Contents lists available at ScienceDirect

### **Electrochemistry Communications**

journal homepage: www.elsevier.com/locate/elecom

#### Short communication

# Interconnected carbon-decorated Tio<sub>2</sub> nanocrystals with enhanced lithium storage performance

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#### ARTICLE INFO

Article history: Received 6 November 2013 Received in revised form 12 December 2013 Accepted 20 December 2013 Available online 29 December 2013

*Keywords:* TiO<sub>2</sub> Anode Lithium ion batteries

#### 1. Introduction

Enormous efforts have been recently made to develop high performance anode materials for next generation lithium ion batteries (LIBs) [1–3]. Nanostructured titania with various polymorphs and morphologies have been considered as promising candidates because of its fast lithium insertion/deinsertion kinetics, high operating voltage, reliable safety, non-toxicity and low cost [4–7]. However, some shortcomings such as crystal aggregation during cycling and limited electric conductivity have been observed in the pure nanostructured TiO<sub>2</sub> anodes [8,9]. In order to overcome these obstacles, carbon-coated titania nanostructures have been widely investigated and performance improvement as anodes in LIBs has been observed [8–10]. TiO<sub>2</sub> micro-/nano-structure was first prepared and subsequent composite formation between TiO<sub>2</sub> and carbon phase was then conducted. Similar methods such as mechanical ball milling with graphite and post-pyrolysis of carbon precursors have been employed. However, it is unavoidable that these two-step methods are complicated and timeconsuming [11-13]. Although one-step hydrothermal route was tried [25], the titania precursors were sensitive to moisture and reproducibility of preparation may be concerned. Hence, facile one-step method using stable precursors is highly required. Furthermore, performance of TiO<sub>2</sub> anode was reported to depend on the preparation temperature [6]. With low-temperature heat-treatment (<500 °C), distorted crystal lattice and deleterious surface functional groups can exist in TiO<sub>2</sub>. These problems result in highly irreversible Li<sup>+</sup> consumption during the first lithiation process and low initial coulomb efficiency (IE) is unavoidable [6]. For the purpose of IE improvement, calcination temperature should

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#### ABSTRACT

A novel direct preparation method was developed to prepare interconnected carbon-decorated  $TiO_2$  nanocrystals. The as-prepared composite exhibited a morphology of interconnected 10 nm  $TiO_2$  nanocrystals with uniform carbon coverage. As anode in lithium ion batteries, high reversible capacity of 184 mAh g<sup>-1</sup> with 87.4% initial coulomb efficiency, outstanding rate performance and stable cycle life were observed, which was attributed to ultrafine particle size, uniform carbon coating and high preparation temperature.

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be as high as possible, which usually brings about crystal aggregation and coarsening of  $TiO_2$  [6].

Herein, we propose a direct preparation of novel  $TiO_2-C$  nanocomposite starting from an environment-friendly and moistureinsensitive precursors, namely interconnected carbon-decorated titania nanocrystals (CDTN). In our novel preparation, highly crystalline  $TiO_2$  nanoparticles with uniform carbon coating survived even after high calcination temperature. Following advantageous features are expected in CDTN anode: i) ultrafine nano- $TiO_2$  crystals possess enhanced lithium uptake capability; ii) uniformly decorated carbon phase makes an improved electrical conductive network; iii) carbon phase avoids the direct contact between individual titania crystals, thus preventing crystal aggregation during battery cycling; iv) highly developed  $TiO_2$  crystal structure with complete removal of surface functional groups ensures an improved IE.

#### 2. Experimental

### 2.1. Preparation of the interconnected carbon-decorated titania nanocrystals (CDTN)

Self-made resol oligomers (Mw < 500) and titanium citrate complex solution were prepared according to reported method [10,15]. Typically, 4 g of 20 wt.% resol ethanol solution and 6 ml of 0.5 M titanium citrate complex ethanol solution was added dropwise into 8 g of ethanol under stirring. After stirring for 2 h, orange solution was poured into a big dish and kept in the ambient environment. After complete solvent evaporation, the dish was transferred into the oven at 100 °C for 24 h and a polymeric film was acquired. The film was carbonized at 750 °C for 2 h (the temperature ramping rate is 5 °C min<sup>-1</sup>). As soon as the temperature was cooled down to the room temperature, the carbonized





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sample was ground into fine powders and indexed as  $carbon-TiO_2$  composite (CT). Then, the CT sample was calcinated at 450 °C for 30 m in air (the temperature ramping rate is 6 °C min<sup>-1</sup>) to produce CDTN.

