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Novel Nanostructured Metal Oxide Materials for Water Treatment and Advanced Rechargeable Batteries

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NOVEL NANOSTRUCTURED METAL OXIDE MATERIALS FOR WATER TREATMENT
AND ADVANCED RECHARGEABLE BATTERIES

by

Jingwan Huo

A Dissertation Submitted in
Partial Fulfillment of the
Requirements for the Degree of

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May 2018
ABSTRACT

NOVEL NANOSTRUCTURED METAL OXIDE MATERIALS FOR WATER TREATMENT AND ADVANCED RECHARGEABLE BATTERIES

by

Jingwan Huo

The University of Wisconsin-Milwaukee, 2018
Under the Supervision of Professor Chris Yuan and Professor Yin Wang

Clean water and energy shortage problems are pressing issues human facing today, as they not only cause many problems including public health and environmental deteriorations, but also threaten sustainable development of human society. It is urgent and critical to address and solve these problems.

Among various technologies, photocatalysis and adsorption applications in water and wastewater catch a lot of attention in recent years, for their low costs and for no requirement for chemical additives or thermal inputs. Photodegradation of contaminants by nanomaterials as catalysts is an important method in water and wastewater treatment. The adsorption is mainly through complexation between dissolved metals and the oxygen in metal oxides. Nano-adsorbents offer significant improvement with extremely high specific surface area and associated sorption sites, short intraparticle diffusion distance, tunable pore size and surface chemistry. Metal oxides such as iron oxide, titanium oxide, aluminum oxide and zirconium oxide are effective, low cost adsorbents for heavy metals and radionuclides.

Same as the situation of clean water supply, energy shortage is another urgent problem. As energy demand and consumption of conventional fossil fuel increase constantly, the resulting energy crisis and environmental problems caused by combustion of fossil fuel drive us to look for
more sustainable energy sources. Renewable energy resources appear to be an effective and promising solution. To overcome the variability for better renewable energy utilization, suitable energy storage devices are required.

Nanotechnology and nanomaterials are actively pursued to both improve existing technologies and develop new technology. TiO$_2$ nanomaterials have been intensively investigated in various applications, as photoelectrode, catalyst, sensor, energy storage. TiO$_2$ nanomaterials possess fascinating properties such as biological and chemical inertness, photostability, low cost, nontoxicity and superior oxidization ability. However, the wide electronic band gap and fast electron recombination rate limit the photocatalytic application and energy conversion efficiency of TiO$_2$ nanomaterials. Though TiO$_2$ can retain capacity at fast charge/discharge rates, and it is also electrochemical stable in common electrolytes and lack of harmful solid-electrolyte interfacial layer, the capacity of TiO$_2$ is relatively low. Zirconium oxide has valuable chemical and physical properties, including high melting point, mechanical and thermal resistance, low electrical conductivity, biocompatibility, chemical inertness (resistant to oxidant agent and acids/bases, nontoxic, and not dissolvable in water) is a widely used inorganic material. ZrO$_2$ is practical applied in fuel-cell technology, catalyst or catalyst support, oxygen sensor, thermal-barrier coatings and so on.

In this study, different modification strategies are carried out to improve the performance of TiO$_2$ and ZrO$_2$ in water treatment and energy storage applications. There are three objectives in this proposal. The first objective is to demonstrate high-efficiency photocatalysts using innovative hybrid nanostructures that consist of Pt nanoparticles and rGO co-modified three-dimensionally ordered microporous (3DOM) TiO$_2$. The excellent charge-separation property and high adsorption capacity of rGO increased the charge carrier lifetime and affinity to organic molecules. The
introduction of Pt nanoparticles increased spectral response to visible light through surface plasmon resonance and suppressed charge recombination. This study entails the synthesis and characterization of Pt/rGO-TiO₂ for application in methyl orange photodegradation. The second objective of this study is to demonstrate high-performance Lithium-ion battery electrode using hybrid nanomaterials consist of Fe₂O₃ nanospindles assembled on 3DOM TiO₂ with carbon coating. The carbon coated TiO₂@Fe₂O₃ material showed good electrochemical performance with exhibits a large reversible capacity about 570 mAh g⁻¹, which is about four times of the reversible capacity of 3DOM TiO₂. In addition to the high reversible capacity, the obtained material also exhibits good cycle performance and superior rate capacity. This rationally designed composite benefits from both good stability of TiO₂, high capacity of Fe₂O₃, and good electron conductivity of carbon. The third objective is to study zirconium modified clays as absorbents of phosphate from aqueous solution. Comparing three clays, zirconium modified MT (2:1), VT (2:1) and KT (1:1) exhibit different structure and surface properties, and thus performance differently toward phosphate adsorption. The adsorption kinetics data of phosphate on zirconium modified clays could be well described by the pseudo-second-order model, indicating that the adsorption was through chemisorption. The experimental equilibrium data of phosphate adsorption on modified clays were fitted better by Langmuir isotherm model than the Freundlich, implying monolayer adsorption. The effect of water chemistry (pH, co-existing anions, ionic strength, DOC) was also studied. These low-cost, abundant and effective are easily synthesized and have potential for practical wastewater treatment.
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<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
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<tbody>
<tr>
<td>3D</td>
<td>Three-dimensional</td>
</tr>
<tr>
<td>3DOM</td>
<td>Three-dimensionally order macroporous</td>
</tr>
<tr>
<td>ACAC</td>
<td>Acetylacetone</td>
</tr>
<tr>
<td>BE</td>
<td>Binding energy</td>
</tr>
<tr>
<td>CB</td>
<td>Conduction band</td>
</tr>
<tr>
<td>CCT</td>
<td>Colloidal crystals templating</td>
</tr>
<tr>
<td>CV</td>
<td>Cyclic voltammogram</td>
</tr>
<tr>
<td>CEC</td>
<td>Cations exchange capacity</td>
</tr>
<tr>
<td>DOC</td>
<td>Dissolved organic carbon</td>
</tr>
<tr>
<td>DRS</td>
<td>UV-vis diffused reflectance spectroscopy</td>
</tr>
<tr>
<td>EDCs</td>
<td>Endocrine disrupting compounds</td>
</tr>
<tr>
<td>$E_g$</td>
<td>Band gap energy</td>
</tr>
<tr>
<td>EIS</td>
<td>Electrochemical impedance spectroscopy</td>
</tr>
<tr>
<td>FCC</td>
<td>Face-centered-cubic</td>
</tr>
<tr>
<td>HA</td>
<td>Humic acid</td>
</tr>
<tr>
<td>HRTEM</td>
<td>High resolution transmission electron microscopy</td>
</tr>
<tr>
<td>ICP-OES</td>
<td>Inductively coupled plasma-optical emission spectroscopy</td>
</tr>
<tr>
<td>KT</td>
<td>Kaolinite</td>
</tr>
<tr>
<td>LIBs</td>
<td>Lithium-ion batteries</td>
</tr>
<tr>
<td>MMA</td>
<td>Methyl methacrylate</td>
</tr>
<tr>
<td>MO</td>
<td>Methyl orange</td>
</tr>
<tr>
<td>MT</td>
<td>Montmorillonite</td>
</tr>
<tr>
<td>NF</td>
<td>Nanofiltration</td>
</tr>
<tr>
<td>NOM</td>
<td>Natural organic matter</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Full Form</td>
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<tr>
<td>--------------</td>
<td>-----------</td>
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<tr>
<td>NPs</td>
<td>Nanoparticles</td>
</tr>
<tr>
<td>PMMA</td>
<td>Poly(methyl methacrylate)</td>
</tr>
<tr>
<td>PPCPs</td>
<td>Personal care products</td>
</tr>
<tr>
<td>PS</td>
<td>Polystyrene</td>
</tr>
<tr>
<td>rGO</td>
<td>Reduced graphene oxide</td>
</tr>
<tr>
<td>SEI</td>
<td>Solid-electrolyte interface</td>
</tr>
<tr>
<td>SEM</td>
<td>Scanning electron microscopy</td>
</tr>
<tr>
<td>SPR</td>
<td>Surface plasmon resonance</td>
</tr>
<tr>
<td>TEM</td>
<td>Transmission electron microscopy</td>
</tr>
<tr>
<td>UV</td>
<td>Ultraviolet</td>
</tr>
<tr>
<td>UV-vis</td>
<td>Ultraviolet-visible</td>
</tr>
<tr>
<td>VB</td>
<td>Valence band</td>
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<td>VT</td>
<td>Vermiculite</td>
</tr>
<tr>
<td>XPS</td>
<td>X-ray photoelectron spectroscopy</td>
</tr>
<tr>
<td>XRD</td>
<td>X-ray diffraction</td>
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</table>
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CHAPTER 1  Introduction and research objectives

1.1 Research background

Clean water supply is a major problem worldwide. As shown in Figure 1-1, the worldwide water demand is increasing from 2000 to 2050, and two-thirds of the world population will face water scarcity by 2025 [1, 2]. About half of the world’s population (2.6 billion) lacks access to basic sanitation, and 1.1 billion people are using water without improved sources [3]. The shortage of clean water causes many problems including public health and environmental deteriorations. Furthermore, with increasing population and industrialization practices, the rate of contamination of water supplies has been accelerated, water pollution becomes more complex and difficult to remove, which amplifies the public and environmental problems [4-6]. There is an urgent need to provide effective and affordable water treatment in developing countries.

Figure 1-1 (a) Global water demand in 2000 and 2050 [1] and (b) predicted water scarcity and stress in 2025 [2]

To address these issues, various water treatment technologies have been developed, including coagulation and flocculation, filtration, disinfection, lime and soda softening, ion exchange, evaporative processes, adsorption, membrane filtration, photodegradation, etc. [7, 8]. Regardless, numerous research has been conducted on the development of more effective and low-
cost methods and materials for water treatment. Nanotechnology is actively pursued to enhance the performance of existing treatment processes and to develop new processes. Nanotechnology-enabled water and wastewater treatment promises not only to overcome major challenges faced by existing treatment technologies, but also to provide new treatment capabilities that could allow economic utilization of unconventional water sources to expand the water supply [9]. Nanomaterial properties that are desirable for water and wastewater applications include high surface area for adsorption, high activity for photocatalysis, and other unique optical and electronic properties that find use in novel treatment processes and sensors for water quality monitoring.

As Professor Richard E. Smalley of Rice University gave in his presentation “Top Ten Problems of Humanity for Next 50 Year” on Energy & NanoTechnology Conference in 2003, energy and water are at the top of the list, followed by food, environment, poverty, etc. Professor Smalley also envisioned several problems are interlinked, such as the lack of people entering the fields of science and engineering, the need of alternative to fossil fuels and the need to address global warming. Same as the situation of clean water supply, energy supply shortage is another challenge that humans are facing today. According to International Energy Outlook 2016 by US Energy Information Administration (EIA), energy demand worldwide is projected to grow significantly over the 28-year period from 2012 to 2040. Total world consumption of marketed energy expands from 549 quadrillion British thermal units (Btu) in 2012 to 629 quadrillion Btu in 2020 and to 815 quadrillion Btu in 2040, a 48% increase from 2012 to 2040, as shown in Figure 1-2(a) [10]. Current primary energy consumption sources are still fossil fuels (oil, gas and coal), which is more than 85% of total energy consumption sources in 2015, according to World Energy Council, Resources 2016 Summary (Figure 1-2(b)). Such ever increasing energy demand could force the world to face energy crisis on non-renewable fossil fuels. What’s more, effluent gas
emissions from combustion of fossil fuels (CO, CO$_2$, SO$_2$, NO$_x$) could potentially damage world environmental healthy and global warming [11]. Renewable energy resources appear to be a promising alternative to fossil fuel, which is an effective and efficient solution to reduce greenhouse emission from fossil fuel consumption and address the global warming problem. As known, renewable energy resources are generated by natural resources, such as sunlight, wind, hydropower and geothermal energy, of which utilization is extensively limited by variable weather conditions. Thus, to overcome its variability and to use renewable energy, the availabilities of suitable energy storage devices, namely batteries, are required [12].

![Figure 1-2](image)

Figure 1-2 (a) World energy consumption, 1990-2040 [10], (b) comparative primary energy consumption over the past 15 years [13]

### 1.2 Water treatment through photocatalysis and adsorption

Removal of inorganic and organic substances in water to meet certain standard for reuse or drink is the goal for water treatment [14]. Typical inorganic solutes include heavy metals, oxyanions, radionuclides, fluoride, and cations that cause water hardness. Heavy metals such as Pb (II), Cd (II), Cr (VI), Cu (II), As (III) and so on, are serious environmental contaminants, due to their environmental persistence, high toxicity and tendency to accumulate in body tissues [15]. Most heavy metals are known to be carcinogenic agents, even a short-term exposure to heavy metal at trace level is a risk for human beings. For instance, the presence of lead in blood can damage various bodily systems, including the nervous and reproductive systems and kidney. Long
term exposure to arsenic via drinking water causes cancer of the skin, lungs, urinary bladder and the kidney [8]. Nitrogen and phosphorus are needed for aquatic organisms to grow, but the excessive presence of nitrogen and phosphorus in water bodies can cause eutrophication. As a result, heavy metals and phosphorus are targets to be removed from water, especially freshwater bodies [16].

Organic matters in water can be categorized as natural organic matter (NOM) and synthetic organic compounds. As an important component of organic matter, NOM is present in all sources of surface water and groundwater as a heterogeneous mixture of a variety of complex organic and slightly water-soluble materials, including humic substances (hydrophobic), hydrophilic acids, proteins, lipids, carboxylic acids, amino acids, and hydrocarbons [17]. In addition to NOM, many synthetic organic micro-pollutants, such as pesticides, endocrine disrupting compounds (EDCs), and pharmaceutical and personal care products (PPCPs) are also found in water sources [18-21]. Removing these types of synthetic organic matters from drinking water has gained increasing attention to reduce the potential risk to human and wildlife.

Among the various treatment techniques and processes for water and wastewater treatment, photodegradation and adsorption are conventional but efficient techniques for removing contaminants from water [8]. Photodegradation of contaminants by nanomaterials as catalysts is an important method in water and wastewater treatment. Utilizing sunlight and nanophotocatalysts to degrade organic contaminants is important in the area of water and wastewater treatment, especially for the wastewater containing small amounts of organic substance [22]. Photodegradation process possesses several advantages over competing processes, such as complete mineralization, no waste disposal problem, low cost and mild temperature and pressure conditions. The initial process for heterogeneous photocatalysis of organic and inorganic
compounds by semiconductors is the generation of electron-hole pairs in the semiconductor particles. Under light irradiation, the photocatalyst absorbs light with energy equal to or greater than the band gap of the semiconductor, an electron is exited from the valance band to the conduction band. The photoinduced electron transfer to adsorbed organic or inorganic compounds at the surface of semiconductor, and reduce dissolved oxygen in solution; while a hole can migrate to the surface, reacting with adsorbed water or surface hydroxyl groups (-OH) to produce hydroxyl radical (•OH) [22, 23].

