Facile Synthesis of Mn_{0.68}Bi_{0.32}OCl Mix-Crystals And Its Supercapacitive Behavior

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Research Article

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Abstract

Mn$_{0.68}$Bi$_{0.32}$OCl mix-crystals for supercapacitor were successfully synthesized via a facile solid-phase method using Bi(NO$_3$)$_3$ and MnCl$_2$ with molar ratio of 1:1 as precursors. The Mn$_{0.68}$Bi$_{0.32}$OCl mix-crystals were characterized by scanning electron microscopy, X-ray diffraction, Brunauer-Emmett-Teller surface area measurements and thermogravimetry and differential scanning calorimetry, respectively. Cyclic voltammetry and galvanostatic charge/discharge technique were performed for the Mn$_{0.68}$Bi$_{0.32}$OCl mix-crystals in 1 M Na$_2$SO$_4$ aqueous solutions; the specific capacitance of Mn$_{0.68}$Bi$_{0.32}$OCl was about 203 F.g$^{-1}$ at the current density of 3 A. g$^{-1}$ with a long life time, owing to the high power density of Mn$_{0.68}$Bi$_{0.32}$OCl mix-crystals and the higher surface area, good conductivity, and high stability of the Mn$_{0.68}$Bi$_{0.32}$OCl mix-crystals.

1. Introduction

Supercapacitor as a next generation of promising electrochemical energy storage device has the advantages of fast charge and discharge speed, long service life, high power density, high safety and good cycle stability. Based on the charge storage mechanism, supercapacitors can be divided into double-layer capacitors, pseudo capacitors and hybrid supercapacitors. The two-layer capacitor consists of activated carbon electrode material, and pseudocapacitive supercapacitors include metal oxide electrode materials and polymer electrode materials. Mn and Bi are environmentally friendly and inexpensive, widely used in the preparation of supercapacitors, and the Mn$_3$O$_4$ [1], MnO [2], MnOx [3], Mn$_2$O$_3$ [4], Bi$_2$O$_3$ [5–8], BiOCl [9], (Cu)BiOCl [10], and BiOBr [11] supercapacitors have been prepared for supercapacitors.

BiOCl as a semiconductor material was often widely used in the field of photocatalysis [12–13] due to its special layered structure, which is composed of [Bi$_2$O$_2$]$_{2+}$ layer and double-layer halogen ion layer staggered along the [001] direction, and its synthesis mainly include hydrothermal / solvothermal method, template method, ultrasonic method, solid-phase method and sol-gel method [14–19]. The band gap of BiOCl is 3.6 eV, the ion doping was a effective for improving the catalytic performance of BiOCl [20], the different molar ratio manganese-doped BiOCl composites (Mn/BiClO) were prepared using Bi (NO$_3$)$_3$ and Mn(CH$_3$COO)$_2$ as precursors by the hydrothermal method with imidazole hydrochloride [HMIM]Cl as the chlorine source and template for the photocatalytic degradation of rhodamine [21], Cao et al. synthesized the Mn$^{3+}$ doping BiOCl in hydrochloric acid and potassium chloride solution using Bi (NO$_3$)$_3$ and MnCl$_2$ with molar ratio of 1:0.015 as the precursors by hydrothermal method for the metronidazole photodegradation [22], the Bi(Mn)OCl could also be synthesized using hydrothermal method by adjusting the pH of MnCl$_2$-BiCl$_3$-HCl solution with ammonia water for the degradation of malachite green dye [23].

In present work, the Mn$_{0.68}$Bi$_{0.32}$OCl was synthesized in one step using Bi (NO$_3$)$_3$ and MnCl$_2$ with molar
electrochemistry, and its supercapacitive behavior was studied.

2. Experimental

2.1. Materials

60% polyvinylidene fluoride (PVDF) water emulsion was provided by Shanghai San-Ai-Fu new material Ltd (Shanghai, China). N. N-dimethylformamide, Na$_2$SO$_4$, Bi (NO$_3$)$_3$·5H$_2$O and MnCl$_2$·4H$_2$O were purchased from Shanghai Analytical Chemicals Company. All chemicals were of analytical grade without further purification. Graphite electrode was used as active material carrier because of its low price, good adsorb ability and electrochemical stability. At the same time, graphite was easy to make flexible electrode. The graphite electrode was prepared by the following method: the graphite electrode of a diameter of 14.0 mm was inserted into the polypropylene-random (PPR) plastic pipe of an inner diameter of 14.00 mm, the surface of graphite electrode was polished using sandpaper of 1000 and 7000 mesh to the mirror, respectively, then further cleaned with polishing cloth, and the other end was connected the with a copper column. The prepared graphite electrode was sonicated with deionized water and ethanol for 30 minutes, respectively, and dried for standby.

