Active and Passive Elec. Comp., 1994, Vol. 16, pp. 113-117 Reprints available directly from the publisher Photocopying permitted by license only 1994 Gordon and Breach Science Publishers S.A. Printed in Malaysia

# MECHANISMS OF THE REVERSIBLE ELECTROCHEMICAL INSERTION OF LITHIUM OCCURRING WITH NCIMs (NANO-CRYSTALLITE-INSERTION-MATERIALS)

### S.D. HAN, N. TREUIL, G. CAMPET,\* J. PORTIER, C. DELMAS

Laboratoire de Chimie du Solide du CNRS, 351 cours de la Libération-33405 Talence--France J.C. LASSEGUES

Laboratoire de Spectroscopie Moléculaire et Cristalline du CNRS, 351 cours de la Libération-33405 Talence-France

A. PIERRE

Department of Mining-Metallurgical and Petroleum Engineering, University of Alberta-Edmonton-Alberta, T6G-2G6-Canada

(Received September 7, 1993; in final form November 8, 1993)

A new family of insertion-compound electrodes, so called NCIMs (Nano-Crystallite-Insertion-Materials), has been proposed: the major requirement is that the electrode materials have to be polycrystalline with a crystallite and particle size as small as possible (the accepted definition being that many crystallites make a particle). Indeed, by minimizing the size of the crystallites, the formation of defects bonds is favored, particularly at the crystallite surface, acting as reversible (de)grafting sites of  $Li<sup>+</sup>$ . Also, the cation-anion bonding is weakened not only in the grain boundary region but also within the crystallite close to its surface: then the electrochemical insertion of  $Li<sup>+</sup>$  takes place through easy bonding rearrangements.

### I. INTRODUCTION

In the last 20 years, much attention has been focused on  $A_xMO_z$ —type intercalation compounds  $(A = Li, Na$  and  $M = Co, Ni, Mn...$ , which are used as positive electrodes in reversible alkali electrochemical cells (see for example refs. 1). However, a very long-term cyclability (i.e., over  $10<sup>3</sup>$  cycles) might be hardly achievable, particularly for corresponding electrodes having a large grain size, probably because the  $Li<sup>+</sup>$  (de)intercalation process slightly perturbs the host lattice.

Some of us have patented, a few years ago, a new strategy and related experiments that have enabled us to put forward a rather new family of insertioncompound electrodes able to sustain long-term  $Li<sup>+</sup>$  electrochemical cyclability<sup>2</sup>. The major requirement is that the electrode materials are polycrystalline with a crystallite and particule sizes as small as possible. Therefore, we later called the polycrystalline electrode materials NCIM<sub>s</sub> (for nano-crystallite-insertion material $)^{3,4}$ .

Author for correspondence.

#### 114 G. CAMPET et al.

Some nanocrystanite insertion materials (TVCTMS)			
Sample	Class	Average grain size $(A)$	Insertion rate x (measured in $LiClO4$ (p.c.), $1.5V \le V(Li) \le 3.5V$
Li <sub>x</sub> SrTiO <sub>3</sub>		80	$0 \leq x \leq 0.3$
Li.CrO <sub>2</sub>		30	$0 \leq x \leq 1$
$LixMn2O3$		50	$0 \leq x \leq 2$
$LixFe2O3$	D	150	$0 \le x \le 0.5$
Li, NiO,		60	$1 \leq x \leq 2$
Li <sub>x</sub> CuO <sub>y</sub>		50	$1 \leq x \leq 2$
$Li$ , $WO$ ,		40	$0 \leq x \leq 2$

TABLE Some nanocrystallite insertion materials (NCIMs)

Table I gives important examples related to mixed-valency metal oxides<sup>2,5</sup>. For clarity, the examples listed in Table <sup>I</sup> have been divided into two classes, <sup>I</sup> and D, according to whether the resistivity tends to increase (class I) or to decrease (class D) upon the electrochemical  $Li<sup>+</sup>$  insertion process.