Adsorption offers flexibility in design and operation, in many cases it will produce high-quality treated effluent. What’s more, most adsorption processes are reversible, the adsorbents can be regenerated for multiple use [24]. The adsorption is mainly through complexation between dissolved metals and the oxygen in metal oxides. The adsorption process can be divided into two steps: fast adsorption of metals ions on external surface of the adsorbent, and rate-limiting intraparticle diffusion along the micropore walls [9]. Nano-adsorbents offer significant improvement with extremely high specific surface area and associated sorption sites, short intraparticle diffusion distance, tunable pore size and surface chemistry. Metal oxides such as iron oxide, titanium oxide, aluminum oxide and zirconium oxide are effective, low cost adsorbents for heavy metals and radionuclides. Besides, clays, activated carbon, zeolites and other nano-structure materials are of great interest in application as support of nano-absorbents in water and wastewater treatment [25].

1.3 Lithium ion batteries

Lithium ion batteries (LIBs) have become the predominant power source for portable electronics for many years, with the advantages of high energy density, long lifespan and environmental benignity. There is desire for LIBs with high energy density and power density,
long cycle life, and high rate capability to meet the increasing demand [26, 27]. Rechargeable LIBs are considered as a reliable and promising source to store energy for hybrid electric vehicles (HEV) and plug-in hybrid electric vehicles (PHEV) [28]. Currently, graphite-based anodes are used in most commercially available rechargeable LIBs, which provides a theoretical capacity of about 372 mAh g\(^{-1}\) [28, 29]. Due to the low theoretical capacity, graphite electrode can only supply relatively low energy-storage ability and cannot meet the increasing power demand. The design and synthesis of new anode materials are necessary to reach higher performance. Transition metal oxides have shown high theoretical capacity (~500-1000 mAh g\(^{-1}\)), which is much higher than graphite material, making them promising anode materials for high performance LIBs. However, most of the transition metal oxides usually suffer from the problem of poor electronic conduction. Another challenge in using transition metal oxide is their poor cycling performance. Therefore, more efforts are needed to improve transition metal oxide anode materials.

1.4 Summary and literature review

1.4.1 Nanostructured TiO\(_2\) material

Titanium is the second most abundant transition metal in the Earth’s crust, and its oxide, TiO\(_2\), finds use in a wide array of applications, such as photoelectrode, catalyst, pigment, sensors, energy storage. Among various oxide semiconductor photocatalysts, TiO\(_2\) has been intensively investigated due to its fascinating properties such as biological and chemical inertness, photo-stability, low cost, nontoxicity and superior oxidization ability [30-32].

Bhatkhande et al. reviewed photocatalytic degradation for environmental applications. In the review they summarized that almost all types of organic and inorganic substances can be degraded using photocatalysis [22]. Introducing photo-catalytic properties into a membrane helps reduce organic fouling by promoting the degradation of organic matters on the membrane surface.
Nanomaterials (e.g., gold nanoparticles, TiO₂) have been widely applied to modify nanofiltration (NF) membranes and their photo-catalytic property have been investigated [33, 34]. Rajesh et al. impregnated poly(amide-imide) and TiO₂ nanoparticle into polysulfone NF membranes and reported a significant improvement in membrane performance including antifouling properties [35]. In a separate study, a TiO₂ nanowire membrane prepared through hydrothermal methods exhibited satisfied photocatalytic activity that nearly 100% humic acid removal was obtained by photocatalytic degradation and filtration [36]. Zhang et al. synthesized anti-fouling Fe₂O₃/TiO₂ nanowire membrane, the photocatalytic activity of hybrid membrane improved humic acid removal under solar irradiation (98% removal in 12 h), much higher than TiO₂ nanowire membrane (55% removal in 12 h) [37]. Modification of TiO₂ NF membrane through grafting methyl, phenyl or phosphonic acid groups onto the membrane surface has been shown to reduce membrane fouling by organic matters [38]. TiO₂ and silver nanoparticles have also been applied to modify polymeric membranes to reduce biofouling [39-42].

However, applications of TiO₂ as photocatalyst is limited by its fast electron recombination rate, which results in low photocatalytic reaction efficiency significantly. What’s more, bare TiO₂ has wide electronic band gap (3.2 eV), so it can only be excited by UV light which only accounts for 2-3% of the solar spectrum [32]. Numerous effects have been done to increase light response in visible light region and to reduce the recombination for enhancing light conversion efficiency. For example, Zhou et al. synthesized Ag₂O/TiO₂ nanobelts, which showed enhanced degradation of methyl orange under ultraviolet and visible light [43]. Ag/reduced graphene oxide (rGO) co-decorated TiO₂ nanotube arrays as a new photocatalyst, exhibited almost 100% photocatalytic removal of typical herbicide 2,4-dichlorphenoxyacetic acid under simulated solar light irradiation.
Under UV light irradiation, carcinogenic Cr(VI) in aqueous solution was photocatalytically reduced to less toxic form of chromium, Cr(III), over ZnO semiconductor catalyst [45].

A variety of TiO$_2$ nanomaterials, such as nanotubes, nanowires, nanoparticles, nanocrystal films, nanotube arrays, and more, have been reported. Three-dimensionally ordered macroporous (3DOM) materials are well-known for uniform pore size, high surface area and controllable, ordered porosity. 3DOM TiO$_2$ has attracted intensively attention in applications as catalyst or catalyst support, adsorptions, battery electrode materials and photonic materials. 3DOM structure has been demonstrated to enhance the light absorption and to improve light conversion efficiency, the periodic structure forbids the propagation of light with certain energies, increase the path length of light through slow-light effect and result in a stop-band reflection because of coherent Bragg diffraction [46-48]. Wu et al. studied optical and photocatalytic properties of 3DOM TiO$_2$ with different pore sizes, and they found when photonic bandgap edges match the electronic bandgap of TiO$_2$, slow photons effect at the photonic bandgap edges improved light absorption [49]. What’s more, when the photonic band gap is close to photocatalysis irradiation wavelength, light absorption by the material and photochemical process could be enhanced [50].

While application of TiO$_2$ in photocatalysis has been extensively studied for years, interests on TiO$_2$ as an anode material in LIBs developed several years ago, due to its electrochemical stability in common electrolytes and the lack of harmful solid-electrolyte interfacial (SEI) layers, resulting in better overcharge protection and safety [51]. Another key advantage of TiO$_2$ is that it can retain its capacity at fast charge/discharge rates [52]. 3DOM TiO$_2$ has been reported to possess excellent initial capacity of 248 mAh g$^{-1}$ at 0.2 C and 208 mAh g$^{-1}$ at 1C, and delivers a stable capacity (over 170 mAh g$^{-1}$ at a rate of C/2) over 100 cycles [51, 52].
Besides the photonic and electrochemical properties, hierarchical porosity of 3DOM systems are beneficial for heterogeneous catalytic applications, as they provide small pores for nanoparticle immobilization and the presence for larger pore networks reduces mass transport limitations [53]. In 3DOM structure, the macro-sized ball-shaped pores are closely packed with a high degree of order and interconnected to each other through small windows. The large pores in these materials have been found to improve the accessibility of reactants to the active sites of the material.

Based on literature review, as functional material or support, TiO$_2$ based nanomaterials have advantages as photocatalyst and anode material in LIBs. There are still some drawback limiting its application, such as large band gap, high recombination rate as photocatalyst, relatively low capacity compared to graphite-based anode materials.

1.4.2 Nanostructured ZrO$_2$ material

Zirconium oxide has valuable chemical and physical properties, including high melting point, mechanical and thermal resistance, low electrical conductivity, biocompatibility, chemical inertness (resistant to oxidant agent and acids/bases, non-toxic, and not dissolvable in water is a widely used inorganic material [54, 55]. ZrO$_2$ is practical applied in fuel-cell technology, catalyst or catalyst support, oxygen sensor, thermal-barrier coatings and so on [56].

Zirconium oxide exhibits specific binding affinity toward fluoride through metal-ligand interaction [57], strong affinity toward phosphate and arsenic through inner-complex [16, 54, 58], as a result, ZrO$_2$ is widely studied as adsorbents. Amorphous ZrO$_2$ demonstrates high adsorptive capacity due to the porous and highly hydrated structure, which allow ions to diffuse into the structure rather than restricted to just the external surface site. Shang’s group studied adsorption removal of arsenic and phosphate onto amorphous ZrO$_2$ NPs, which were synthesized by a simple
In their studies, the am-ZrO$_2$ NPs were found to have very strong adsorption on arsenic species and phosphate, through inner-complex mechanism. Mesoporous ZrO$_2$ was synthesized and showed maximum phosphorus adsorption capacity of 29.71 mg P/g [59]. Hydrous ZrO$_2$ NPs were reported strong arsenic adsorption for both As (III) and As (V) in both lab synthesized and natural water samples [58].

Usually presented as fine or ultrafine particles, zirconium oxide as absents are difficult to separate and recover from water by filtration or centrifugation. Also, when directly used in fixed-bed or any other flow-through systems, the pressure drop during operation baring technical bottleneck [57]. Therefore, supporting nanosized ZrO$_2$ onto substrate provides an attractive alternative to overcome the disadvantage of NPs. What’s more, dispersion of metal oxides over support with high surface area is an effective method to enhance the adsorption capacities of active species. Magnetic mesoporous SiO$_2$, SBA-15, ion exchanger, cotton fiber, graphite oxide, polymer resin have been studied as support of ZrO$_2$ nanomaterials, in environmental applications [57, 60-64]. Nano-hydrous zirconium oxide was encapsulated as the active species on an anion exchange resin, the supporting resin and hydrous zirconium oxide NPs captured phosphate through nonspecific electrostatic affinity and inner-sphere complexes, respectively [62]. The nanocomposite adsorbent exhibited highly preferable removal of phosphate from water, even other commonly occurring anions at greater levels presented in aqueous solution. Clays are known as low cost, abundance, high sorption properties and potential for ion-exchange, clays minerals are interesting materials for use as adsorbents, and have been studied as support for nanoparticles [65-69].
1.5 Objectives of study

The overall objective for the proposed study is to synthesize and modify nanostructured TiO\textsubscript{2} and ZrO\textsubscript{2}, in order to improve the performance in water treatment and energy storage applications. The first objective is to demonstrate novel TiO\textsubscript{2}-based photocatalysts with hybrid composites structures and enhanced activity for organic compounds degradation in water, and contribute more information about the reaction mechanism. The second objective is to incorporate TiO\textsubscript{2} and Fe\textsubscript{2}O\textsubscript{3} with novel structure to obtain a stable anode material for LIBs. The third objective is to prepare zirconium modified clays for phosphorus removal from water through adsorption, adsorption performance as kinetics and equilibrium isotherms was studied to understand the process. To achieve the research objectives, the work is divided into three major tasks, which are summarized below.

Task 1. Nanostructure Pt/rGO-TiO\textsubscript{2} photocatalyst for degradation of MO (Chapter 2)

The objective of this task is to synthesize Pt/rGO-TiO\textsubscript{2} nanocomposite, with Pt and rGO decoration on 3DOM TiO\textsubscript{2} simultaneously, which could help improve visible light absorbance and facilitate electron-hole separation to enhance photocatalytic activity. Three subtasks will be carried out:

- 3DOM TiO\textsubscript{2} will be synthesized by CCT and sol-gel method.
- Pt nanoparticles will be deposited on TiO\textsubscript{2} using impregnation method, and rGO will be thermo-reduced from GO, to obtain Pt/rGO-TiO\textsubscript{2} photocatalyst. And Pt-TiO\textsubscript{2} and rGO-TiO\textsubscript{2} will be synthesized as comparison.
- The resulting nanomaterials will be characterized using various techniques, such as scanning electron microscopy (SEM), transmission electron microscopy (TEM), high-resolution TEM (HRTEM), X-ray diffraction (XRD), X-ray photoelectron spectroscopy
(XPS), and UV-vis diffuse reflectance spectra. Finally, the nanomaterials will be applied to methyl orange (MO) photodegradation as model pollutant.

**Task 2. Carbon coated Fe$_2$O$_3$/TiO$_2$ anode material for LIBs (Chapter 3)**

The objectives of this task are to fabricate and characterize carbon coated Fe$_2$O$_3$/TiO$_2$ composite and apply the materials to LIBs anode. The electrochemical performance of Carbon coated Fe$_2$O$_3$/TiO$_2$ hybrid material will be evaluated. Three subtasks will be carried out:

- 3DOM TiO$_2$ will be synthesized by CCT and sol-gel method.
- Fe$_2$O$_3$ nanospindles will be assembled onto 3DOM TiO$_2$, and the resulted material will be carbon coated using glucose as carbon source.
- The obtained materials will be characterized using various techniques, such as scanning electron microscopy (SEM), transmission electron microscopy (TEM), high-resolution TEM (HRTEM), X-ray diffraction (XRD), X-ray photoelectron spectroscopy (XPS), Raman spectrum. And the nanomaterials will be tested as anode material for LIBs.
- Electrodes and half-coin cells will be prepared, and their performance will be investigated.

**Task 3. Zirconium modified clays for phosphorus removal from aqueous solution through adsorption (Chapter 4)**

The objective of this task is to fabricate and characterize zirconium modified clays, which could adsorb phosphate in water as low-cost adsorbents. To study phosphate removal performance of zirconium oxide on different supports, three clays with different structure are chosen to be studied. Three subtasks will be carried out:

- A series of zirconium modified clays with different Zr/clay ratios will be synthesized through simple reaction, using ZrOCl$_2$ as zirconium source.
• The resulting materials will be characterized using various techniques, such as scanning electron microscopy (SEM), X-ray diffraction (XRD), zeta-potential measurement.

• Phosphorous adsorption capacity of different zirconium modified clays will be evaluated by equilibrium isotherm and kinetic study at neutral pH. The equilibrium isotherms and kinetic results will be analyzed by appropriate models to understand the adsorption process.

• The impact of clay structure as support of zirconium oxide will be discussed.
CHAPTER 2  Pt/rGO-TiO$_2$ photocatalyst for degradation of MO

2.1 Introduction

Semiconductor-based heterogeneous photocatalysis has been increasingly applied to deal with environmental pollution problems [30]. Photocatalytic oxidation of organic pollutants in water is a promising technique for water and wastewater treatment [70]. Under light irradiation, the photocatalyst absorbs light with energy equal to or greater than the band gap of the semiconductor, an electron is exited from the valance band (VB) to the conduction band (CB), leaving a positively charged hole (h$^+$) at the band edge of the VB. The photoinduced electrons ($e^-$) transfer to adsorbed organic or inorganic compounds at the surface of semiconductor, and reduce dissolve oxygen in solution; while a hole can migrate to the surface, reacting with adsorbed water or surface hydroxyl groups (-OH) to produce hydroxyl radical (•OH) [22, 23]. Among various oxide semiconductor photocatalysts, titanium oxide (TiO$_2$) has been intensively investigated due to its fascinating properties such as biological and chemical inertness, photostability, low cost, nontoxicity and superior oxidization ability [30-32]. However, applications of TiO$_2$ as photocatalyst is limited by its fast electron recombination rate, which results in low photocatalytic reaction efficiency significantly. What’s more, bare TiO$_2$ has wide electronic band gap (3.2 eV), so it can only be excited by UV light which only accounts for 2-3% of the solar spectrum [32]. To achieve effective light conversion efficiency with TiO$_2$-based photocatalysts, more effects are still required to extend light response in visible light region and reduce the recombination.