2.2. Synthesis of Mn$_{0.68}$Bi$_{0.32}$OCl

0.01M Bi (NO$_3$)$_3$·5H$_2$O and MnCl$_2$·4H$_2$O were dissolved with 0.01 M HNO$_3$ of 30 ml in a porcelain crucible and heated in a 100 °C oven for 24 hours for nearly dry, then the mixture was transferred into a muffle furnace, heated from room temperature to 500 °C for 0.5 h, kept at 500 °C for 4 h, from 500 to 600 °C for 0.5 h, kept 600 °C for 1 h, and cooled to room temperature for use.

2.3. Material characterization

The nitrogen adsorption and desorption experiments of Mn$_{0.68}$Bi$_{0.32}$OCl were carried out at 77 K using SA3100 surface area and pore size analyzers (Beckman Coulter, Inc. USA). The morphologies of Mn$_{0.68}$Bi$_{0.32}$OCl were examined by scanning electron microscope (SEM) (QUANTA FEG 450, USA), equipped with an EDAX OCTANE PRO energy dispersive spectrometer (EDS) (FEI, USA). The thermogravimetry (TG) and differential scanning calorimetry (DSC) were performed using a NETZSCH STA 449F3 simultaneous thermal analyzer (German). X-Ray diffraction (XRD) analysis was performed on the Mn$_{0.68}$Bi$_{0.32}$OCl with a Switzerland ARL X’TRA X-ray diffractometer rotating anode with Cu-Kα radiation source (λ = 0.1540562 nm).

2.4. Electrochemical characterization

To investigate the supercapacitive behavior of the nanomix-crystals, the active material, carbon black (CB) and PVDF were taken in the weight ratios of 100 : 10 : 10. The mixture of Mn$_{0.68}$Bi$_{0.32}$OCl of 10.0 mg, CB of 1.0 mg and PVDF of 1.0 mg was firstly dispersed in N. N-dimethylformamide of 10.00 ml, ultrasound for 30 minutes, then the mixture of 100 µl evenly dispersed on the surface of the clean loading mass of active material was 0.10 mg.
Land-CT2001A battery analyzer (Wuhan, China) was utilized for the charge/discharge cycling life tests of 
Mn$_{0.68}$Bi$_{0.32}$OCl. The electrochemical characterization of the prepared capacitive electrodes was also
carried out with a CHI660e electrochemical analyzer (CHI, USA) in a three-electrode cell system. The
prepared capacitive electrode was used as working electrode, a about 0.80 cm graphite rod in diameter of
1.40 cm were counter and reference electrodes, and the length of reference graphite electrode was
selected according to the equal electric quantity of charge and discharge curve. The electrochemical
impedance spectroscopy (EIS) was measured at the open-circuit potential over the frequency range of
0.02 to $10^5$ Hz with an a.c. amplitude of 5 mV. All electrochemical measurements were carried out at
room temperature.

3. Results And Discussion

3.1 Specific surface area of Mn$_{0.68}$Bi$_{0.32}$OCl

As shown in Fig. 1, the isotherm of Mn$_{0.68}$Bi$_{0.32}$OCl was classified as type IV with an H3 hysteresis loop.
The specific surface area (SSA) was calculated using the Brunauer Emmett-Teller (BET) method. The
pore-size distributions (PSDs) of Mn$_{0.68}$Bi$_{0.32}$OCl were also computed by the Barrett Joyner Halenda
(BJH) plots. The peak in PSDs (shown in Fig. 1 inset) was centered at 37.3 and 242.2 nm. The SSA
values of Mn$_{0.68}$Bi$_{0.32}$OCl calculated by BET method were 14.6 m$^2$ g$^{-1}$, which was more than 13.28 m$^2$
g$^{-1}$ of Mn-doped BiOCl [22], and conducive to the penetration of electrolyte into the surface of
electroactive substances.

3.2 Micrographs and EDS spectrum of Mn$_{0.68}$Bi$_{0.32}$OCl

The SEM micrographs and the EDS spectrum of Mn$_{0.68}$Bi$_{0.32}$OCl are shown in Fig. 2 and Fig. 3,
respectively. The SEM micrographs in Fig. 2 showed that the smooth surface, neat edges and layered
particles could be ascribed to the Bi (Mn) OCl mix-crystals. The EDS measurements in Fig. 3 further
demonstrated the chemical composition of products. The EDS spectra and element analysis revealed
that the rate of O : (Mn+ Bi) : Cl was 1.42 : 0.73 : 1, while the rate of Mn: Bi was 2.09 : 1. From Table
the measurement errors of O and Cl elements were more than those of Mn and Bi elements, suggesting that
the chemical formula of the so-prepared product was Mn$_{0.68}$Bi$_{0.32}$OCl.