Rather similar considerations were reported by Barloux et al. and concern the spinel LiMn<sub>2</sub>O<sub>4</sub><sup>6</sup>. Also apparently related to that, Kumagai et al.<sup>7</sup> have reported that the positive electrode MnO<sub>2</sub>.  $vV_2O_5$  was formed by incorporation of  $V_2O_5$  into  $MnO<sub>2</sub>$  matrices and the crystallinity of the oxide decreased with increase in  $V<sub>2</sub>O<sub>5</sub>$ content incorporated. They have shown that the amount of  $Li<sup>+</sup>$  ions that can be reversibly electrochemically (de)insered increased with increasing y value, i.e., with decrease in the crystallinity; it reached about  $1 Li<sup>+</sup>$  per mole of the oxide with y  $= 0.6<sup>7</sup>$ .

In this paper, the framework of the model accounting for the reversible electrochemical  $Li^+$  insertion occurring in the NCIM<sub>s</sub> is presented. We also show, for the first time, that the model accounts for the evolution of the open circuit voltage of the electrodes, vs the fraction,  $x$ , of the alkali.

## II. MECHANISMS OF THE REVERSIBLE ELECTROCHEMICAL INSERTION OF LITHIUM OCCURING WITH NCIMs

First of all, by minimizing the size of the crystallites we tend to:

- (i) favor the formation of defect bonds, particularly at the crystallite surface (of its vicinity), such as anion adjacent to cation vacancies: *these defects act* as reversible (de)grafting sites for  $Li^{+}$ .
- (ii) weaken the cation-anion bonding not only on the grain boundary region but also within the grain close to its surface: then the electrochemical insertion of  $Li^+$  occurs through easy bonding rearrangements<sup>3</sup>.

That is depicted below for  $SrTiO<sub>3</sub>-NCIM$ , taken as a non-limiting but illustrative example. First of all, Fig. 1 illustrates the electron conduction via  $[Ti:3d]_{sub}$  or  $[Ti:3d]_{bw}$  energy states.

 $\bullet$  [Ti:3d]<sub>sub</sub> represents deep subband-gap energy states arising from cation defects adjacent to an anion vacancy. They are lowered below the  $\pi^*$  conduction



FIGURE 1 Simplified band energy scheme of  $SrTiO<sub>3</sub>-NCIM$ .

band of  $Ti^{4+}$ :  $3d^{0}(t_{2g})$  parentage. Conversely, anion defects adjacent to cation vacancies occur. They introduce acceptor states  $[O:2p]_{sub}$  arising from the  $O^{2-}$ :  $2p^6$  valence band. According to the model, the latter defects act as reversible (de)grafting sites for  $Li^+$ , (see (i)).

 $\bullet$  the [Ti: 3d]<sub>bw</sub> and [O: 2p]<sub>bw</sub> energy states originate from Ti-O bond weakening. This bond weakening induces  $Li<sup>+</sup>$  (de)insertion as mentioned above (see (ii)).

We will see, now, that the model accounts for the differences observed between the open-circuit voltage (OCV) vs x (the fraction of the alkali) curves related to polycrystalline electrodes having different sizes of crystallites. For sake of simplicity, such a behavior is illustrated only for two n-type electrodes  $Li<sub>x</sub>SnO<sub>2</sub>$  and  $Li<sub>x</sub>WO<sub>3</sub>$ (Fig. 2a, b).



FIGURE 2 Equilibrium OCVs vs x for some Li/LiCF<sub>3</sub>SO<sub>3</sub>/NCIMs (a: Li<sub>x</sub>SnO<sub>2</sub>; b: Li<sub>x</sub>WO<sub>3</sub>).

The concentration of the "sub" and b.w." states increases as the crystallite size is reduced. This obviously causes, only for the lower x values, a "pushing" of the Fermi-energy  $(E_F)$  and thereby of OCV towards "cathodic" values. Indeed, for the lower <sup>x</sup> values, the OCV are higher for the electrodes having the smallest crystallite size (Fig. 2a and 2b for  $x \le 0.15$ ).