Among a variety of TiO$_2$ nanomaterials, three-dimensionally ordered microporous (3DOM) structure is drawing increasing attention for its advantageous properties, including unique periodic structure, optical properties and favored mass transfer. 3DOM materials, also known as inverse opal, are commonly synthesized through colloidal crystals templating (CCT) method. The
colloidal crystals are self-assembled from monodisperse microspheres, such as poly(methyl methacrylate) (PMMA), polystyrene (PS), silica, to a face-centered close-packed or opal arrangement. A liquid precursor penetrates the opal and fills the voids of the templates. After solidification of the precursor, the colloidal templates are removed by calcination or extraction, yielding an inverse opal replica of the array of ordered spheres. Where the original microspheres made contact, the macropores are interconnected by windows.

3DOM structure has been demonstrated to enhance the light absorption and improve light conversion efficiency, the periodic structure forbids the propagation of light with certain energies, increase the path length of light through slow-light effect and result in a stop-band reflection because of coherent Bragg diffraction [46-48]. Wu et al. studied optical and photocatalytic properties of 3DOM TiO$_2$ with different pore sizes, and they found when photonic bandgap edges match the electronic bandgap of TiO$_2$, slow photons effect at the photonic bandgap edges improved light absorption [49]. What’s more, when the photonic band gap is close to photocatalysis irradiation wavelength, light absorption by the material and photochemical process could be enhanced [50]. Besides the photonic properties, hierarchical porosity of 3DOM systems are beneficial for heterogeneous catalytic applications, as they provide small pores for nanoparticle immobilization and the presence for larger pore networks reduces mass transport limitations [53].

In order to reduce the electron recombination and widen the light absorption spectrum of TiO$_2$, studies have been done about modifications on TiO$_2$. Doping with other elements or incorporating with electron-accepting materials, such as loading noble metals like Au or Pt to enhance the separation of the photogenerated electrons and holes, or carbon nanotubes or graphene to increase the visible-light adsorption [71, 72]. Owing to its excellent catalytic property, Pt is the most widely used active catalyst, and decoration of TiO$_2$ by Pt has been studied [73]. In aspect of
photocatalyst, Pt nanoparticles respond strongly to visible light due to their surface plasmon resonance (SPR), which refers to a collective oscillation of conduction electrons induced by the electric field of light [74-76]. By incorporating Pt nanoparticles on the surface of inverse TiO$_2$ opals, more light is absorbed and the lifetime of the light-excited electron and holes are extended. Liang et al studied synthesized 3DOM Pt/TiO$_2$ as catalyst for water-gas shift reaction, and they observed very good catalytic performance due to better mass transfer of 3DOM porous structure and high intrinsic activity of Pt/TiO$_2$ [77]. Synergy of slow photon effect and chemically amplified photochemistry in Pt loaded 3DOM TiO$_2$, as photocatalyst of methyl orange (MO) degradation. The improved photocatalytic activity is attributed to the existence of metallic Pt, which facilitates the scavenging of the excited electron and increases efficient electron-hole separation [71].

However, the catalytic activity of noble metal/TiO$_2$ materials is severely limited by the ultrafast relaxation of hot electrons (<160 fs) and rapid charge recombination. Coupling with graphene - an electron acceptor material with high electron mobility, has been reported to maximize the charge separation [73, 78]. Graphene is known with excellent conductivity, large specific surface area, and high hydrophobicity, not only can significantly suppresses recombination of photogenerated electron-hole pairs of TiO$_2$ and increasing charge transfer rate of electrons, but also can extend light absorption spectrum, and increase surface-adsorption capacity to organic molecules [32, 44, 79, 80]. Combining noble metal nanoparticles and graphene materials to improve TiO$_2$ catalytic activity has been studied, based on different TiO$_2$ structure, including nanoparticles, nanotubes and 3DOM TiO$_2$ [44, 78, 81, 82]. The ternary hybrid catalysts based on different structures of TiO$_2$ (3DOM, nanotube arrays and nanosheets) all showed significantly improved catalytic activity and stability, which provides some new hints for improving TiO$_2$ photocatalytic activity under visible light.
Recently Boppella et al. reported Au nanoparticles and reduced graphene oxide (rGO) incorporated TiO$_2$ inverse opal photoelectrodes, focusing on solar-to-energy application as photoelectrochemical (PEC) water splitting devices [78]. The literature reports provide a guide for rational design a novel and effective visible-light driven photocatalyst in water treatment. In the present work, a new Pt/rGO-TiO$_2$ photocatalyst was designed and prepared by Pt nanoparticles and rGO co-modification on 3DOM TiO$_2$. 3DOM TiO$_2$ was synthesized through colloidal crystals templating (CCT) method using poly(methyl methacrylate) (PMMA) as sacrificial template and sol-gel method using Ti(OBu)$_4$ as precursor. The obtained 3DOM TiO$_2$ was modified with Pt nanoparticles and reduced graphene oxide for improved visible-light photocatalytic ability, as illustrated by the photocatalytic degradation of methyl orange (MO) as a representative dye. Pt/rGO-TiO$_2$ showed higher light absorption and photocatalytic degradation efficiency toward MO under visible light, compared to 3DOM TiO$_2$, Pt-TiO$_2$ and rGO-TiO$_2$. In the stability test, the prepared photocatalyst maintained high photocatalytic activity after 3 cycles. Thus, this new ternary hybrid photocatalyst has potential applications in environmental purification.

2.2 Experimental

2.2.1 Chemicals and materials

All chemicals were reagent grade and used without further purification. Methyl methacrylate (MMA), titanium butoxide (Ti(OBu)$_4$), acetylacetone (ACAC), ethanol, chloroplatinic acid hexahydrate (H$_2$PtCl$_6$·2H$_2$O), methyl orange (MO) were purchased from Sigma-Aldrich. Single layer graphene oxide was purchased from ACS Material. Potassium persulfate (K$_2$S$_2$O$_8$) was purchased from Alfa Aesar. Ultrapure water (resistivity > 18.0 MΩ) was used for all experiments.
2.2.2 Preparation of PMMA template

Monodisperse PMMA nanospheres were synthesized using surfactant-free emulsion polymerization according to literature techniques [83]. In a typical synthesis, a mixture of ultrapure water (165 mL) and methyl methacrylate (MMA, 18.6 mL) was prepared in a three-neck round-bottom flask, with a water-cooled condenser. The mixture was stirred at 350 rpm, while being heated to 70 °C and purged with nitrogen gas to deactivate the inhibitors. After stabilization of the temperature, K₂S₂O₈ (0.5 x 10⁻³ mol/L, 0.0236g) was added into mixture as initiator dissolved in 10 mL water. The reaction was sustained for 2 hours, producing colloidal PMMA spheres. PMMA colloidal crystals were formed by centrifuging the colloid at 1500 rpm for 24 h, decanting the water and allowing the solid dry for 3 days.

2.2.3 Preparation of 3DOM TiO₂

TiO₂ precursor was prepared by mixing Ti(OBu)₄, ACAC, H₂O and ethanol in molar ratio: 1:1:3:20 in open air at room temperature for 2 hours. The obtained solution was stored in brown glass bottle for future use. Dried PMMA colloidal crystals were crushed to a powder and deposited in millimeter-thick layers on filter paper in a Böchner funnel. With suction applied to the Böchner
funnel, the precursor solution was applied dropwise to completely wet the PMMA powder. The infiltration was repeated by 5 times. After dried at room temperature for 2 hours, the samples were calcined at 300 °C at a step of 2 °C per min for 3 hours to remove PMMA template and then at 450 °C at a step of 2 °C per min for 4 hours to calcine TiO₂.

2.2.4 Preparation of Pt-TiO₂

Pt was loaded on 3DOM TiO₂ with impregnation method using H₂PtCl₆ solution as precursor [77, 84, 85]. 1.5 mL of 7mM H₂PtCl₆ aqueous solution was dropped onto 0.1 g 3DOM TiO₂ sample and dried under air at 70 °C for 12 h and subsequently calcined at 500 °C for 3 h.

2.2.5 Pt/rGO-TiO₂ and rGO-TiO₂ composite catalysts

In a typical synthesis, desired amount of graphene oxide (0.127 mL, 10 mg/L) solution were made into 20 mL aqueous solution, and the mixture was ultrasonicated for 1 hour (Branson, 2510). Then certain amount of Pt-TiO₂ (0.0636g) were added in to GO solution and ultrasonicate for 1 hour, the ratio of GO/Pt-TiO₂ was 2 wt%. Pt/GO-TiO₂ was filtered with FP-200 membrane (Paul Life Sciences, PVDF) and dried in air. The obtained material was calcined at 500 °C for 2 h to reduce GO to graphene. rGO-TiO₂ was obtained by the same method by adding TiO₂ instead of Pt-TiO₂.

2.2.6 Material characterization

The morphology and of sample were imaged with scanning electron microscopy (SEM) using a Hitachi S-4800 SEM microscope and Hitachi H9000NAR transmission electron microscopy (TEM). The composition of the obtained material was examined by Energy-dispersive X-ray spectroscopy (EDS) integrated with SEM. The content of Pt was analyzed by dissolving Pt nanoparticles in HCl and HNO₃ completely and measure the concentration of Pt in solution using inductively coupled plasma-optical emission spectroscopy (ICP-OES) (Perking Elmer Optima
The crystal structure and phase state of Pt-TiO$_2$ were determined by X-ray diffractometry (XRD) using Bruker D8 Discover A25 diffractometer with copper K$_\alpha$ radiation. Raman spectra were taken on a Reinshaw 1000B Raman spectrometer with a 632.8 nm HeNe laser source. X-ray photoelectron spectroscopy (XPS) was employed to characterize the chemical states of prepared photocatalyst using Perkin Elemer PHI 5440 ESCA system with an Al K$_\alpha$ X-ray source. The UV-Vis diffuse reflectance spectra were tested using a Shimadzu UV-2600 UV-Vis spectrophotometer, with BaSO$_4$ as reference.

### 2.2.7 Photocatalytic activity test

The photocatalytic activity of 3DOM TiO$_2$, Pt-TiO$_2$, rGO-TiO$_2$ and Pt/rGO-TiO$_2$ was quantified by monitoring the degradation of MO under visible light irradiation. A 300 W Xenon lamp was used as the light source (CEL-HXF300, Beijing China Education Au-light Co., Ltd). The photocatalytic reaction was performed in a reactor, which was surrounded by water-cooled jacket. A cut-off filter (>420 nm) was mounted before the output of light to removes UV light and admit only visible light to enter the reactor. For these MO degradation tests, 20 mL aqueous suspensions of MO (10 mg/L) and 20 mg photocatalyst powders were placed in reactor. The dispersions were kept in the dark for 30 minutes with magnetic stir to achieve adsorption-desorption equilibrium. Aliquots (1 mL) were collected and centrifuged to remove catalyst particles every 15 min during visible-light illumination. The concentration of residual MO was determined by monitoring its UV-visible absorption peak at absorption at 464 nm [86]. The concentration of MO is proportional to its absorbance. MO degradation was expressed as $C/C_0$ vs irradiation time, where $C_0$ is the concentration of the original MO solution after 30 min adsorption, and $C$ is the concentration of MO solution after every 15 min reaction.
To test the reusability of the Pt/rGO-TiO$_2$ sample, the photocatalytic test was repeated three times. The experimental conditions were the same as previously described, except the reaction time was limited to 90 min. After each round of the reaction, the mixture was recovered by centrifuged and rinsed by ultrapure water.

2.3 Results and discussion

2.3.1 Synthesis and characterization of materials

Pt/rGO-TiO$_2$ composite was synthesized in three steps (Figure 2-1): (1) synthesis of 3DOM TiO$_2$ through CCT and sol-gel method, (2) Pt nanoparticles were loaded onto 3DOM TiO$_2$ through impregnation method using H$_2$PtCl$_6$ as precursor and (3) modification and reduction of GO in Pt/rGO-TiO$_2$ composite. A combination of techniques, described individually in detail in the Experimental Section, were used in the synthesis process, including CCT, sol-gel processing, impregnation method.

As shown in Figure 2-2(a), 3DOM TiO$_2$ exhibits a highly ordered microporous structure, which replicate the three-dimensional closely-packed PMMA opals in Figure S2-1 (Appendix), with pore diameter about 210 nm and 25 nm wall thickness. The pore size of 3DOM TiO$_2$ is smaller than PMMA spheres (~330 nm) due to a shrinkage during the calcination. The small windows between the opal pores can be clearly observed in Figure 2-2(b), which replicates from the contact bridges between the neighboring PMMA spheres. The underlying layer of pores and porous walls can be clearly observed, indicating that the sample possesses a three-dimensionally well-open, ordered and interconnected microporous network. Figure 2-2(b) shows that Pt nanoparticles have been successfully deposited on the surface of 3DOM TiO$_2$ by the impregnation method. Pt nanoparticles are well dispersed on TiO$_2$ surface and uniform in size. Pt content of 1.7 wt% was confirmed by ICP-OES, which is close to the experimental design (2 wt%). After deposition and
reduction of GO on TiO$_2$, a continuous and transparent rGO film is formed on the surface of 3DOM TiO$_2$ (Figure 2-2(c)). When using Pt-TiO$_2$ instead of bare 3DOM TiO$_2$, the ternary Pt/rGO-TiO$_2$ composite was obtained (Figure 2-4(d)).

The Pt-TiO$_2$ sample was further characterized using TEM to confirm the phase of TiO$_2$ and the existence of Pt nanoparticles. From Figure 2-3(a) the well-defined 3DOM structure with overlapped pores can be easily observed, and the porous skeleton of 3DOM TiO$_2$ are composed of many nanoparticles. At higher magnification (Figure 2-3(b)), it is clear to see Pt nanoparticles are highly dispersed on 3DOM TiO$_2$ wall and sizing range from 3-6 nm. HRTEM image (Figure 2-3(c)) shows the lattice fringes of both TiO$_2$ and Pt, which indicates the highly crystalline nature of TiO$_2$ and Pt nanoparticles. The lattice fringe with a d-spacing of 0.352 nm can be assigned to the
(101) lattice plane of anatase TiO$_2$, and the fringe with d-spacing of 0.224 nm belongs to the (111) lattice plane of Pt nanoparticles.