The reaction mechanism should be as follows: The Bi$^{3+}$ in the dilute nitric acid was hydrolyzed to
BiONO$_3$, then which was substituted by the Mn$^{2+}$ and Cl$^-$. The sublimation temperatures of MnCl$_2$ and
BiCl$_3$ are 1190 and 430$^\circ$C, respectively [24], respectively, so the product contained less Bi. The reaction
equations were given as
$$\text{Bi(NO}_3\text{)}_3 + \text{H}_2\text{O} \rightarrow \text{BiONO}_3 \downarrow + 2\text{HNO}_3$$

$$\text{BiONO}_3 + \text{MnCl}_2 + \text{O}_2 \rightarrow \text{Mn}_{0.68}\text{Bi}_{0.32}\text{OCl} + \text{BiCl}_3 \uparrow + \text{NO}_3^-$$

$$4\text{HNO}_3 \rightarrow 4\text{NO}_2 \uparrow + 2\text{H}_2\text{O} + \text{O}_2 \uparrow$$

### Table 1
Elemental analysis of Mn$_{0.68}$Bi$_{0.32}$OCl

| Element | Weight% | Atom% | Intensity | Error % |
|---------|---------|-------|-----------|---------|
| O K     | 16.97   | 45.29 | 14.08     | 14.59   |
| BiM     | 36.67   | 7.49  | 79.66     | 3.69    |
| ClK     | 26.20   | 31.56 | 41.75     | 11.46   |
| MnK     | 20.16   | 15.67 | 34.52     | 4.23    |

### 3.3 XRD pattern of Mn$_{0.68}$Bi$_{0.32}$OCl

The XRD pattern of Mn$_{0.68}$Bi$_{0.32}$OCl is shown in Fig. 4. The diffraction peaks of Mn$_{0.68}$Bi$_{0.32}$OCl matched well with the structure of BiOCl (PDF #85-0861), the peaks of $12.14^\circ$, $24.28^\circ$, $25.99^\circ$, $32.63^\circ$, $33.58^\circ$, $36.68^\circ$, $41.02^\circ$, $46.78^\circ$, $49.56^\circ$, $49.83^\circ$, $54.20^\circ$, $55.24^\circ$, $58.74^\circ$, $60.51^\circ$, $60.78^\circ$, $63.15^\circ$, $68.19^\circ$, $69.59^\circ$, $75.09^\circ$, $77.62^\circ$, $77.82^\circ$ could be assigned to the diffraction plane of (001), (002), (101), (110), (102), (003), (112), (200), (004), (113), (211), (10 4), (212), (114), (203), (005), (220), (221), (214), (310) and (302), respectively, the (001) peak of the Mn$_{0.68}$Bi$_{0.32}$OCl showed a clear shift of $\sim 0.112^\circ$ to the right compared to the BiOCl, and no other impurity peaks were found, demonstrating that the Bi$^{3+}$ ions of BiOCl were easy to be replaced by Mn$^{3+}$ ions to produce mix-crystal Mn$_{0.68}$Bi$_{0.32}$OCl.

### 3.4 TG and DSC of Mn$_{0.68}$Bi$_{0.32}$OCl

The TG and DSC curves of Mn$_{0.68}$Bi$_{0.32}$OCl are shown in Fig. 5. When the temperature raised to 700 °C, the mass of sample increased by about 1.8 %, which could be attributed to the oxidation of MnOCl on the surface of Mn$_{0.68}$Bi$_{0.32}$OCl to Mn$_2$O$_3$ due to the stable Mn$_2$O$_3$ [25, 26]; when the temperature raised from 832 °C the mass of 86.8 % was remained, and the mass in the temperature range of 832 to 898 °C slowly decreased to 86.0%, which could be attributed to the oxidative decomposition of Mn$_{0.68}$Bi$_{0.32}$OCl into Mn$_2$O$_3$, Bi$_2$O$_3$, and BiCl$_3$ [27] because BiOCl was steady below 600°C [13], while when the temperature reaches 1000 °C the mass of 79.3 % was remained, reveling that Mn$_{0.68}$Bi$_{0.32}$OCl was a higher safe material.

### 3.5. Electrochemical performance of Mn$_{0.68}$Bi$_{0.32}$OCl
The electrochemical performance of Mn$_{0.68}$Bi$_{0.32}$OCl was examined with cyclic voltammetry (CV) in Fig. 6 and galvanostatic charge/discharge in 1 M Na$_2$SO$_4$ aqueous solutions in a fixed voltage window of -0.50 ~ 0.50 V vs. graphite in Fig. 7. From Fig. 6(a) it could be seen that the oxidation current increased with the increase of voltage, and a oxidation peak was found at 2.556 V, which could be attributed to Mn$^{3+}$/Mn$^{4+}$, indicating that the change of electrode potential in the range of -0.5 ~ 0.5 V mainly came from the diffusion of ions, and the electrode of Mn$_{0.68}$Bi$_{0.32}$OCl with the richer O and Cl elements of higher electronegativity, had higher adsorption capacity for the Na$^+$ ions. Fig. 6(b) indicated that the electrodes in the range of -0.5 ~ 0.5 V had the characteristics of supercapacitor at low sweep speed.

