)-MnO\(_2\) was discussed by the voltage profile and cyclic voltammetry curves. To confirm that basic medium fabrication is compatible to conventional acidic medium based MnO\(_2\), the capacity values are compared.
Table 1. Comparison of current work to literature reports on MnO₂ in lithium ion batteries.

| References       | Synthesis Method                     | Initial Capacity (Current Density) | Cycle Retention | Rate Capacity          | Electrode |
|------------------|--------------------------------------|------------------------------------|-----------------|------------------------|-----------|
| Chen [8]         | KMnO₄ + HCl Hydrothermal             | 1119 mAh g⁻¹ (100 mA g⁻¹)          | 96% (100 cycles)| 1000 mAh g⁻¹ (200 mA g⁻¹) | Anode     |
| Vernardou [1]    | KMnO₄ + Nitric Acid Hydrothermal     | 943 mAh g⁻¹                         | N/A             | N/A                    | Anode     |
| Walanda [20]     | Hydrothermal of Mn₂O₃ in H₂SO₄       | ~1100 mAh g⁻¹ (20 mA g⁻¹)           | N/A             | N/A                    | Anode     |
| Li [17]          | MnSO₄ + H₂SO₄ Hydrothermal           | 750 mAh g⁻¹ (270 mA g⁻¹)            | 64% (40 cycles) | N/A                    | Anode     |
| Dai [18]         | MnSO₄ + H₂SO₄ Ozone Oxidation        | 210 mAh g⁻¹ (20 mA g⁻¹)             | 82.5% (20 cycles)| N/A                    | Cathode   |
| Abuzeid [5]      | KMnO₄ + H₂SO₄ green-like synthesis method | 236 mAh g⁻¹ (20 mA g⁻¹)            | 73% (20 cycles) | 75 mAh g⁻¹ (200 mA g⁻¹) | Cathode   |
| Our Study        | KMnO₄ + NH₄OH Stirring and precipitation | 800 mAh g⁻¹ (50 mA g⁻¹)            | 91% (200 cycles)| 600 mAh g⁻¹ (200 mA g⁻¹) | Anode     |
|                  |                                      | 250 mAh g⁻¹ (50 mA g⁻¹)            | 93% (200 cycles)| 100 mAh g⁻¹ (200 mA g⁻¹) | Cathode   |
2. Experimental

2.1. Synthesis of MnO₂

All chemicals purchased from Sigma-Aldrich (Copenhagen, Denmark) were used without further purification. The synthesis was conducted under an ambient atmosphere. For preparation of α-MnO₂ nano flakes, 0.2 g KMnO₄ (1.28 mmol) was dissolved in 10 mL deionized water and a slurry of 0.5068 g of MnCO₃·H₂O (3.84 mmol) in 15 mL 25% NH₃ solution was added. Constant stirring for 1 h at 80 °C were applied and then additional 5 mL 25% NH₄OH were added and continued stirring for 1 h. The resulting brown precipitate was collected by decantation, washed and centrifuged with three portions of 10 mL of distilled water and dried in air overnight followed with heat treatment at 400 °C in air (heating up with the rate of 10 °C/min to 400 °C from room temperature and keeping then at this temperature for 4 h. The time was optimized experimentally by ex-situ X-ray Diffraction (XRD, Stockholm, Sweden) studies after different annealing times—2, 3, 4 and 6 h). The yield of resulting greyish black powder is 0.3781 g (86%).

2.2. Material Characterization

XRD studies of the powders were carried out with D8 SMART CCD Apex-II diffractometer (Bruker Nordic, Stockholm, Sweden) in rotation mode. SEM energy-dispersive X-ray spectroscopy (EDS) investigation was performed using Hitachi TM-1000-μ-DEX instrument (Spectral Solutions AB, Stockholm, Sweden). TEM investigation was made with Phillips CM12 instrument (Philips, Sweden). AFM studies have been performed with Bruker FastScan with ScanAsyst microscope (Blue Scientific, Stockholm, Sweden). FTIR studies were made with Spectrum 100 instrument (Perkin-Elmer, Upplands Väsby, Sweden). Thermogravimetric investigation was carried out with Pyris-1 thermobalance (Perkin-Elmer, Upplands Väsby, Sweden).