Figure 2-3 (a)-(b) TEM images of Pt-TiO$_2$ with different magnifications and (c) HRTEM image of Pt-TiO$_2$
Figure 2-4 XRD patterns of as-prepared catalysts with 2 theta between (A) 10° and 70° and (B) 37° to 50°; Raman spectra of as-prepared catalysts with wavelength (C) 250-1800 cm⁻¹ and (D) 1100-1800 cm⁻¹. a) TiO₂, b) Pt-TiO₂, c) rGO-TiO₂, d) Pt/rGO-TiO₂.

The structure of catalysts was characterized by XRD, as shown in Figure 2-4(A). XRD patterns show the main phase of TiO₂ in all four samples are anatase, characteristic peaks at 25.3°, 36.9°, 37.8°, 48.0°, 53.1°, 55.1°, 62.7°, 68.8° can be assigned to the plane (101), (004), (200), (105), (211), (204), (116) of anatase (JCPDS 21-1272), respectively. The peaks at 27.5°, 36.1°, 41.2°, 44.1°, 56.6° represent planes (110), (101), (111), (210), and (220) of rutile (JCPDS 21-1276). With both Pt-TiO₂ and Pt/rGO-TiO₂ samples, the weak peak at 39.7° and 46.2° could be indexed to the (111) and (200) planes of the Pt nanoparticles (JCPDS 65-2868), respectively.
The weak peak at 22.8° in rGO-TiO$_2$ and Pt/rGO-TiO$_2$ deviates from the characteristic (002) plane reflections of graphitic carbon at 26.3° [87]. The broad diffraction peak of rGO indicates poor ordering of the sheets in the stack direction, which may due to mainly single or only a few layers of rGO in composites [88]. The main characteristic peak of rGO (ca. 25) has low intensity and overlap with the peak of anatase TiO$_2$ at 25.3° [89].

The crystalline structure of TiO$_2$ and rGO were further investigated by Raman spectroscopy, as shown in Figure 2-4(C)-(D). For all samples, the characteristic peaks at 146 (E$_g$), 396 (B$_{1g}$), 515 (A$_{1g}$) and 637 cm$^{-1}$ (E$_g$) are attributed to anatase TiO$_2$, which corroborate XRD results [30]. The peaks at 146 cm$^{-1}$ is derived from the bending vibration of O-Ti-O bond, the other three peaks are related to Ti-O-Ti bending types [48]. The weak peaks at 197 cm$^{-1}$ are assigned as rutile TiO$_2$. Characteristic bands at 1340 and 1590 cm$^{-1}$ were assigned to D and G peaks for rGO. D band represents the dispersive and defect-induced vibrations and G peak is related to the in-plane vibration of sp$^2$ bonded carbon atoms. The similarity of D and G peaks in intensity indicates the rGO contains a large amount of disordered sp$^2$ carbons, which are related to the defects that may be generated during the reduction process [90, 91].

XPS was employed to study the chemical states of elements and the interaction between TiO$_2$ and rGO in Pt/rGO-TiO$_2$ composite. The curve fitting of O 1s, C 1s, Ti 2p and Pt 4f peaks of XPS spectra for the composite are shown in Figure 2-5(a)-(d). The O 1s spectrum was fitted with two peaks at 530.1 eV and 531.9 eV. The former is the characteristic of the lattice oxygen of TiO$_2$, and the other one with 1.8 eV higher binding energy can be assigned to the surface-adsorbed component of the hydroxyl group (Ti-OH) [92]. Two types of carbon with different binding energies at 285.2 eV and 287.4 eV were observed, which were assigned to the sp2-hybridized C, the carbonyl C (C=O), respectively. The absence of C in C-O bonds and carboxylate C (O=C-O)
and low intensity of carbonyl C (C=O) indicate that GO was mainly reduced by removing C-O and C=O bonds and a high crystalline degree of graphene [78, 81, 90, 93]. There are two characteristic peaks at 459.3 eV (Ti 2p$_{3/2}$) and 465.0 eV (Ti 2p$_{1/2}$) on Ti 2p XPS spectrum, which correspond to spin-orbital splitting photoelectrons in anatase TiO$_2$ structures. The peaks shifted toward higher binding energy end comparing to pure TiO$_2$ (458.5 eV and 464.2 eV), the red-shift suggested the formation of strong electronic interactions between TiO$_2$ and rGO. The electronic interactions caused change in chemical state and/or coordination environments of Ti$^{4+}$, suggesting a charge transfer from TiO$_2$ to rGO [78, 94, 95]. The Pt 4f spectrum was fitted into three sets of peaks. The strongest set at 70.9 eV and 74.3 eV are assigned to metallic Pt. The second set of peaks at 72.7 eV and 75.9 eV is due to Pt (II), and the last set with lowest intensity at 75.2 eV and 77.6 eV could be from a small amount of Pt (IV) [71, 84].

The optical property of 3DOM catalysts, UV-vis diffuse reflectance spectroscopy (DRS) studies were performed on all samples, the results are shown in Figure 2-6(A). 3DOM-TiO$_2$ exhibits a wider light absorption region comparing to the commercial P25, owing to the slow photon effect and multiple scattering of the 3DOM structure. As shown in reflectance spectrum (Figure 2-6(A)-a), 3DOM TiO$_2$ has a strong stop band maximum at 400 nm which is near the electronic absorption edge. The strong reflection peak indicates a high-quality inverse opal structure in wide range [96]. According to modified Bragg equation, the photonic stop band of 3DOM TiO$_2$ was estimated at about 527 nm (Appendix) [97, 98]. When stop-band is in visible range, the visible light absorption by the material and photochemical process would be enhanced [50].
The modification of Pt obviously increased light adsorption in the visible light range owing to the surface plasmon resonance (SPR) effect and photosensitizing effect of Pt. Unlike other plasmon metal, no Pt SPR peak observed, because small Pt nanoparticles always exhibit broad extinction below 450 nm, without observable peaks [74-76]. It can be observed a red-shift in absorption edge and enhanced light absorption across the visible region for rGO-TiO₂. The addition of rGO extends visible light adsorption properties, may be attributed to the chemical bonding between TiO₂ and rGO, formation of Ti-O-C bond, similar to C-doped TiO₂ composites [79, 99]. Visible light adsorption is further improved by adding rGO into Pt-TiO₂, due to the
overlapping of extended adsorption properties by rGO and Pt plasmonic adsorption. This simultaneous modification of Pt and rGO could attribute to enhance the photoexcitation efficiency of TiO$_2$ and therefore its photocatalytic activity in theory.

Figure 2-6 UV-vis diffuse reflectance spectra (A) and Kubelka-Munk transformed reflectance spectra (B) of a) TiO$_2$, b) Pt-TiO$_2$, c) rGO-TiO$_2$, d) Pt/rGO-TiO$_2$. (A) shows simultaneously the absorbance spectra of 3DOM catalysts (left) and the reflectance spectrum of 3DOM TiO$_2$ with stop band maxima at 400 nm (right)

The band gap energy ($E_g$) of photocatalysts are estimated according to the Kubelka-Munk equation: $\alpha h\nu=\text{const}(h\nu-E_g)^2$, where $\alpha=(1-R)^2/2R$, $R=10^{-A}$ and $A$ is an optical absorption [78, 99]. Plotting $(\alpha h\nu)^{1/2}$ vs $h\nu$ based on the spectral response in Figure 2-6(A) gives the extrapolated intercept corresponding to the $E_g$ value, as shown in Figure 2-6(B). $E_g$ value of 3DOM TiO$_2$ was calculated as 2.90 eV, which is smaller than anatase TiO$_2$ (3.2 eV). $E_g$ values of Pt-TiO$_2$, rGO-TiO$_2$ and Pt/rGO-TiO$_2$ catalysts are 2.75, 2.59 and 2.67 eV, which are much smaller than that of 3DOM TiO$_2$ (2.90 eV) [48, 100]. After modification of Pt or rGO, the band gap was significantly narrowed in these hybrid materials, due to electronic interaction between TiO$_2$ and Pt nanoparticles and/or rGO [79].

2.3.2 Photodegradation of methyl orange

The photocatalytic performance of the samples has been examined in terms of degradation of MO in an aqueous solution under visible-light irradiation. The degradation curves of MO over
time are shown in Figure 2-7(A). The photocatalytic performance can be affected by many factors, including light adsorption, electron-hole pairs life, morphology, crystallinity, crystallite size, etc. The overall photocatalytic performance results from a synergistic effect of several key factors [101]. 3DOM structure not only extends TiO$_2$ light adsorption, but also provides more active surface area and improved mass transfer, therefore, in 120 min about 25% MO degraded over 3DOM TiO$_2$. 3DOM TiO$_2$ modified by only Pt NPs or rGO showed enhanced photocatalytic activity, and by comparison, rGO-TiO$_2$ (53% removal) exhibited better photocatalytic performance than Pt-TiO$_2$ (32% removal), which was presumably benefited from the high charge separation and adsorption capacity of rGO relative to Pt NPs. Pt/rGO-TiO$_2$ showed highest photocatalytic degradation efficiency about 80% after 120 min, indicating a synergetic effect between Pt NPs and rGO.

A first-order kinetic model was employed to fit the degradation data by using the linear transformation: $\ln(C_0/C) = kt$, where $C_0$ and $C$ are MO concentrations initially and after time $t$ respectively and $k$ (min$^{-1}$) is the kinetic constant. The rate constants were determined from plot of $\ln(C_0/C)$ versus irradiation time and presented in Figure 2-7(B). As can be seen, MO degradation
kinetic constants over 3DOM TiO$_2$, Pt-TiO$_2$, rGO-TiO$_2$ and Pt/rGO-TiO$_2$ are about 0.0021, 0.0032, 0.0062 and 0.0129 min$^{-1}$, respectively. The photodegradation reaction kinetic constant over Pt/rGO-TiO$_2$ was improved about 6, 4 and 2 times, comparing to those of 3DOM TiO$_2$, Pt-TiO$_2$ and rGO-TiO$_2$, respectively. In accord with UV-vis diffuse reflectance spectra, Pt nanoparticle and rGO modified 3DOM TiO$_2$ showed improved photocatalytic activity, resulted from extended light response. What’s more, simultaneous modification of Pt nanoparticles and rGO further improved light absorption and charge transfer, improved MO photodegradation by 6 times than pristine 3DOM TiO$_2$.

Figure 2-8 Recycling test of MO degradation over Pt/rGO-TiO$_2$ within 90 min visible light irradiation

The reusability of Pt/rGO-TiO$_2$ catalyst was tested by the photodegradation of MO under visible light. Pt/rGO-TiO$_2$ catalyst exhibits stable photocatalytic activity as shown in Figure 8, after three runs, the photodegradation efficiency reached 66% within 90 min, which is 5% lower than the first round. The recycling test demonstrating the good stability and high reusability of the catalyst, which may favor practical applications in water treatment.
2.3.3 Mechanism

As shown in results, the enhanced photocatalytic activity of MO photodegradation over Pt/rGO-TiO$_2$ composite can be ascribed to improved visible light absorbance and improved charge transfer efficiency.

First, the slow photon enhancement of 3DOM structure plays an importance role for improving photocatalytic activity. Based on the UV-vis DRS spectra, the introduction of Pt NPs and rGO can effectively extend the spectral response from UV to visible region. TiO$_2$ can be excited by visible light to generate electrons and holes. In the composite system, rGO acts as an acceptor of photogenerated electrons from TiO$_2$ and transporter to separate the photogenerated electron-hole pairs [30, 31, 79, 89, 90, 95, 102]. The work function of rGO (4.4 eV) is higher than the energy level of conduction band (CB) of anatase TiO$_2$ (4.2 eV), charge transfer from TiO$_2$ to rGO is thermodynamically favorable than recombination with holes [103, 104]. Furthermore, when Pt and rGO are in contact, the photogenerated electrons transfer from rGO to Pt is energetically favorable, due to higher work function of Pt (5.64 eV) than rGO [103, 104]. Being a high work function material with excellent electron-accepting and transport properties, rGO acts as a superhighway for the transportation of the electrons, leading to hole-electron separation. Consequently, the vectorial electron transfer route of Pt/rGO-TiO$_2$ can be realized along TiO$_2$→rGO→Pt [105].

Meanwhile, due to the formation of Schottky barrier at the interface of Pt nanoparticles and TiO$_2$, and larger working function of Pt than CB energy level of TiO$_2$, Pt nanoparticles can capture the generated electrons from TiO$_2$ [23, 96]. As a result, the electron-hole pairs efficiently separate and lifetime of the charge carriers is effectively lengthened. The assumed synergism between Pt
nanoparticles, rGO and 3DOM TiO₂ and the possible charge transfer mechanism in the Pt/rGO-TiO₂ composite is presented in Figure 2-9.

![Diagram of charge separation and transportation in Pt/rGO-TiO₂ photocatalyst under light illumination.](image)

Figure 2-9 Schematic illustration of charge separation and transportation in Pt/rGO-TiO₂ photocatalyst under light illumination.

Briefly, photogenerated electrons transfer from TiO₂ through rGO to Pt nanoparticles, or directly to Pt nanoparticles though junction of Pt/TiO₂. The accumulated electrons are good reductants that could be transferred to dissolved oxygen to form superoxide anion radical (O₂⁻). Then O₂⁻ subsequently transformed to active oxygen species, such as •OH, HO₂⁻ and H₂O₂ [22, 70]. The remaining holes at the valence band (VB) of TiO₂ with a strong oxidation power reacted with adsorbed water or surface hydroxyl groups (-OH) to produce •OH [96, 106]. These active species possess a high oxidation power that will result in the oxidative degradation of MO [107].
Under visible light irradiation, holes play a predominant role over hydroxyl radicals in degradation of dye molecules [101]. The reaction processes are summarized as follows:

\[
\begin{align*}
\text{TiO}_2 + hv & \rightarrow e^- + h^+ \\
e^- + \text{rGO} & \rightarrow \text{rGO}(e^-) \\
e^- + \text{Pt} & \rightarrow \text{Pt}(e^-) \\
\text{rGO}(e^-) + \text{Pt} & \rightarrow \text{rGO} + \text{Pt}(e^-) \\
\text{Pt}(e^-) + O_2 & \rightarrow O_2^* \\
(2e^- + O_2 + 2H^+ & \rightarrow H_2O_2) \\
(H_2O_2 + O_2^* & \rightarrow \cdot OH + OH^- + O_2) \\
(O_2^* + H^+ & \rightarrow HO_2^*) \\
O_2^* + H_2O & \rightarrow \cdot OH + HO_2^* + H_2O \\
h^+ + H_2O/-OH & \rightarrow \cdot OH + H^+ \\
\cdot OH/h^+ + \text{MO} & \rightarrow \text{degradation products}
\end{align*}
\]

As discussed in the proposed mechanism, both rGO and Pt nanoparticles play important roles in improving the photocatalytic activity of 3DOM TiO\textsubscript{2}. Moreover, MO molecules are concentrated near the catalyst surface due to the high adsorption capacity of rGO, further promoted photocatalytic reaction over the ternary Pt/rGO-TiO\textsubscript{2} composite [31, 44, 79, 108].