For higher x values  $(x \gg 0.15)$  and when the inside-crystallite structure is well adapted for the reversible intercalation of lithium as it occurs for  $Li<sub>x</sub>WO<sub>3</sub>$ , an inversion of the OCV is observed (Fig. 2b): indeed for  $x \ge 0.15$  all the subband gap energy states  $[W^{6} : 5d^0]_{sub}$  and  $[\tilde{W}^{6} : 5d^0]_{low}$  (the "twin states" of  $[Ti^{4} : 3d^0]_{sub}$ and b.w. reported in Fig. 1) are filled with electrons. Therefore, the lithium intercalation within the nanocrystallites can now take place; it is accompanied with <sup>a</sup> "delocalization" of the injected electrons in the conduction band. On the other hand, it is well established that the band-energy width increases as the crystallite size decreases  $[8]$ ; therefore the  $WO_3$  electrodes having the smallest crystallite size have their conduction-band edge shifted towards "anodic" values: this causes a "decrease" of  $E_F$ , and thereby of OCV, towards "anodic" values (as it is illustrated on Fig. 2b for  $x \ge 0.15$ .

# III. THE NCIM HAVE BEEN INVESTIGATED IN DIFFERENT FORMS: EITHER POWDERED ELECTRODES (A) OR COMPOSITE ELECTRODES (B) OR THIN FILM ELECTRODES (c).

#### 1. Powdered Electrodes or Composite Electrodes

They can be efficiently used as positive electrodes for rechargeable lithium batteries of high energy density. Figure 3 shows the charge-discharge curves for powdered and composite electrodes.



FIGURE 3 Charge-discharge curves of: (a) Li/LiClO<sub>4</sub>-p.c./WO<sub>3</sub>-NCIM (50Å size), (b) Li/LiClO<sub>4</sub>p.c./"soft" composite electrode consisting of  $WO_3$ -NCIM (50Å size) dispersed in conductive polymer.



FIGURE 4 Transmission ( $\lambda = 550$  nm) vs time for repeated coloring and bleaching cycles with the NCIM (50 Å) counter electrodes  $Li_2xNiO_2$  (c). The Li<sup>+</sup> insertion rate, x, varies within 0 and 1. The coloring and bleaching potentials Vc and Vb are measured vs  $Li$  in  $LiClO<sub>4</sub> p.c.$  electrolyte.

### 2. Thin Film Electrodes

They can be efficiently used either as positive electrodes for thin-film rechargeable batteries of high energy density or as counter electrodes for electrochromic windows. Transmission vs time for repeated coloring and bleaching cycles is shown in Figure 4.

#### **REFERENCES**

- 1. J. Rouxel, in E Levy (ed.), Physics and Chemistry of Layered Materials, vol. VI, Reidel, Dordrecht, 1979.
	- K:. Mizushima, P.C. Jones, P.J. Wiseman and J.B. Goodenough, Solid State Ion., 3-4 (1981) 171.
	- C. Delmas, J. Braconnier, A. Maazaz and P. Hagenmuller, Rev. Chem. Miner., 19 (1982) 343. J. Molenda, Solid State Ion., 21 (1986) 263.
	- S. Miyazaki, S. Kikkawa and M. Koizumi, Synth. Met., 6 (1983) 211.
- 2. J.P. Couput, G. Campet, J.M. Chabagno, M. Bourrel, D. Muller, R. Garrié, C. Delmas, B. Morel, J. Portier and J. Salardenne, Int. Appl. Publ. under PCT. Int. Pat. Class GO2F 1701, FOI G9/ 00, C 23C 14/34, WO 91/01510, 1989.
- 3. N. Treuil, G. Campet, unpublished results: DEA report of N. Treuil. Bordeaux (1993).
- 4. G. Campet, S.D. Han, N. Treuil, MCR Shastry, J. Portier, C. Delmas, J.C. Lassègues, Mat. Sciences and Eng. B (submitted for publication).
- 5. B. Morel, Doctoral thesis, University of Bordeaux I (1991).
- 6. P. Barloux, J.M. Tarascon and EK. Shokoohi, J. Solid State Chem. 94 (1991) 185.
- 7. N. Kumagai, S. Tanifuji, T. Fujiwara and K. Tanno, Electrochim. Acta, 37(6) (1992) 1039.
- 8. P.E. Lippens and M. Lanoo, Physical Review B, 39 (1989) 15,



International Journal of Rotating

http://www.hindawi.com Volume 2014 Chemical Engineering International Journal of **Antennas and** 



http://www.hindawi.com Volume 2014

http://www.hindawi.com Volume 2014 Active and Passive Electronic Components

Shock and Vibration

http://www.hindawi.com Volume 2014

Acoustics and Vibration Advances in http://www.hindawi.com Volume 2014