2.3. Electrochemistry

The electrochemical properties of α-MnO₂ electrode materials were tested in half cells configuration with lithium-metal as counter-electrode as an anode (loading mass: 0.3 mg cm⁻²) and cathode (loading mass: 0.3 mg cm⁻²). 70% of prepared α-MnO₂ was mixed with 20% conducting carbon (Timcal Super P, MTI Corporation, Richmond, CA, USA) and 10% PVDF (MTI Corporation, Richmond, CA, USA) binder in N-methyl-2-pyrrolidone (NMP) solvent. Obtained slurry was coated on aluminum (for cathode) and copper (for anode) current collectors. After drying overnight in a vacuum oven set at 80 °C, the prepared working electrode is assembled into 2032 coin cells (CR 2032) inside a glove box. Lithium foil (MTI Corporation, Richmond, CA, USA) was used as a counter-electrode, Cellgard 2500 (Charlotte, NC, USA) as a separator, and 1 M lithium hexafluorophosphate (LiPF₆) salt in ethylene carbonate (EC)/diethyl carbonate (DEC) (Sigma Aldrich, St. Louis, MO, USA, 1:1 volume ratio) as an electrolyte. Galvanostatic charge-discharge profile was measured in the potential range of 0.01 V to 2 V for anode and 1.5 V to 3.5 V for cathode (BST8-MA, MTI, Richmond, CA, USA). In addition, cyclic voltammetry was performed by Gamry Reference 600+ equipment (Warminster, PA, USA) at a scanning rate of 0.1 mV s⁻¹.

3. Results and Discussion

3.1. Synthesis and Structure of the Material

In preparation of manganese dioxide in basic medium, the reduction of potassium permanganate with concentrated ammonia was conducted. The proposed route involves setting together a solid phase, MnCO₃(s), together with the solution of KMnO₄ and NH₃(aq). The reaction in the liquid phase provides amorphous KₓMnO₂₋ₓY (and N₂(g) nitrogen), with oxidation state for Mn higher than +IV. In contact with MnCO₃, this results apparently in seeding of MnO₂(s) on the surface of the carbonate crystals. In subsequent thermal treatment, the whole mass is transformed into small-size α-MnO₂.
crystallites, both the amorphous part crystallizes with release of adsorbed water and breakdown of surface hydroxide groups and the solid MnCO$_3$ is broken down with release of CO$_2$. It is noteworthy that the thermal treatment of MnCO$_3$ only gives as revealed by XRD in this work a complex mixture of Mn$_2$O$_3$, Mn$_3$O$_4$ and MnO$_2$ (see Supplementary Figure S1) and also the heat-treatment of the amorphous mass, resulting from reduction of KMnO$_4$ by NH$_3$ on reflux in concentrated solution offers a complex mixture of phases after heat-treatment at over 600 °C (Figure S2). It is the combination of the two systems that results in practically phase-pure MnO$_2$. Addition of a stoichiometric amount of manganese (II) carbonate, MnCO$_3$, provides the necessary reducing agent not causing inclusion of other transition metal cations [27].

The product obtained from the initial reaction step in solution revealed a low-intensity X-ray diffraction pattern corresponding to MnCO$_3$ together with an amorphous phase (see Supplementary Figure S3). Subsequent thermal treatment in 2 h at 400 °C led to a practically amorphous material according to XRD, while further treatment in 2 h resulted in a crystalline sample with XRD pattern corresponding to α-MnO$_2$ phase (Figure 1). The average crystallite size calculated from Scherrer equation using the most intense line at 2θ = 16.987° (with Miller indices hkl = 211) is 108 nm. This is as stated only an average value as both smaller and bigger crystallites (over 100 nm) could be observed by TEM (see Figure 2). The peak at 2θ = 14.707° is corresponding to minor impurity of the residual MnCO$_3$ (the most intense hkl = 104 line).

