Phosphorus-Derived Compounds for Li and Na Ion Batteries

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Abstract

In this review, progress in Phosphorus-based electrode materials for lithium, and sodium batteries, including Olivine, Tavorite, NASICON, Phosphite and phosphorene types, is briefly summarized. Such Phosphorus-based electrode materials will be attractive for next generation energy storage devices owing to its structural features which provide ultimate safety guarantee compared to other electrode materials.

Introduction

Rechargeable batteries have been paid remarkable attention and are regarded as the most promising energy storage system, especially for mobility (electric vehicle) and large-scale (grid level) energy storage applications, owning to the high energy density, power density and cycle life compared to other storage systems. Noted that during the last 10 years, Lithium and sodium ion battery research has been focused on which is made possible to deliver low cost and high energy density devices. Considerable research efforts have been made to investigate on high voltage, high-capacity and stable electrode materials for alkali metal ion batteries. Overall electrochemical performance of battery is truly depends on the characteristic features of electrode materials. Currently, Phosphorus-based compounds such as, metallo-phosphates, pyrophosphates, fluorophosphates, Oxy-fluorophosphates, mixed phosphates and phosphites have been considered as thermally and chemically stable materials for alkali metal ion batteries [1,2]. These compounds have an open phosphate structure, which provides an easy motion of alkali metal ions and the strong bonding of oxygen atoms with phosphorus helps to avoid oxidation with the electrolyte. Furthermore, elemental phosphorus (P) was found to be electrochemically active and used as an alkali metal ion battery anode. The three-electron-transfer reaction of phosphorus with alkali metal provides an extremely high theoretical capacity of 2595mAhg⁻¹, which is almost seven times higher than that of graphite. Therefore, these compounds have the ability to prevent an internal short circuit in the energy storage devices and high storage capacity; however, they have been suffering for low electronic & ionic conductivities and large volume expansion which are needed to be addressed for sustainable use of the materials.

Here we present a comprehensive mini review on current research progress that has been focused on the development of phosphates and phosphorus based materials for non-aqueous alkali metal ion battery electrodes. An overview of synthesis and electrochemical properties of polyanionic phosphates and phosphites versus Li & Na is provided.

Discussion

In 1997, Padhi et al. [3] demonstrate the electrochemical activity of LiFePO₄ olivine structure based on the two-phase reaction \( \text{LiFePO}_4 \rightarrow \text{FePO}_4 + \text{Li}^+ + e^- \) with a first-order phase transition and \( \text{Fe}^{3+}/\text{Fe}^{2+} \) redox potential of 3.4V [3]. All Olivine LiMPO₄ (M= Mn, Fe, Co and Ni) materials have low intrinsic electronic conductivity which are obstacles for the lithium ion motion in system. These drawbacks may be due to the (i) Li/M anti-site defects (ii) M ion on an Li site and a lithium vacancy and (iii) M⁺⁺ hole center (small polaron) and a lithium vacancy. Some of the defect mechanisms are unfavorable for lithium ion migration in
the crystal structure. In order to enhance the electronic conductivity and electrochemical properties of LiMPO$_4$ cathode materials, researchers have been playing around the microstructure of the electrode materials by carbon coating, doping with metal as well as the synthesis of nano-sized particles to obtain unfavorable defect free phase. The reported voltage ranges for LiFePO$_4$, LiMnPO$_4$, LiCoPO$_4$, and LiNiPO$_4$ are 3.4V, 4.1V, 4.8V and 5.1V vs Li/Li$^+$, respectively. Alkali metal oxy-phosphates AMOPO$_4$ (A= Li & Na; M= V & Ti) are considered as a potential candidate for high energy battery applications due to their high theoretical capacity. Among all Polyanionic phosphates, tavorite-type structures are capable of very high rates with one-dimensional lithium diffusion path. Another one, $\text{A}_{x}\text{MPO}_4\text{Y}$ Tavorite-type compounds (with $\text{A}=\text{Li}$, $\text{Na}$, $\text{Mg}$...; $\text{M}=\text{Ti}$, $\text{V}$, $\text{Mn}$, $\text{Fe}$; and $\text{Y}=\text{O}$, $\text{OH}$, $\text{F}$, and a mixture of them) are reported for the energy storage applications. Incorporated fluoride is replacing oxides on phosphorus site of metal phosphate frameworks, which changes both the structural and chemistry of materials. Fluoride rich framework have the ability to stabilize the structure and higher oxidations states on a transition metal due to the formation of bridge between $\text{F}^-$ and $\text{MO}$ in different transition metal centers and the formation of terminal fluoride sites on $[\text{MO}_n\text{F}_{6-n}]^{-}\text{m}^-$ and $[\text{PO}_4-p\text{F}_p]^{-}\text{q}^-$ polyhedral. These fluorophosphates are exhibit high voltage and excellent energy densities with very high cycling stability [4]. Furthermore, red and black phosphorous are the most used anode materials for both lithium and sodium ion batteries after silicon. The single-layered black phosphorus, named phosphorene, has been widely studied in battery application due to the graphite-like layered structure. In phosphorene, phosphorus atom is covalently bonded with three adjacent phosphorus atoms to form a puckered honeycomb structure [5]. As per the electrochemical activity, elemental phosphorus has enough small atomic weights and great Li-uptake ability (Li$_{3}$P, 2596 mAh g$^{-1}$) like silicon. The operating potential is much safer than graphite and delivers high capacity, which can be a suitable anode material for next generation alkali-metal-ion batteries.

**Summary and future outlook**

In summary, this short overview showed the amount of research efforts towards the development of phosphorus based active electrode materials for Li and Na ion batteries. Figure 1 & 2 Show the Average voltage and energy density (Wh kg$^{-1}$) versus gravimetric capacity (mAh g$^{-1}$) for phosphorus-based electrode materials for lithium and sodium ion batteries, respectively. The major drawback of phosphorus-based electrode materials is its poor electronic conductivity. To overcome these low intrinsic electronic conductivity and long diffusion path natures of phosphorus-based electrode materials, different strategies (like, carbon coating, composite with reduced graphene oxide, aliovalent transition metal doping and nano size particles) were adopted successfully. Altogether, it is clear that Phosphorus based materials will bring many technological breakthroughs in the future when they are employed in next generation alkali metal ion battery applications.

**Figure 1:** Phosphates for Lithium ion battery applications..
Figure 2: Phosphates for Sodium ion battery applications.

References