2.4 Conclusion

In summary, by incorporating Pt nanoparticles and rGO with 3DOM TiO\textsubscript{2}, a new effective photocatalyst was facilely synthesized. The excellent charge-separation property and high adsorption capacity of rGO increased the charge carrier lifetime and affinity to organic molecules. The introduction of Pt nanoparticles increased spectral response to visible light through surface plasmon resonance and suppressed charge recombination. The synergism between rGO and Pt
nanoparticles results in the novel ternary photocatalyst effective to remove MO under visible light, the photodegradation rate toward MO of resulting material is 6 times higher than 3DOM TiO$_2$. Moreover, the photocatalysis process is stable and reusable, making it a promising photocatalyst for practical water treatment.
CHAPTER 3 Carbon coated Fe₂O₃/TiO₂ composite as anode material for LIBs

3.1 Introduction

As mentioned in Chapter 1.3, titanium dioxide (TiO₂) is an anode material in LIBs that shows promise in high-power applications due to its highly reversible lithiation process, lack of solid electrolyte interface (SEI) formation, and relatively small volume expansion. TiO₂ is also a highly abundant material, relatively non-toxic, and chemically stable [109]. 3DOM TiO₂ has an interconnected structure which allows for efficient mass transfer, electron transfer, and contact between the electrode and electrolyte [109].

Among transition metal oxides, iron oxides are attractive anode materials for rechargeable LIBs because their high theoretical capacities of about 900-1000 mAh g⁻¹, nontoxicity, low cost and worldwide abundance. The lithium storage mechanism of iron oxides is based on a redox conversion reaction, where iron oxides are reduced to metallic nanoclusters dispersed in a Li₂O matrix upon lithiation and are then reversible restored to their initial oxidation states during delithiation. For example, the reaction mechanism of Fe₂O₃ can be described as: Fe₂O₃ + 6Li⁺ + 6e⁻ ↔ 3Li₂O + 2Fe [110]. However, capacity degradation of Fe₂O₃ anode due to its large volume expansion, low conductivity limits its application in LIBs. To overcome capacity degradation of iron oxide, optimization on Fe₂O₃ nanomaterials with different structures and modification have been studied as anode material for LIBs. Mesoporous α-Fe₂O₃ showed a high specific capacity of 1360 mAh g⁻¹ with excellent cycling stability (50 cycles) and high rate capacity [111]. Reddy et al. synthesized α-Fe₂O₃ nanoflakes on Cu foil through thermal treatment, the resulting material exhibited a stable capacity of 680 mAh g⁻¹ with no noticeable capacity fading up to 80 cycles [112]. Lin et al. Reported hydrothermally synthesized single-crystalline α-Fe₂O₃ nanorods, which
retained capacities after numerous cycles. The nanorod structure provided a short path for lithium-ion diffusion and effective accommodation of the strain generated from volume expansion [113].

Three-dimensional (3D) nanostructure enhances diffusion kinetics for lithium storage by ensuring better lithium ion transport across the electrode and more efficient electron transport. 3D structure also provides tolerance for the volume variation of the entire electrode during lithium storage. Yu employed TiO$_2$ nanotube arrays as 3D substrate for Fe$_2$O$_3$ nanorods, and such hierarchical architecture exhibited capacity of 400 mAh g$^{-1}$ over at for 50 cycles [114]. Nanosized Fe$_2$O$_3$ were loaded on single-walled carbon nanotube membrane were used as a flexible, binder-free anode of LIBs, showed a high reversible capacity of 1243 mAh g$^{-1}$ at a current density of 50 mA g$^{-1}$ and an excellent cyclic stability over 90 cycles at 500 mA g$^{-1}$ [115].

The introduction of carbonaceous materials has been generally accepted due to their high electronic conductivity, excellent buffering effect and mechanical strength [116]. Carbon coated Fe$_2$O$_3$ nanoparticles was loaded onto graphene nanosheets, the resulting material showed reversible capacity of 900 mAh g$^{-1}$ after 50 cycles, and 82% capacity retention rate [116]. Carbon coated Fe$_3$O$_4$ coaxial nanotubes showed excellent cycling performance with a specific capacity of 1020 mAh g$^{-1}$ after 150 cycles. The carbon coating on the surface of Fe$_3$O$_4$ nanotube improved the electronic conductivity, also prevented direct contact between organic solvent and Fe$_3$O$_4$, which contributed to high capacity retention [117]. Besides, carbon coating on Fe$_2$O$_3$ rods provided a protection for active materials from structural destruction and agglomeration, porous Fe$_2$O$_3$@C rods exhibited rate capability and cyclic stability as anode of LIBs, capacity retained at 639 mAh g$^{-1}$ after 500 cycles at 500 mA g$^{-1}$ [118].
Herein, we design a new anode material for LIBs, in which nanosized Fe$_2$O$_3$ are assembled onto 3DOM TiO$_2$ and further modified with carbon coating, for a high energy storage capacity and long-term stability.

3.2 Experimental

![Diagram showing fabrication processes of carbon coated TiO$_2$@Fe$_2$O$_3$ material](image)

Figure 3-1 Schematic illustration of fabrication processes of carbon coated TiO$_2$@Fe$_2$O$_3$ material

3.2.1 Chemicals and materials

All chemicals were reagent grade and used without further purification. Methyl methacrylate (MMA), titanium butoxide (Ti(OBu)$_4$), acetylacetone (ACAC), ethanol, iron chloride hexahydrate (FeCl$_3$.6H$_2$O), glucose were purchased from Sigma-Aldrich. Potassium persulfate ($K_2S_2O_8$) was purchased from Alfa Aesar. Deionized (DI) water (resistivity > 18.0 MΩ) was used for all experiments.

3.2.2 Preparation of PMMA and 3DOM TiO$_2$

Preparation of PMMA template and 3DOM TiO$_2$ are the same with method mentioned in Chapter 2.2.2 and 2.2.3

3.2.3 Preparation of TiO$_2$@Fe$_2$O$_3$

The growth of β-FeOOH nanospindles on 3DOM TiO$_2$ was conducted by immersing the as-obtained 3DOM TiO$_2$ in 250 mL of aqueous FeCl$_3$ solution (80 mM) at 80 °C for 6 h. The final products were washed with DI water several times before being fully dried in air at 60 °C. Then the TiO$_2$@FeOOH composite is annealed in air at 400 °C for 3h with a slow ramp rate of 1 °C/min.
3.2.4 Preparation of carbon coated TiO$_2$@Fe$_2$O$_3$

Certain amount of TiO$_2$@Fe$_2$O$_3$ was immersed in 10 mL glucose solution (containing same amount of glucose) for 24 h, and dried at 60 °C in air, followed by annealing at 500 °C for 3 h under Ar flow, with ramp rate of 5 °C min$^{-1}$. The carbonization temperature was chosen as 500 °C, because at this temperature the generated carbon could be partially graphitized to benefit the LIB performance [29].

3.2.5 Coin cell fabrication

The Li-ion battery electrode was made by mixing the carbon coated TiO$_2$@Fe$_2$O$_3$ composite with carbon black and alginate binder to form a uniform slurry in a weight ratio of 8:1:1 and then spread on a copper foil using a stainless-steel blade. The electrode was dried at 60 °C in a vacuum oven for 10 h. CR2032 coin cells then were assembled in an Ar-filled glovebox using the as-prepared carbon coated TiO$_2$@Fe$_2$O$_3$ anodes as working electrodes and lithium metal foil as counter electrodes. The electrolyte was 1 M LiPF$_6$ in the ethylene carbonate/ethyl methyl carbonate in 1:1 volume ratio. A Celgard 20 µm-thick monolayer polyethylene membrane was used as a separator.

3.2.6 Material Characterization

The morphology of sample was imaged with a Hitachi S-4800 scanning electron microscope (SEM) and a Hitachi H9000NAR transmission electron microscope (TEM). The crystal structure and phase state of obtained composites were determined by X-ray diffractometry (XRD) using Bruker D8 Discover A25 diffractometer with copper K$_\alpha$ radiation. Raman spectra were taken on a Reinshaw 1000B Raman spectrometer with a 632.8 nm He-Ne laser source. X-ray photoelectron spectroscopy (XPS) was employed to characterize the chemical states of prepared photocatalyst using Perkin Elemer PHI 5440 ESCA system with an Al K$_\alpha$ X-ray source. The
charge-discharge cycles were performed between 0.05-3 V at a current of 50 mAh g\(^{-1}\) for 100 cycles with LANHE CT2001A (Land, China).

3.3 Results and discussion

3.3.1 Synthesis and characterization

The synthesis process of carbon coated TiO\(_2@\)Fe\(_2\)O\(_3\) composite is straightforward, as schematically illustrated in Figure 3-1. 3DOM TiO\(_2\) is firstly synthesized as nanostructured 3D substrate through CCT and sol-gel method. Then β-FeOOH nanospindles are spontaneously grown and assembled in microporous structure of 3DOM TiO\(_2\) by hydrolysis of FeCl\(_3\). After thermal dihydroxylation of β-FeOOH nanospindles on 3DOM TiO\(_2\) at 400 °C, the resulting composite is coated with carbon using glucose as carbon source. A combination of techniques, described individually in detail in the Experimental Section, were used in the synthesis process, including CCT, sol-gel processing, hydrolysis.

The morphological characteristics of 3DOM TiO\(_2\), TiO\(_2@\)Fe\(_2\)O\(_3\) and carbon coated TiO\(_2@\)Fe\(_2\)O\(_3\) are characterized using SEM, as shown in Figure 3-2. PMMA (330 nm) spheres are uniform in size and effectively packed into face-centered-cubic (FCC) structure. Each colloidal particle is connected with 6 surrounding particles through 6 bridges on the two-dimensional plane (Figure S2-1). 3DOM TiO\(_2\) exhibits a highly ordered microporous structure, which replicate the three-dimensional closely-packed PMMA opals in Figure 3-2(a-b), with pore diameter about 210 nm and 25 nm wall thickness. The pore size of 3DOM TiO\(_2\) is smaller than PMMA spheres (~330 nm) due to a shrinkage during the calcination. The small windows between the opal pores can be clearly observed in Figure 3-2(b), which replicates from the contact bridges between the neighboring PMMA spheres. The underlying layer of pores and porous walls can be clearly
observed, indicating that the sample possesses a three-dimensionally well-open, ordered and interconnected microporous network.

Figure 3-2 SEM images of (a)-(b) 3DOM TiO$_2$, (c)-(d) TiO$_2$@Fe$_2$O$_3$, (e)-(f) carbon coated TiO$_2$@Fe$_2$O$_3$

After the reaction of FeCl$_3$ solution and calcination, the surface of 3DOM TiO$_2$ is covered with large quantities of Fe$_2$O$_3$ nanospindles as a result of the hydrolysis of Fe$^{3+}$ and simultaneously
heterogeneous growth of FeOOH on 3DOM TiO$_2$, as shown in Figure 3-2(c-d). The Fe$_2$O$_3$ nanospindles were with diameter about ~10 nm, while length varied from ~30 to ~100 nm. From magnified SEM image Figure 3-2(d), it can be seen that Fe$_2$O$_3$ nanospindles well dispersed on to TiO$_2$ macropores and walls, which provide open channels and more contact area for the transportation of lithium ions and electrons [119]. The TiO$_2$@Fe$_2$O$_3$ composite was further coated with a thin layer of amorphous carbon using glucose as carbon source, as seen in Figure 3-2(e-f), while the structure of Fe$_2$O$_3$ nanospindles and 3DOM TiO$_2$ can still be observed.

TEM analysis was also conducted to further investigate the morphology of TiO$_2$@Fe$_2$O$_3$ and carbon coated TiO$_2$@Fe$_2$O$_3$. From Figure 3-3(a) the macropores in hexagonal arrangement can be easily observed, and the porous skeleton of 3DOM TiO$_2$ are composed of many nanoparticles. In the pores of 3DOM TiO$_2$, well separated Fe$_2$O$_3$ nanospindles are grafted on the wall of pores, the size is consistent with observation in SEM images. HRTEM image (Figure 3-3(b)) shows the lattice fringes of both TiO$_2$ and Fe$_2$O$_3$, which indicates the highly crystalline nature of TiO$_2$ nanoparticles and Fe$_2$O$_3$ nanospindles. The lattice fringe with a d-spacing of 0.352 nm can be assigned to the (101) lattice plane of anatase TiO$_2$, and the fringe with d-spacing of 0.36 nm belongs to the (012) lattice plane of $\alpha$-Fe$_2$O$_3$ nanospindles. After carbon coating, there is no apparent change in the morphology of the products, as shown in Figure 3-3(c). TEM characterization indicates that the entire surface of TiO$_2$@Fe$_2$O$_3$ composite has been covered with a uniform and continuous amorphous carbon over with a thickness of a few nanometers. HRTEM image (Figure 3-3(d)) shows a high crystallized nanospindle covered by a thin layer of amorphous carbon, the lattice fringe with d-spacing of 0.25 nm, which can be assigned to (110) lattice plane of $\alpha$-Fe$_2$O$_3$ [111].
Figure 3-3 (a) TEM image and (b) HRTEM image of TiO$_2$@Fe$_2$O$_3$, (c) TEM image and (d) HRTEM image of carbon coated TiO$_2$@Fe$_2$O$_3$

The structure of obtained composites was characterized by XRD, as shown in Figure 3-4(a). XRD patterns show the main phase of TiO$_2$ in all samples are anatase (JCPDS 21-1272), characteristic peaks at 25.3°, 36.9°, 37.8°, 48.0°, 53.1°, 55.1°, 62.7°, 68.8° can be assigned to the plane (101), (004), (200), (105), (211), (204), (116) of anatase, respectively. After hydrolysis of Fe$^{3+}$ ions, the new peaks are assigned to β-FeOOH (JCPDS 75-1594), characteristic peaks at 11.9°,
16.9°, 26.9°, 34.2°, 39.4°, 46.5°, 55.4° and 61.3° can be assigned to the plane (110), (200), (130), (400), (211), (301), (411) (620) and (002) of β-FeOOH, respectively. The transform from β-FeOOH to hematite (JCPDS 33-0664) was confirmed by XRD result, the peaks at 24.1°, 33.1°, 35.6°, 40.8°, 49.5°, 54.1°, 57.6°, 62.5° and 64.0°, which are corresponding to plane (012), (104) (110), (113), (024), (116), (018), (214) and (300) of hematite.