The SEM images (see Figure 2A,B) show strongly aggregated small particles forming low mass density solid material. The morphology of the prepared MnO$_2$ revealed by TEM shows a flake-like structure with 0.491 nm 200 plane fringe pattern, which agrees to the Cryptomelane type α-MnO$_2$ structure (Figure 2D). Low-charge cations are required to stabilize the Cryptomelane MnO$_2$ structure—the formula of the natural Cryptomelane mineral is KMn$^{IV}_7$Mn$^{III}_{16}$O$_{16}$ [28]. The α-MnO$_2$ obtained in this work is apparently stabilized by the minor residual content of potassium in its composition. The Mn to K ratio was determined by EDS analysis and was found to be Mn:K = 32:1, i.e., only 25% of the additional positions in the Cryptomelane structure are thus occupied by the K-cations in the classic mineral structure, which here can be formulated as K$_{0.25}$Mn$_8$O$_{16}$.

The flakes are connected in a very open interpenetrated network (Figure 2C) that is apparently formed through the thermal decomposition process associated with evolution of gases—both CO$_2$ from MnCO$_3$ crystals and water vapor from K-containing amorphous hydrated precipitate surrounding the carbonate crystals in the initial product of solution reaction. The process is straightforward and easy to carry out. The Brunauer-Emmett-Teller (BET) analysis was conducted to further confirm the interconnected structure (Figure S4). The pore distribution profile shows existence of various pore size in the prepared materials. This is attributed to the pore existing between interconnected MnO$_2$
nanoparticles. In addition, the surface area (61.5 m$^2$ g$^{-1}$) is relatively higher than reported bulk MnO$_2$ nanoparticles, which supports the interconnected structure as well [5,21].

![Figure 2. SEM (A,B) and TEM (C,D) images and EDS analysis of the α-MnO$_2$ obtained (E).](image)

Direct confirmation to the proposed mechanism could be obtained in the high resolution TEM and AFM studies of the intermediate reaction mixtures. As can be seen in Figure 3, the amorphous flakes containing K-cations and produced by decomposition of KMnO$_4$ in the presence of ammonia commence to crystallize forming nuclei of the Cryptomelane type phase. The crystallites of MnCO$_3$ are turned amorphous by the thermal treatment but keep their shape. The AFM images (Figure 3B,C) show clearly the rhombic shape blocks derived from the initial MnCO$_3$ crystallites embedded into a less dense matrix. The final material is produced by crystallization accompanied with diffusion of potassium into these bigger blocks as the final material is single phase and has uniform potassium distribution in it according to EDS. The overall mechanism of the process can thus be described by Scheme 1.

The occurring chemical transformations can be described by the following reaction equations:

\[
3\text{KMnO}_4(aq) + (4 - 2x)\text{NH}_3(aq) \rightarrow 3\text{KMnO}_{2+x}(s) + (2 - x)\text{N}_2(g) + 6\text{H}_2\text{O} \quad (1)
\]

\[
\text{MnCO}_3(s) \rightarrow \text{MnO}_{2-y} + \text{CO}_2(g) \quad (2)
\]

\[
\text{KMnO}_{2+x}(s) + 3\text{MnO}_{2-y}(s) \rightarrow K_{0.25}\text{MnO}_2(s) \quad (3)
\]
Figure 3. TEM (A) and AFM (B,C) images of the material obtained in the course of thermal transformation of the reaction mixture into $\alpha$-MnO$_2$.

Scheme 1. Reaction pathway in producing hollandite type $\alpha$-MnO$_2$ phase.

3.2. Stability Test of Produced $\alpha$-MnO$_2$

FTIR (Figure S5) shows the chemical information of samples kept at ambient condition. Minor degree of hydration ($\delta$(O–H) at about 1600 and $\nu$(O–H) at 3700 cm$^{-1}$) and the network of metal-oxygen bonds present as characteristic bands (around 536 and 585 cm$^{-1}$).

The TGA (Figure S6) shows the loss of hydrating water starting at room temperature and finishing at about 200 °C with the loss of 2.4–2.7% of the total weight in both air and nitrogen atmosphere. This process is followed by thermal decomposition with reduction of MnO$_2$ into Mn$_3$O$_4$ in air at 500–620 °C (corresponding weight loss 6.1%) and into first Mn$_3$O$_4$ at 470–560 °C (weight loss of about 6%) and then Mn$_2$O$_3$ at 660–800 °C (further weight loss of about 3%) in N$_2$ atmosphere. This confirms that MnO$_2$ was produced when the heating was carried out to 400 °C with subsequent annealing and shows considerable thermal stability of the obtained material.