#### 2.2. Characterization and anode performance investigation

The X-ray diffraction (XRD) patterns were obtained with a Rigaku D/ Max-3C diffractometer equipped with a rotating anode and Cu  $K_{\alpha}$ radiation ( $\lambda = 0.15418$  nm). The external morphology of materials was examined using a scanning electron microscope (SEM, Philips XL30S FEG), whereas the pore images were obtained by a transmission electron microscope (TEM, TECNAI G2 T-20S). The anode performance was analyzed with a coin-type (CR2016) two-electrode cell with Li foil counter electrode (Cyprus Co.) and electrolyte of 1.0 M LiPF<sub>6</sub> in ethylene carbonate (EC)/dimethyl carbonate (DMC) (1:1 volume ratio) (Techno Semichem). The galvanostatic charge-discharge testing with a voltage range of 1.0–2.5 V vs. Li/Li<sup>+</sup> was conducted at the current rate from 0.2C to 10C (1C was in our experiments defined as 150 mA  $g^{-1}$ ) on a WBCS-3000 battery cycler (WonATech) at ambient temperature. The cycle performance for 100 cycles was recorded at 0.2C. Cyclic voltammetry was measured on IviumStat Electrochemical Interface (Ivium Technologies) with a voltage range of 1.0–2.5 V vs. Li/Li<sup>+</sup> at  $0.2 \text{ mV s}^{-1}$ .

#### 3. Results and discussion

Fig. 1 schematically shows formation process of the CDTN material. The resol oligomers and titanium citrate complex are initially dispersed in the ethanol and interacted each other. With process of ethanolevaporation, polymerization and carbonization, a carbon– $TiO_2$  nanocomposite (CT) with ~60 wt.% carbon is obtained. Afterward, CT is carefully calcinated under air, which produces interconnected carbondecorated TiO<sub>2</sub> nanocrystals (CDTN). The selection of precursors plays a key role for successful synthesis of CDTN. Titanium citrate complex, with a special cage-like titanium-centered complex molecular structure, strongly interacts and crosslinks with resols to form Ti-containing hybrid polymer [10,14–16], which probably retards organic–inorganic phase separation. Isolated Ti-containing nanophases can be homogeneously located within the resol polymeric matrix [14,15]. With calcination in nitrogen and air atmosphere sequentially, the Ti-containing nanophases are converted into ultrafine TiO<sub>2</sub> nanoparticles, while carbon derived from the resol polymer can coat on the TiO<sub>2</sub> surface. Because of strong confinement of these isolated Ti phases, fine nanostructure survives even after high-temperature heat-treatment.

In Fig. 2(a), TEM images of CT material show that high-contrast nanocrystals were uniformly dispersed in low-contrast carbon phase. In CDTN, morphology became mesoporous structure after carbon burning as shown in Fig. 2(b). From high resolution TEM images in Fig. 2(c) and (d), estimated TiO<sub>2</sub> crystal size was 5 and 10 nm for CT and CDTN, respectively, which coincided with average crystal size estimated by XRD patterns in Fig. 2(f) [17]. This indicated that fine titania nanocrystals formed in the initial step and severe crystal coarsening was effectively prohibited in CDTN. As shown in Fig. 2(f), main titania phase in both samples was anatase while rutile and brookite phase appeared as minor ones, which is associated with further heating process. In Fig. 2(e), SEM analysis shows external morphology of stacked TiO<sub>2</sub> nanoparticles in CDTN, implying that interstitial spaces between nanoparticles generated mesopores. Also, the inset EDS mapping demonstrated that



Fig. 1. Scheme of direct synthesis of the interconnected carbon-decorated titania nanocrystals (CDTN).

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