Figure 3-4 (a) XRD patterns of TiO₂, TiO₂@FeOOH and TiO₂@Fe₂O₃, (b) Raman spectrum of carbon coated TiO₂@Fe₂O₃.

The Raman spectrum of carbon coated TiO₂@Fe₂O₃ shown in Figure 3-4(b) contains characteristic wide of carbon D and G bands of carbon at approximately 1350 and 1590 cm⁻¹ [120, 121]. The D band is associated with disorder, allowing by zone edge modes of the graphite structure to become active due to the lack of long-range order in amorphous and quasi-crystalline forms of carbon materials. The G band corresponds to the stretching vibrations of sp² bonded carbon atoms in the basal plane of the crystalline graphite. From the Raman spectrum, the intensity of G band is higher than that of D band, showing the presence of sp² carbon in the carbon coating on surface of TiO₂@Fe₂O₃ composite [118]. The peak intensity ratio I_D/I_G can be used to evaluate
the degree of crystallinity of carbon materials [122]. The \( \frac{I_D}{I_G} \) obtained from Raman spectrum is 0.8, indicating of the high carbonization, which contributes to electronic conductivity.

Figure 3-5 Fitted XPS spectra (a) Fe 2p, (b) Ti 2p, (c) C 1s and (d) O 1s of carbon coated TiO2@Fe3O4 composite

To further characterization the chemical composition of the resulting carbon coated TiO2@Fe3O4 composite, XPS was conducted. As exhibited in XPS spectrum (Figure 3-5(a)), the peaks at 710.6 and 724.1 eV in the Fe 2p spectrum are attributed to Fe 2p3/2 and Fe 2p1/2, respectively [123, 124]. In addition, weak shakeup satellites are also observed at the higher binding energy side of the main peaks in the Fe 2p spectrum, as indicated by arrows in Figure 3-5(a), which is characteristic of Fe\(^{3+}\) in Fe3O4 [125]. Figure 3-5(b) shows the Ti 2p XPS spectrum, in which two
peaks at 458.7 and 464.6 eV can be assigned to Ti 2p3/2 and 2p1/2, respectively [126]. The C 1s spectrum, shown as Figure 3-3(c), can be fitted into three component peaks centered at 284.5, 286.0 and 287.9 eV, corresponding to sp2-bonded carbon (C-C), epoxy/hydroxyls (C-O-C/C-O-H) carbon atoms and carbonyls carbon (C=O), respectively [120, 127]. As shown in Figure 3-5(d), the O 1s spectrum ranging from 536 to 528 eV can be deconvoluted into two components. The dominant peak at 529.9 can be assigned to lattice oxygen connected to Ti4+ an Fe3+. The peak at 532.1 eV can be assigned to C=O bond [128].

3.3.2 Battery performance

To evaluate the electrochemical performance of obtained composites, the electrodes were prepared following practical industrial procedures: mixed with conductor (carbon black) and binder, then coated onto copper foil using automatic thick film coater.

The principal reaction of reversible lithium insertion/extraction into anatase TiO2 can be described as: TiO2 + xLi+ + xe− ↔ LixTiO2 (3.1), with the maximum insertion coefficient x determined to be 0.5 due to its poor electronic conductivity and lithium ion diffusivity [129]. The theoretical charge storage capacity of anatase TiO2 is estimated as 167.5 mAh g⁻¹ [130].

The electrochemical reactions involved in the above processes can be described as follows:

\[
\begin{align*}
\text{Fe}_2\text{O}_3 + \text{Li}^+ + e^- & \leftrightarrow \text{LiFe}_2\text{O}_3 & (3.2) \\
\text{LiFe}_2\text{O}_3 + \text{Li}^+ + e^- & \leftrightarrow \text{Li}_2\text{Fe}_2\text{O}_3 & (3.3) \\
\text{Li}_2\text{Fe}_2\text{O}_3 + 4\text{Li}^+ + 4e^- & \leftrightarrow 2\text{Fe} + 3\text{Li}_2\text{O} & (3.4)
\end{align*}
\]

Cyclic voltammetry (CV) was used to examine the behavior for the lithium ion intercalation of obtained composites. Figure 3-6 displays the representative cyclic voltammograms (CVs) for the first four cycles of 3DOM TiO2, TiO2@Fe2O3 and carbon coated TiO2@Fe2O3 electrodes, at a scan rate of 1 mV s⁻¹ in the voltage window of 0.05-3.2 V. In Figure 3-6(a), two
well defined current peaks of TiO$_2$ are observed at 1.55 V (cathodic sweep) and 2.2 V (anodic sweep). The peak at 1.55 V corresponds to the biphasic transition from tetragonal anatase to orthorhombic Li$_{0.5}$TiO$_2$ when the insertion coefficient $x$ reaches $\sim 0.5$. This result is consistent with the results reported in the literature [129-131]. For TiO$_2$@Fe$_2$O$_3$, the cathodic peaks (first sweep) located around 0.45 V and 1.25 V are observed, the peak at 1.25 eV corresponds to the lithium intercalation into Fe$_2$O$_3$ (Equation (3.1) and (3.2)), the second peak at 0.45 V is attributed to the further reduction of the Li$_2$(Fe$_2$O$_3$) into Fe$^0$ and the formation of LiO$_2$ and SEI film [27, 119].

There are two oxidation peaks at 1.7 and 2.4 V corresponding to the extraction of Li ions from Fe$_2$O$_3$ and TiO$_2$ from TiO$_2$@Fe$_2$O$_3$ electrode. In the subsequent cycles, the reduction peak at 1.2 V disappears and the intensity of peak at 0.6V decreases, indicating the occurrence of some irreversible processes in the electrode materials in the first cycle ascribed to the formation of SEI film.

The CV curves of carbon coated TiO$_2$@Fe$_2$O$_3$ showed similar peaks to that of TiO$_2$@Fe$_2$O$_3$, as shown in Figure 3-6(c). At the first cycle, three reduction peaks at 0.4, 0.8 and 1.2 V are observed, corresponding to the insertion of lithium into the TiO$_2$@Fe$_2$O$_3$ and complete reduction of Fe$_2$O$_3$ to metallic Fe and SEI film formation, respectively [114]. The peaks at 1.7 and 2.2 V are ascribed to the restoration of iron oxides and the delithiation of TiO$_2$. The slight shift of the two cathodic peaks to higher potentials after the first cycle is probably related to some possible activation process of the Li ion insertion in the first cycle. The subsequent curves show good reproducibility in the following cycle, demonstrating the good reversibility of the TiO$_2$@Fe$_2$O$_3$ electrode. The lithium storage mechanism in carbon coated TiO$_2$@Fe$_2$O$_3$ is similar to that in iron oxide and TiO$_2$ electrodes.
Figure 3-6 CVs at a scan rate of 1 mV s$^{-1}$ for the first four cycles of (a) 3DOM TiO$_2$, (b) TiO$_2$@Fe$_2$O$_3$, (c) carbon coated TiO$_2$@Fe$_2$O$_3$ electrodes
In order to test the applicability of the new composites in LIBs, the electrochemical lithium storage properties of carbon coated TiO$_2$@Fe$_2$O$_3$ were evaluated. The electrochemical properties of bare 3DOM TiO$_2$ and TiO$_2$@Fe$_2$O$_3$ were also investigated for comparison. Figure 3-7(a) shows the cycling performance of TiO$_2$, TiO$_2$@Fe$_2$O$_3$ and carbon coated TiO$_2$@Fe$_2$O$_3$ at a current density of 50 mA g$^{-1}$. The bare 3DOM TiO$_2$ electrode only delivers a low capacity with its initial discharge and charge capacity around 636 and 322 mAh g$^{-1}$, respectively. In the first 20 cycles, the discharge and charge capacity decreased to about 150 mAh g$^{-1}$. However, after 20 cycles the 3DOM TiO$_2$ electrode showed excellent cycling stability, which still delivered a capacity around 150 mAh g$^{-1}$. The TiO$_2$@Fe$_2$O$_3$ electrode on the one hand has relatively high initial discharge and charge capacities capacity around 1280 and 645 mAh g$^{-1}$, respectively, on the other hand, it suffered from degradation. As seen in Figure 3-7(a), the discharge capacity of the TiO$_2$@Fe$_2$O$_3$ electrode firstly decreased from 645 to 275 mAh g$^{-1}$ in 25 cycles. After the carbon coating, the cyclic stability was significantly enhanced. The carbon coated TiO$_2$@Fe$_2$O$_3$ electrode showed comparable initial discharge and charge capacities to that of TiO$_2$@Fe$_2$O$_3$, about 1155 and 570 mAh g$^{-1}$, respectively.
From the second cycle onwards, the discharge and charge capacity of carbon coated TiO$_2$@Fe$_2$O$_3$ retained about 575 and 565 mAh g$^{-1}$ after 100 cycles, respectively. It is well known that the electrochemical performance is not only highly dependent on the intrinsic crystalline texture and surface properties, but also greatly related to the morphology and assembled structure of active materials. In carbon coated TiO$_2$@Fe$_2$O$_3$ structure, the Fe$_2$O$_3$ nanospindles are largely separated from each other because they are grown directly on the 3DOM TiO$_2$, which improves the accessibility of lithium ions in the electrolyte [26]. The continuous and elastic amorphous carbon overlayer further strengthens the structural integrity of TiO$_2$@Fe$_2$O$_3$ structure, which will improve the cycle life.

In addition to the cycling performance, the high-rate capacity is another important parameter for LIB application, especially for high-power batteries. To further evaluate the rate capability, the batteries using TiO$_2$, TiO$_2$@Fe$_2$O$_3$ and carbon coated TiO$_2$@Fe$_2$O$_3$ as active materials were tested under various current density from 50 mA g$^{-1}$ to 800 mA g$^{-1}$, as Figure 3-7(b) presents. The carbon coated TiO$_2$@Fe$_2$O$_3$ electrode exhibits a reasonably good cycling response at various current rates. Even at a high current density of 800 mA g$^{-1}$, carbon coated TiO$_2$@Fe$_2$O$_3$ electrode can also deliver a reversible capacity of 250 mAh g$^{-1}$, which is much higher compared to TiO$_2$@Fe$_2$O$_3$ and TiO$_2$ (~100 mAh g$^{-1}$). The improved performance is contributed to rationally designed hierarchical nanostructures, the excellent stability of TiO$_2$ and the presence of carbon coating.

Figure 3-8 presents the discharge-charge profiles of 3DOM TiO$_2$, TiO$_2$@Fe$_2$O$_3$ and carbon coated TiO$_2$@Fe$_2$O$_3$ electrodes at a rate of 50 mA g$^{-1}$, at 1, 5, 10, 25, 50, 75, 100 cycles.
Figure 3-8 Galvanostatic discharge/charge profiles of (a) 3DOM TiO$_2$, (b) TiO$_2$@Fe$_2$O$_3$, (c) carbon coated TiO$_2$@Fe$_2$O$_3$ electrodes at a constant current density of 50 mA g$^{-1}$. 
For the 3DOM TiO\textsubscript{2} electrode, the first lithium insertion capacity is observed to be 635 mAh g\textsuperscript{-1} from Figure 3-8(a). The larger capacity obtained may be ascribed to a shorter diffusion length for both the electron and Li\textsuperscript{+}, and a larger electrode/electrolyte contact area of 3DOM TiO\textsubscript{2} compared with traditional materials, which facilitate the lithium ions’ insertion and extraction. The discharge plateau in the discharge/charge curve can be attributed to the electrochemical behavior of the thin-film electrode [130].

As shown in Figure 3-8(b), in the discharge curve of the first cycles, there are three voltage plateaus (~1.75, ~0.8, ~0.6 V), resulting from the lithium reactions with TiO\textsubscript{2} and Fe\textsubscript{2}O\textsubscript{3} nanomaterials [127]. The discharge-charge profiles are in good agreement of CV measurement. The first plateau at ~1.75 V is associated with the phase transition between the tetragonal TiO\textsubscript{2} and orthorhombic Li\textsubscript{x}TiO\textsubscript{2} phases. The second plateau at ~0.8 V attributed to lithium insertion into the crystal structure of Fe\textsubscript{2}O\textsubscript{3} and the formation of cubic LiFe\textsubscript{2}O\textsubscript{3}. An extended potential plateau appears at 0.6 V, corresponding to the reduction of Fe\textsuperscript{2+} to Fe\textsuperscript{0} and the formation of amorphous Li\textsubscript{2}O. In the following delithiation process, a broad voltage plateau centered at ~1.7 V is observed, indicating the oxidation of Fe\textsuperscript{0} to Fe\textsuperscript{3+} [128]. It can be seen that the first discharge and charge capacities are ~1280 and ~645 mAh g\textsuperscript{-1}, respectively. The electrode suffered from discharge and charge capacity degradation during cycling. In the first 25 cycles, the capacities dropped fast and maintained at a lower capacity about 230 mAh g\textsuperscript{-1}.

In Figure 3-8(c), the plateau voltages of carbon coated TiO\textsubscript{2}@Fe\textsubscript{2}O\textsubscript{3} electrode in the first cycle are at (~0.4, ~0.8, ~1.2 V) were in good agreement with the oxidation and reduction peaks the CV curves (Figure 3-6(c)). The profiles of charge-discharge were rather different during the first cycles but became similar during the following cycling. Starting from the second cycle the discharge and charge capacities were stable at around 570 mAh g\textsuperscript{-1}. 

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3.4 Conclusion

A novel composite was successfully prepared by grafting Fe$_2$O$_3$ nanospindles onto 3DOM TiO$_2$, followed by carbon coating using glucose as carbon source. The carbon coated TiO$_2$@Fe$_2$O$_3$ material was applied as electrode mater in LIBs and showed good electrochemical performance. The composite exhibits a large reversible capacity about 570 mAh g$^{-1}$, which is about four times of the reversible capacity of 3DOM TiO$_2$. In addition to the high reversible capacity, the obtained material also exhibits good cycle performance and superior rate capacity. This rationally designed composite benefits from both good stability of TiO$_2$, high capacity of Fe$_2$O$_3$, and good electron conductivity of carbon.
CHAPTER 4  Adsorption removal of phosphorus by zirconium modified clays

4.1 Introduction

Wastewater from municipalities and industries may contain trace amounts of phosphate [59]. Phosphorus is an essential macronutrient that encourages the growth of photosynthetic algae and cyanobacteria. Excessive amounts of phosphorus entering water bodies could cause eutrophication and subsequently stimulate blue-green algae blooms under proper conditions [132]. The removal of phosphorus is essential to prevent eutrophication of natural water bodies, especially freshwaters.

As reviewed in Chapter 1, adsorption is one of the most effective method for the removal of phosphate in water, and nanosized metal oxides have been found wide environmental applications in the adsorptive removal of contaminants in water. Among a various of metal (hydr)oxides, zirconium (hydr)oxide has strong surface complexing ability for phosphate, which make it a superior adsorbent toward phosphate [61].