The galvanostatic charge/discharge curves are shown in Fig. 7. The specific capacitance (SC) value was calculated by the formula as $SC = \frac{it}{m\Delta V}$. The SC value of Mn$_{0.68}$Bi$_{0.32}$OCl electrode (WE) at current density of 3, 5, 7, and 10 A g$^{-1}$ was 113.9, 63.0, 23.0 and 7.7 F g$^{-1}$, respectively, which was significantly affected by the current density. Compared with the bare electrode (BE) and the CBE (working electrode of absent Mn$_{0.68}$Bi$_{0.32}$OCl), an SC value of 5.8 F g$^{-1}$ for the BE and 9.0 F g$^{-1}$ for the CBE at the current density of 3 A g$^{-1}$ was less than that of the WE, revealing that the capacitance was mainly derived from the Mn$_{0.68}$Bi$_{0.32}$OCl.

The cycling stabilities of Mn$_{0.68}$Bi$_{0.32}$OCl were evaluated at a current density of 3 A g$^{-1}$ for 10000 cycles in Fig. 8. From Fig. 8 the SC values from cycle 1 to 12 at the current density of 3 A g$^{-1}$ increased due to the penetration of electrolyte into electrode active material, and the SC value of the first cycle was 124.2 F g$^{-1}$ at the current density of 3 A g$^{-1}$, which was close to the value of 113.9 F g$^{-1}$ obtained with CHI660e electrochemical analyzer. And from cycle 13 to 10000 the SC values at the current density of 3 A g$^{-1}$ were in the range of 211.8 ~ 224.0 F g$^{-1}$ with cycle efficiencies (n%) of 94.5 ~ 106.0 %, indicating that the electrode materials had higher stability.

The EIS analysis was also used to predict the behavior of the capacitive electrode. As shown in Fig. 9 the Faradic charge transfer resistances ($R_{ct}$) corresponding to the diameter of the semicircle in the plot, were closely related to the surface area and conductivity of the electrode, and the straight sloping line was associated with the ions diffusion. And the intercepts of BE, CBE, and WE in the Nyquist plots, which were related to the electrical resistance of the electrolyte ($R_{ele}$), were about 0.53, 0.79, 4.64 Ω with smaller $R_{ct}$, indicating that the electrolyte ions easily accessed the surface of active material, and the electroactive material on the electrode had stronger depolarization ability, which could promote the transfer of ions and electrons. Also, the slope of ions diffusion was steeper, demonstrating that Mn$_{0.68}$Bi$_{0.32}$OCl had faster diffusion of electrolyte ions.

4. Conclusions

In this paper the Mn$_{0.68}$Bi$_{0.32}$OCl were prepared using Bi (NO$_3$)$_3$ and MnCl$_2$ of molar ratio of 2:1 as precursors by hot solid-phase method. The Mn$_{0.68}$Bi$_{0.32}$OCl with simple preparation and low cost showed...
a distinctly improved electrochemical performance of supercapacitor because of the high SSA, good electrical conductivity and absorption and perfect stable crystallinity of Mn$_{0.68}$Bi$_{0.32}$OCl. The improved rate performance and good cycling stability make it as a promising electrode material for supercapacitor.

**Declarations**

**Acknowledgment**

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Figures
Figure 1

Nitrogen adsorption/desorption isotherms and BJH pore-size distribution of curves (inset) of Mn0.68Bi0.32OCl
Figure 2

SEM images of Mn_{0.68}Bi_{0.32}OCl.

Figure 3

EDS spectra of Mn_{0.68}Bi_{0.32}OCl
Figure 4

XRD pattern of Mn0.68Bi0.32OCl (a) and BiOCl (b)

Figure 5
Figure 6

Cvs of Mn0.68Bi0.32OCl at different voltage window (a) and scan rate of 5, 10, 30, and 50 mV s⁻¹ (b).
Figure 7

Galvanostatic charge/discharge curves of WE at the current density of $3 \times 10 A \, g^{-1}$ (a) and compared electrode at the current density of $1 A \, g^{-1}$ (b).
Figure 8

Plot of variation in the SC values of Mn0.68Bi0.32OCl at a current density of 3 A g⁻¹
EIS of Mn$_{0.68}$Bi$_{0.32}$OCl (a: BE; b: CBE; c: WE)