3.3. Electrochemical Performance

The $\alpha$-MnO$_2$ electrode was assembled and tested as anode and cathode separately vs. lithium (half cells) for lithium ion batteries. Figure 4A shows the voltage profiles of MnO$_2$ for the initial two cycles at a current density of 20 mA g$^{-1}$. It exhibits an initial discharge and charge capacity as 2369 and 968 mAh g$^{-1}$, respectively, with a 59.1% first cycle Coulombic efficiency. The loss of capacity at the first cycle is attributed to the irreversible conversion reactions of lithium and MnO$_2$ and the formation of solid electrolyte interphase (SEI) layer, which is obvious by the first cycle voltage
profile slope and plateau in 1.5 to 0.7 V range [8]. The cyclic voltammetry (CV) was conducted to understand the kinetics (Figure 4C). The initial two cycles are scanned at a range of 0.01 to 2.0 V at a 0.1 mV s$^{-1}$ scan rate. The CV curve for the first cycle shows a cathodic peak near 1.5 V which is not observed at the second cycle. The SEI layer and the irreversible conversion reaction between lithium and MnO$_2$ shows the observed cathodic peak [16]. In addition, a broad peak appearance from 1.5–0.5 V corresponds to the reduction of MnO$_2$ to Mn$^{2+}$, and the peak near 0.1 V corresponds to the reduction of Mn$^{2+}$ to metallic Mn$^0$, which is easily found in other transition metal compounds for lithium ion batteries. An anodic peak appearance at 1.30 and 2.0 V is a response to the oxidation of Mn$^0$ and Mn$^{2+}$, respectively [8]. A shift in the second cycle explains a possible structure change induced by the formation of metallic manganese.

Figure 4. Electrochemical data of $\alpha$-MnO$_2$ anode electrode. (A) charge-discharge profiles for first two cycles, (B) rate study, (C) cyclic voltammetry curve, and (D) constant current (100 mA g$^{-1}$) cycling.

Figure 4B explains the rate performance of MnO$_2$ as an anode at different current density. The rate study was conducted at various current densities, from 50 mA g$^{-1}$ to 500 mA g$^{-1}$. Stable capacities of 791 mAh g$^{-1}$ and 517 mAh g$^{-1}$ at current densities 50 mA g$^{-1}$ and 500 mA g$^{-1}$, respectively is obtained. The comparison of initial and final capacities at current densities 100 mA g$^{-1}$ (711 mAh g$^{-1}$ vs. 716 mAh g$^{-1}$) shows excellent rate recovery after high rate performance. The cycle retention is 91% over 200 cycles (Figure 4D, 0.045% per cycle) which is relatively stable compared to other reports on MnO$_2$ as anode (Table 1).

Numerous reports also focused on the possibility of using acid medium fabricated $\alpha$-MnO$_2$ as cathodic materials. To further ascertain that basic medium fabrication produced $\alpha$-MnO$_2$ is successful, its cathodic activities are tested at higher potential and compared (Table 1). The charge-discharge profile at a current density of 20 mA g$^{-1}$ displayed an initial discharge and charge capacity as 477 and
317 mAh g⁻¹, respectively, with a 33.4% capacity drop (Figure 5A). A plateau region occurring at a broad range over 2.5 to 2.2 V is attributed to the reduction of Mn⁴⁺ ions. The cyclic voltammetry (CV) shows analogous result to voltage profile, exhibiting an anodic peak at a similar range (oxidation of Mn²⁺ to Mn⁴⁺) with voltage profile (Figure 5C). The discharge capacity of 171 mAh g⁻¹ recovered to 155 mAh g⁻¹ after rate study (91% recover for cathode). The low capacity number at high current density can be explained by the sluggish kinetics of Li ion. Numerous reports compared the influence of Li ion conductivity on rate performance, showing low conductivity results in low rate performance [29,30]. The cycle retention of cathode is measured and displayed in Figure 5D. The capacity retention is 93% over 200 cycles (0.035% decrease per cycle).