Defined by the joint nomenclature committees (JNCs), clay is naturally occurring material composed primarily of fine-grained minerals, which is generally plastic at appropriate water contents and will harden when dried or fired [133]. Clays are hydrous aluminosilicates broadly defined as those minerals that make up the colloid fraction (<2µm) of soil, sediments, rocks and water [134]. The particles of clay minerals may be crystalline or amorphous, platy or fibrous. Clay structure is layered with interlayer space and the layers may be electrically neutral or charged [135]. Large specific surface area, chemical and mechanical stability, layered structure, high cations exchange capacity (CEC), low-cost, an abundant availability, non-toxic nature, etc., have made the clays excellent adsorbent materials.
Clay minerals are phyllosilicates in which the individual layers are composed of a tetrahedral (T) and octahedral (O) sheet. The tetrahedral sheet is a silica tetrahedral layer in which the silicon atom is equidistant from four oxygens or possibly hydroxyls arranged in the form of a tetrahedron with the silicon atom in the center. These tetrahedrons are arranged to form a hexagonal network repeated infinitely in two horizontal directions to form the silica tetrahedral sheet, shown in Figure 4-1(a). The octahedral sheet is comprised of closely packed oxygens and hydroxyls in which aluminum, iron, and magnesium atoms are arranged in octahedral coordination Figure 4-2(b).

![Diagrammatic sketch of the (a) tetrahedral sheet and (b) octahedral sheet](image)

**Table 4-1 Classification of clay minerals according to Grim [136]**

<table>
<thead>
<tr>
<th>Classification</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>I. Amorphous</td>
<td>Allophane group</td>
</tr>
<tr>
<td>II. Crystalline</td>
<td></td>
</tr>
<tr>
<td>A. Two-layer type (sheet structures composed of units of one layer of silica tetrahedrons and one layer of alumina octahedrons)</td>
<td>Kaolinite group, halloysite group</td>
</tr>
<tr>
<td>B. Three-layer type (sheet structures composed of two layer of silica tetrahedrons and one central dioctahedral or trioctahedral layer)</td>
<td>Smectite group, Sodium montmorillonite, Calcium montmorillonite, Bedillite, Vermiculite, Illite group</td>
</tr>
<tr>
<td>C. Regular mixed-layer type (ordered stacking of alternate layers of different types)</td>
<td>Chlorite group</td>
</tr>
<tr>
<td>D. Chain-structure types (hornblende-like chains of silica tetrahedrons linked together by octahedral groups of oxygens and hydroxyls containing Al and Mg atoms)</td>
<td>Sepiolite, Palygorskite (attaoulgite)</td>
</tr>
</tbody>
</table>
Clays are divided into classes or groups such as smectites (montmorillonite, saponite), mica (illite), kaolinite, vermiculite, serpentine, pyrophyllite and sepiolite etc. Table 4-1 show Grim’s classification and lattice structure of clay minerals [136], which gives a basis for the nomenclature and differences between the various clay minerals [137]. The layer structures of different clays are shown as Figure 4-2. Interlayer space is identified as the space between two tetrahedral sheets which can be occupied by cations (such as Na⁺, Ca²⁺, Mg²⁺) and water.

![Diagram of clay structures](image)

**Figure 4-2 Classification of clays [138]**

Modified clays have been extensively studied in various applications, including catalysts and adsorbents. Iron pillared clay was prepared by intercalation of low polymerization degree hydroxy-Fe ions into silicate layer of vermiculite [139]. The iron pillared vermiculite showed favorable photocatalytic properties and long-term stability as photo-Fenton catalyst for removal of synthetic dyes. Two zirconium pillared Portuguese clays were synthesized and studied for methane and carbon dioxide separation, both pillared clays presented high specific surface area and good thermal stability. The zirconium intercalated clays showed higher adsorbed amount of carbon dioxide, which were suitable for methane/carbon dioxide separation [140]. As for applications in water and wastewater treatment, zirconium pillared clay to remove phosphate, arsenic, dyes, and
heavy metals ions. A series of zirconium pillared montmorillonite with different zirconium contents were prepared, the formation of microporous system in the interlayer spaces improved the specific area, pore volume, and basal spacing. The modified montmorillonite was effective to remove arsenate with a high adsorption capacity of over 120 mg/g [141]. An inorganic modified bentonite with Fe ions in the interlayer space was prepared, the phosphate adsorption capacity was improved by ~350% compared to unmodified bentonite at neutral pH [132]. Montmorillonite was intercalated with aluminum and zirconium hydroxy-polycations, the resulting clays were evaluated as adsorbents for the removal of Orange II and methylene blue from aqueous solutions. These pillared clays showed potential as low-cost adsorbents for dye removal from aqueous solutions [65].

To understand the effect of clay structure as support of ZrO₂, three clays were chosen in this study: montmorillonite (MT, 2:1, max expansion), vermiculite (VT, 2:1, some expansion) and kaolinite (KT, 1:1, no expansion).

Montmorillonite is a three-layer (2:1) clay, which is composed of unites made up of two silica tetrahedral sheet with a central alumina octahedral sheet. The net layer charge of montmorillonite is -0.8 charge/unit cell, the resulting negative net charge is balanced by exchangeable cations adsorbed between the unit layers and around their edges. The bonding between two tetrahedral sheets is very week, substitutions mainly take place in the octahedral sheets. Water and exchangeable ions can enter between the two tetrahedral sheets, where is separated from the interlayer spacing where cations balancing the excess negative charge are present [15].

Vermiculite also has a 2:1 type structure, with a medium shrink-swell capacity. Both montmorillonite and vermiculite have high cation exchange capacity due to the substitution of Si⁴⁺
and Al$^{3+}$ [142]. In vermiculite, most of the cation substitutions take place in tetrahedral sheets, such arrangement limits the potential of this mineral to expand its interlayer space.

Kaolinite is a two-layer (1:1) clay, which consists of one tetrahedral sheet of SiO$_4$ and one octahedral sheet with Al$^{3+}$ as the octahedral cation. Both the sheets combine to form a common layer such as the tips of the silica tetrahedra point towards the octahedral layer. There is no interlayer swelling of kaolinite, and the charges within the structural unit are balanced. There is very litter substitution in the kaolinite lattice and kaolinite is the least reactive clay.

In this study, zirconium oxide modified clays with different zirconium oxide contents were prepared and characterized. The phosphate removal performance of zirconium modified clays was examined in detail by both the kinetics and equilibrium adsorption isotherm study, the differences in behavior of MT, VT and KT and their zirconium oxide derivatives towards adsorption of phosphate from aqueous solution.

4.2 Experimental

4.2.1 Chemicals and materials

Kaolinite (KT) was purchased from Millipore, Ca-rich montmorillonite (stx-1b, MT) was obtained from Clay Minerals Society, vermiculite (VT) was obtained from Palabora Mining Co. (South Africa). Zirconium oxychloride octahydrate (ZrOCl$_2$.8H$_2$O, Alfa Aesar) was used as source of zirconium. Sodium dihydrogen phosphate (NaH$_2$PO$_4$, Fisher Scientific), sodium hydroxide (NaOH), sodium carbonate (Na$_2$CO$_3$, Sigma-Aldrich) were used as purchased without further purification. Humic acid (HA) was purchased from Alfa Aeaser. All solution used were prepared with ultrapure water (resistance $> 18.2$ m$\Omega$). Working solutions of NaH$_2$PO$_4$ were prepared from a stock solution of 100 mg P/L to give the required initial concentrations for each experiment.
4.2.2 Pretreatment of vermiculite

Vermiculite was pretreated followed reported method [143]. Briefly, 20 g vermiculite was added into 200 mL of 0.1 M HNO₃ solution, then the mixture was stirred at 90 °C for 3 h. The obtained solid was washed with water until neutral pH and then added into 200 mL of 8 g/L sodium carbonate solution, and kept at 80 °C with vigorous agitation for 3 h. The obtained Na-vermiculite (Na-VT) was washed by ultrapure water until neutral pH. Na-VT was dried at 105 °C and ground.

4.2.3 Preparation of Zirconium modified clays

Zirconium modified clay was prepared according to reported method [141]. Briefly, 5 g clay was firstly dispersed into 100 mL of ultrapure water and continuously stirred for 10 h to obtain a uniform clay aqueous mixture, and followed by adding desired amount of ZrOCl₂.8H₂O into suspension. The ratios between zirconium and clay were varied at 0.31 mmol/g, 0.62 mmol/g, 1.24 mmol/g and 2.48 mmol/g. The resulting suspension was stirred for 8 h. Then the pH of the suspension was adjusted to 8 by adding NaOH. After that, the suspension was stirred for another 12 h. The solid was separated by centrifugation and washed with ultrapure water, and dried in an oven at 60 °C and ground. Finally, clays modified with different Zr/clay ratios were obtained to study the effect of zirconium content on adsorption capacity. Zirconium modified clays are marked as Zr-MT, Zr-VT and Zr-KT. For example, montmorillonite with Zr/clay ratio 0.62 mmol/g was marked as Zr0.62-MT.

4.2.4 Material characterization

The morphology and of sample were imaged with scanning electron microscopy (SEM) using a Hitachi S-4800 SEM microscope. The interlayer structure of different absorbents was determined by X-ray diffractometry (XRD) using Bruker D8 Discover A25 diffractometer with copper Kα radiation. Zeta-potential measurements were analyzed by a Malvern Zetasizer Nano ZS.
4.2.5 Adsorption experiments

Equilibrium adsorption isotherm experiments were carried out in a set of 52 mL polypropylene tubes at room temperature, 40 mL of NaH$_2$PO$_4$ solutions with different initial concentrations and 0.04 g (1 g/L) of adsorbent added to each solution. The NaH$_2$PO$_4$ solutions were kept on an orbital shaker at 300 rpm (Thermo Scientific) to ensure good contact between absorbents and phosphate in water. The pH of phosphate solution was adjusted with HCl and NaOH. For equilibrium adsorption isotherm study, the initial phosphate concentration ranged from 1 to 20 mg P/L, the solution pH was about 7 and the contact time was 24 h. For kinetic study of phosphate adsorption on zirconium modified clays, 100 mL initial concentration of phosphate of 5 mg P/L was contained in 150 mL glass beaker with 0.1 g absorbent, the solution pH was adjusted to 7 and magnetic stirred at 300 rpm. The contact time was from 0 h to 24 h. To invest the effect of pH on the phosphate adsorption capacities, 0.04 g of the adsorbent was added into 40 mL of 5 mg P/L phosphate solution of different initial pH values, ranging from 3.0 to 11.0. The initial pH of phosphate solution was adjusted with 1M NaOH or HCl solution. To study the effect of coexisting anions on the phosphate adsorption capacities, 0.04 g of the adsorbent was added into 40 mL of 5 mg P/L phosphate solution containing 5 mM and 10 mM coexisting ions, which were prepared by dissolving sodium salt forms of Cl$^-$, NO$_3^-$, SO$_4^{2-}$ and CO$_3^{2-}$ into 5 mg P/L phosphate solution. To study the effect of ionic strength on the phosphate adsorption capacities, 0.04 g of the adsorbent was added into 40 mL phosphate solution with same salinity of freshwater, brackish water and seawater, by adding certain amount of NaCl into 5 mg P/L phosphate solution.

After the adsorption experiment, the absorbent was removed by a syringe filter (Millipore PES 0.22 µm membrane). Phosphate concentration in the supernatant solution was analyzed on SEAL AutoAnalyzer 3HR, Continuous segmented Flow Analyzer (SEAL Analytical).
The phosphorus adsorption capacity for zirconium modified clays at equilibrium ($q_e$, mg P/g) was calculated using the Equation (4.1):

$$q_e = \frac{(c_0 - c_e)V}{m} \quad (4.1)$$

where $q_e$ is amount of phosphorus adsorbed on adsorbent at equilibrium (mg P/g),

$c_0$ is initial concentration of phosphorus solution (mg P/L),

$c_e$ is concentration of phosphorus at equilibrium (mg P/L),

$m$ is mass of adsorbent used (g),

$V$ is volume of phosphorus solution (L).

4.3 Results and discussion

4.3.1 Characterization of absorbents

The SEM images of MT, VT, KT and zirconium modified clays with Zr/clay ratio 0.62 mmol/g are shown in Figure 4-3. As shown in the SEM images, all original and zirconium modified clays had coarse and irregular surfaces. The clays remained in the primitive crystal form and crystalline size of original clays. The sample MT exhibits a typical lamellar morphology with a large amount of foliated flakes, well stacked layer structure of MT can be easily observed in some regions of SEM image (Figure 4-3(a)). After zirconium modification, the layer structure of MT appeared more obvious and smoother this may be attributed to the partial delamination of MT layers, after the Zr$^{4+}$ exchange with interlayered cations. The morphology of raw VT and KT are similar to that of MT (Figure 4-3(b-c)), however, the surface of Zr0.62-VT and Zr0.62-KT appears rougher and covered by some large particles on the surface of both VT and KT, which are assumed to be deposition of ZrO$_2$ on clay surface.
Figure 4-3 SEM images of (a) MT, (b) Zr0.62-MT, (c) VT, (d) Zr0.62-VT, (e) KT, (f) Zr0.62-KT
Figure 4-4 XRD patterns of zirconium modified (a) MT, (c) VT, (e) KT between 2 theta 3-70° and (b) MT, (d) VT, (f) KT between 2 theta 3-20°
The layer structures of zirconium modified MT, VT and KT were analyzed by XRD and the patterns of between 2 theta 3-70º and 3-20º are shown in Figure 4-4. For original MT, a diffraction peak at 5.8 º was observed, which is attributable to the (001) reflection. The basal spacing (d_{001}) is 1.52 nm according to the Bragg equation (2d sinθ=nλ), which is normal for Ca-montmorillonite [143, 144]. The interlayer spacing was obtained by subtracting 0.96 nm, the thickness of the clay unit layer, from the XRD (001) basal spacing of clays [145, 146]. After loading with zirconium with Zr/clay ratio 0.31 mmol/g, the basal spacing of Zr0.31-MT slightly decreased to 1.50 nm, which may be caused by replacing Ca^{2+} ions with Zr^{4+} ions between clay layers. Since Zr^{4+} ion radius (0.086 nm) is smaller than Ca^{2+} ion radius (0.114 nm), the interlayer spacing decreased [147]. When Zr/clay ratio increased to 0.62, the basal spacing of Zr0.61-MT further decreased to 1.30 nm, which may be attributed to more Ca^{2+} ions replaced by Zr^{4+} ions in interlays. For Zr1.24-MT, the main diffraction peaks became very broad and shifted to 4.33º. The corresponding basal spacing is 2.04 nm, the increase in basal spacing proves that zirconium species entered into the interlayer region of MT after the modification [148]. The intensity of (001) refection indicates the extent of the long-range ordering of the stacking of the silicate layer [146]. With more loading of zirconium, the main peak was barely observed for Zr2.48-MT, indicating the disorder of clay layer structure after certain amount of zirconium intercalation. Other than the change of (001) reflection, no new peaks were visible after zirconium pillarization, suggesting no surface precipitation, at least crystalline forms were form [149].