Post-diagnostic analysis was carried out to determine whether the basic medium synthesized α-MnO₂ is stable after long-term cycling (200 cycles). SEM and Raman analysis were performed on before and after cycled electrodes (both anode and cathode). The as-prepared electrode is composed of α-MnO₂ (active material, conductive carbon (Super P), and PVDF binder (Figure 6A). It is observed that a uniform, homogeneous electrode is fabricated using α-MnO₂ powder. The SEM image of cycled anode (Figure 6B) and cathode electrode (Figure 6C) shows that there is no obvious physical change in laminate. Raman spectroscopy of pristine electrode shows a peak positioned at 650, 1330, and 1598 cm⁻¹, indicating presence of α-MnO₂ (650 cm⁻¹) and super P (conductive carbon), respectively [31–33]. For both cycled electrodes, the new Raman band appeared and the peak shows a broad feature compared to pristine electrode Raman peak (Figure 6D). The broad peak can be explained by the partial disordered phase of MnO₂ after lithiation and delithiation process. After cycling, new Raman band appears at 484 and 430 cm⁻¹, which agrees to reported band position of Li-birnessite in Raman [34]. The similarity of Raman spectra suggests that during the
repeated lithiation and delithiation, the hollandite MnO$_2$ structure partially transformed to layered birnessite-type Li-MnO$_2$ [35,36].

![Image](image_url)

Figure 6. SEM images of (A) un-cycled electrode, (B) anode cycled for 200 cycles, (C) cathode cycled for 200 cycles, and (D) Raman spectra of each electrode.

4. Conclusions

The newly proposed approach, combining solution redox reaction and seeding on the surface of introduced MnCO$_3$ solid, offers a facile approach to α-MnO$_2$ phase stabilized by residual potassium content and shaped as nano flakes. Compared to acidic medium synthesis, this study shows advantages in, on one hand, operating with a phase-pure Mn(IV) derived material, and, on the other hand, exploiting a nanostructure of small particles with an average size of 10 nm, connected into an interpenetrated network created by the thermal decomposition synthesis associated with evolution of gases. A stable capacity of 791 and 243 mAh g$^{-1}$ at 50 mA g$^{-1}$ were obtained for α-MnO$_2$ tested as anode and cathode, respectively. The rate performance was stable as well, showing 718 and 155 mAh g$^{-1}$ at 100 mA g$^{-1}$ for anode and cathode, respectively. From this study, we demonstrated that MnO$_2$ obtained from basic medium is a promising electrode material for anode as well as cathode. This method can be further utilized in preparing materials that are sensitive to acid medium.

Supplementary Materials: The following are available online at http://www.mdpi.com/2079-4991/8/8/608/s1, Figure S1: XRD pattern of decomposition products of MnCO$_3$ with characteristic peaks for Ramsdellite MnO$_2$ (ICPDS Card 82-2169), Mn$_3$O$_6$ (39-1218), Hausmannite Mn$_3$O$_4$ (80-382), Akhtenskite-MnO$_2$ (30-0820) and Rhodochrosite MnCO$_3$ (86-0173) marked in the image, Figure S2: XRD of the material obtained by heat-treatment of the precipitate from the reaction KMnO$_4$ with NH$_3$ in aqueous medium at 600 °C in 2 h in air with characteristic peaks indicated for K$_{0.5}$Mn$_2$O$_{4.5}$H$_2$O (ICPDS Card 42-1317), Crypromelane KMn$_{8}$O$_{16}$ (34-0168) and K$_6$Mn$_2$O$_6$ (ICDD PDF 01-070-1271), Figure S3: XRD of the initial product of solution synthesis featuring the pure MnCO$_3$ phase ICPDS Card No. 86-0173, Figure S4: BET analysis of (a) N2 absorption isotherm and (b) pore size distribution profile of prepared α-MnO$_2$ nanomaterial, Figure S5: FTIR analysis for produced α-MnO$_2$ nanomaterial, Figure S6: TGA (solid line) and DTG (dashed line) analysis of produced α-MnO$_2$ nanomaterial in the air (A) and in the nitrogen (B) atmosphere.

Author Contributions: G.A.S. designed the synthetic procedure, carried out the synthesis and characterization (SEM, AFM, FTIR, TGA) of oxide materials, and contributed to writing of the manuscript, being one of
the corresponding authors, K.H. prepared the electrodes, performed the electrochemical measurements, post-cycled battery analysis (Raman and SEM), and contributed to writing, G.D. carried out the TEM studies, V.G.K. contributed to discussion especially concerning the formation mechanism, V.G.P. has initiated the study, made strong contribution to writing and was acting as corresponding author.

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