Similar to MT, XRD pattern of original VT showed (001) reflection peak at 8.9º, which basal spacing was calculated as 0.99 nm. After zirconium modification, two new peaks showed up at 7.35º and 7.93º for all Zr/clay ratios, basal spacing are calculated as 1.2 nm and 1.11 nm, respectively. These two peaks are generated by zirconium species pillarized into the interlayer
region of VT after the modification. Moreover, the intensity of the peaks at 7.35° and 7.93° increase with increase of Zr/clay ratio from 0.31 to 2.48 mmol/g.

Different to MT and VT, there is no change in peak locations for KT before and after zirconium modification, and the basal spacing is 0.72 nm (2θ=12.25°), which is typical of kaolinite. Combining the non-expansible property of kaolinite as a 1:1 clay, it can be concluded that the modification of zirconium were all on the external surfaces of KT rather than intercalation between layers [137].

![Figure 4-5 Zeta potentials of unmodified and zirconium modified clays](image)

The zeta potentials of unmodified clays and zirconium modification clays with Zr/clay ratio 0.62 at various pH values are shown in Figure 4-5. Zeta-potential values are shown in Table 4-1. As can be seen from Figure 4-5(a), in the pH range from 3-11, MT, VT and KT are negatively charged, and the zeta potential values become more negative with increase of pH. After modified with zirconium, zeta-potential of modified clay increased within whole pH range from 3-11 due to the introduction of zirconium species, for all MT, VT and KT. Zeta-potential values of Zr0.62-MT remained below zero from pH 3-11. However, expect at pH 3, Zr0.62-VT and Zr0.62-KT showed zeta-potential above zero, these two modified clays remained below zero from pH 5-11. Thus,
under practical water and wastewater treatment conditions, the modified clays are mainly negatively charged.

Solution pH investigated in this study is 7, the dominate species of phosphate in the solution are $\text{H}_2\text{PO}_4^-$ and $\text{HPO}_4^{2-}$. Zirconium modified clays have negative surface charge, thus the coulomb repulsive interaction between negatively charged absorbents surface and the negatively charged phosphate could decrease the phosphate adsorption on zirconium modified clay.

Table 4-2 Zeta potentials of unmodified and zirconium modified clays

<table>
<thead>
<tr>
<th>pH</th>
<th>MT</th>
<th>Zr0.62-MT</th>
<th>VT</th>
<th>Zr0.62-VT</th>
<th>KT</th>
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<td>-41.53</td>
<td>-51.47</td>
<td>-51.83</td>
</tr>
</tbody>
</table>

4.3.2 Equilibrium adsorption isotherm studies of on phosphorus adsorption by absorbents

The adsorption capacity of zirconium modified clays on phosphorus was investigated by the equilibrium adsorption isotherm study, and the results are shown in Figure 4-6(a-c). Clays without zirconium modification showed very low phosphorus adsorption capacity with an initial phosphorus concentration of 20 mg P/L, for all three clays MT (0 mg P/g), VT (0 mg P/g) and KT (0.24 mg P/g). It clearly shows that all of the zirconium modified clays possess higher phosphorus removal capacities, as compared to untreated clays. Furthermore, adsorption isotherms of three clays followed the same trend with increase Zr content in resulting adsorbents. Take MT for example, the phosphorus uptake capacities increase from Zr0.31-MT to Zr2.48-MT, which is attributed to the greater Zr loading in the resulting adsorbents by increasing Zr/MT ratios in the initial synthetic solution.
To further evaluate the phosphorus adsorption capacity, the adsorption data was fitted with both Langmuir and Freundlich as given in Equation (4.2) and Equation (4.3) [150]:

$$q_e = \frac{q_{\text{max}}K_Lc_e}{1+K_Lc_e} \quad (4.2)$$

$$q_e = K_Fc_e^{1/n} \quad (4.3)$$

where $q_e$ is the amount (mg P/g) of phosphorus adsorbed at equilibrium,

$c_e$ is the equilibrium phosphorus concentration (mg P/L) in the solution,

$K_L$ is the Langmuir constant (L/mg), related to energy or net enthalpy of adsorption,

$q_{\text{max}}$ is the Langmuir monolayer adsorption capacity (mg P/g),

$K_F$ is the Freundlich constant, related to adsorption capacity,

$n$ is the Freundlich constant, measuring the adsorption intensity.

The parameters obtained in fitting the experimental data are summarized in Table 4-2. Three values agree with the two possible models of adsorption and they could be used to explain the significant capacity of the modified clays to quantify the adsorption. Overall, Langmuir model fitted the adsorption data better than Freundlich model, with most $R^2$ higher than 0.99. The Langmuir adsorption capacity ($q_{\text{max}}$) for the same type of clay increased with higher Zr/clay ratio. For Zr2.48-MT, VT, KT was determined at about 15.50, 9.95 and 9.60 mg P/g. Zr2.48 -MT showed best phosphate adsorption performance, it can also be observed that the phosphate amount that Zr2.48-MT adsorbed at low equilibrium concentration was quite high. For example, when the equilibrium concentration was just 0.01 mg P/L, the phosphate amount that Zr2.48-MT adsorbed was about 9.77 mg P/g. Such a performance is beneficial to its potential applications in case the phosphate concentration is not very high.
Figure 4-6 The adsorption equilibrium isotherm of phosphorus on zirconium modified clays at pH 7 and 25 °C. (a) MT, (b) VT, (c) KT (adsorbent dose 1g/L, pH 7, 25 °C)
Table 4-3 Langmuir and Freundlich isotherm constants for phosphorus adsorption on zirconium modified clays at pH 7 and T about 25 °C

<table>
<thead>
<tr>
<th>Adsorbents</th>
<th>Langmuir coefficients</th>
<th>Freundlich coefficients</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>R²</td>
<td>K_L (L/mg)</td>
</tr>
<tr>
<td>MT</td>
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<td>-</td>
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<tr>
<td>Zr0.31-MT</td>
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<td>Zr1.24-MT</td>
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<td>5.78</td>
</tr>
<tr>
<td>Zr2.48-MT</td>
<td>0.999</td>
<td>26.88</td>
</tr>
<tr>
<td>VT</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Zr0.31-VT</td>
<td>0.9765</td>
<td>0.48</td>
</tr>
<tr>
<td>Zr0.62-VT</td>
<td>0.9816</td>
<td>0.90</td>
</tr>
<tr>
<td>Zr1.24-VT</td>
<td>0.9962</td>
<td>2.23</td>
</tr>
<tr>
<td>Zr2.48-VT</td>
<td>0.9984</td>
<td>3.31</td>
</tr>
<tr>
<td>KT</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Zr0.31-KT</td>
<td>0.9743</td>
<td>2.98</td>
</tr>
<tr>
<td>Zr0.62-KT</td>
<td>0.9975</td>
<td>4.63</td>
</tr>
<tr>
<td>Zr1.24-KT</td>
<td>0.9985</td>
<td>4.24</td>
</tr>
<tr>
<td>Zr2.48-KT</td>
<td>0.9997</td>
<td>9.65</td>
</tr>
</tbody>
</table>

4.3.3 Kinetic studies on phosphorus adsorption by absorbents

The kinetics of the phosphorus adsorption from aqueous solution onto zirconium modified clays are shown in Figure 5. It is apparent that the adsorption process could be divided into two stages, a rapid stage at the very beginning followed by a gradually slower stage until the adsorption equilibrium was achieved with the increase of the adsorption time. For example, the adsorption rate of phosphorus on Zr1.24-MT at very beginning was fast. Within just 1 h, about 92.3% phosphorus was adsorbed onto zirconium modified MT, and the remaining phosphorus concentration dropped to about 0.37 mg/L, which satisfied the discharge requirement of 0.5-1.0 mg/L. With Zr/clay ratio 0.62, phosphorus adsorption capacity was in order MT>KT>VT. With Zr ratio increased to 1.24, zirconium modified MT and KT showed similar adsorption kinetics, due to increase of q_max. From the kinetic study, it could be seen that 8 h was adequate for the adsorption
equilibrium to occur on Zr/clay ratio 1.24. For Zr/clay 0.68, it took longer to reach equilibrium, however, after 8 h contact time the phosphorus uptake became very close to those at equilibrium.

The adsorption kinetic data could be fitted into various rate models to provide the understanding of the adsorption mechanism. The linear pseudo-first-order rate equation could be given as Equation (4.4):

$$\log(q_e - q_t) = \log q_e - \frac{k_1 t}{2.303} \quad (4.4)$$

And the pseudo-second-order rate equation could be expressed as Equation (4.5):

$$\frac{t}{q_t} = \frac{1}{k_2 q_e^2} + \frac{t}{q_e} \quad (4.5)$$

where \(t\) is the contact time (h), \(q_e\) and \(q_t\) are the amount of phosphorus adsorbed at equilibrium and at time \(t\), respectively. And \(k_1\) (h\(^{-1}\)) and \(k_2\) (g mg\(^{-1}\) h\(^{-1}\)) are the rate constants for pseudo-first-order and pseudo-second-order adsorption processes, respectively. The fitting curves by the pseudo-second-order rate model were demonstrated in Figure 4-7(b) and Figure 4-7(d), for Zr/clay ratio 0.62 and 1.24, respectively. The kinetics parameters obtained in fitting the experimental data are summarized in Table 4-3. It was found that the kinetics of phosphorus adsorption onto zirconium modified clays could be better fitted into the pseudo-second-order model than pseudo-first-order.

The pseudo-second-order rate model indicates a chemisorption occurred between phosphate and zirconium modified clays, involving valency forces through sharing or exchange of electrons between sorbent and sorbate, such as the replacement of \(-\text{OH}\) by phosphate \([16, 148]\). The initial high rate of phosphorus adsorption as shown in Figure 4-7, may be attributed to the existence of the bare surface. As the coverage increases, the number of available surface sites for adsorption decreases and the rate decreases until it reaches equilibrium \([135]\).
Figure 4-7 The phosphorus uptake of zirconium modified clays with the change of contact time, (a) Zr/clay ratio 0.62 mmol/g, (c) Zr/clay ratio 1.24 mmol/g. The fitting curves by the pseudo-second-order rate model of the kinetic study (b) Zr/clay ratio 0.62 mmol/g, (d) Zr/clay ratio 1.24 mmol/g (adsorbent dose at 1g/L, pH 7, 25 °C)

Table 4-4 The kinetic model parameters for the adsorption of phosphorus on zirconium modified clays at pH 7 and T about 25 °C

<table>
<thead>
<tr>
<th>Adsorbents</th>
<th>Pseudo-first-order model</th>
<th>Pseudo-second-order model</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$q_t = q_e (1 - e^{-k_1 t})$</td>
<td>$q_t = k_2 q_e^2 t (1 + k_2 q_e t)$</td>
</tr>
<tr>
<td></td>
<td>$R^2$</td>
<td>$k_1$ (h$^{-1}$)</td>
</tr>
<tr>
<td>Zr0.62-MT</td>
<td>0.987</td>
<td>2.176</td>
</tr>
<tr>
<td>Zr1.24-MT</td>
<td>0.9974</td>
<td>3.441</td>
</tr>
<tr>
<td>Zr0.62-VT</td>
<td>0.932</td>
<td>1.385</td>
</tr>
<tr>
<td>Zr1.24-VT</td>
<td>0.997</td>
<td>1.657</td>
</tr>
<tr>
<td>Zr0.62-KT</td>
<td>0.982</td>
<td>2.679</td>
</tr>
<tr>
<td>Zr1.24-KT</td>
<td>0.992</td>
<td>3.107</td>
</tr>
</tbody>
</table>
4.3.4 Influence of solution chemistry on phosphorus adsorption

Figure 4-8 Effect of pH on phosphorus adsorption by Zr0.62-MT (initial phosphorus concentration at 5 mg P/g, adsorbent dose at 1 g/L, 25 °C, contact time 24 h)

The solution pH effect on phosphorus adsorption by Zr0.62-MT was demonstrated in Figure 4-8. The phosphate adsorption on Zr0.62-MT was dependent on solution pH. Over the examined pH range from 3.0 to 11.0, increasing pH led to monotonically decreased phosphate adsorption. It is known that phosphate acid can dissociate to form different ionic species of $\text{H}_2\text{PO}_4^-$, $\text{HPO}_4^{2-}$ and $\text{PO}_4^{3-}$, depending on the pH of solution [61, 150, 151]. In the test pH range (3.0-11.0), $\text{H}_2\text{PO}_4^-$ and $\text{HPO}_4^{2-}$ are the dominant anionic species.

\[
\begin{align*}
\text{H}_3\text{PO}_4 & \rightleftharpoons \text{H}_2\text{PO}_4^- + \text{H}^+ & \text{pK}_{a1} = 2.15 \\
\text{H}_2\text{PO}_4^- & \rightleftharpoons \text{HPO}_4^{2-} + \text{H}^+ & \text{pK}_{a2} = 7.20 \\
\text{HPO}_4^{2-} & \rightleftharpoons \text{PO}_4^{3-} + \text{H}^+ & \text{pK}_{a3} = 12.33
\end{align*}
\]
In the pH range from 3.0 to 7.0, the phosphate adsorption capacity of Zr0.62-MT slightly decreased from 5.0 mg P/g to 4.39 mg P/g, indicating the suitability of Zr0.62-MT as an efficient phosphate adsorbent in a wide pH range. In the acidic pH region, the protonation of adsorbent surface is favored, which could enhance the electrostatic attraction between the adsorbent surface and the phosphate anions to facilitate the phosphate adsorption. The phosphate adsorption on Zr0.62-MT is a chemisorption dominated process, and the electrostatic attraction is not a major contributor in the adsorption process. When the pH value of solution increased from 7.0 to 9.0, the phosphate adsorption capacity decreased sharply from 4.39 mg P/g to 0.61 mg P/g. At pH 11.0, the phosphate adsorption capacity decreased slightly to 0.57 mg P/g, which is 89% lower than that at pH 3.0. In the pH range from neutral to basic region, the surface charge becomes more negative with pH increases, the coulomb repulsive interaction between negatively charged Zr0.62-MT surface and negatively charged phosphate species could decrease the phosphate adsorption. Moreover, at alkaline pH, OH⁻ ions might compete phosphate ions for the adsorption sites on Zr0.62-MT surface [59, 148]. So, less adsorption occurred at higher pH region.

Coexisting anions, such as Cl⁻, NO₃⁻, SO₄²⁻ and CO₃²⁻, generally present and may interfere with phosphate adsorption in natural water and wastewater. Figure 4-9 shows the effects of coexisting, including Cl⁻, NO₃⁻, SO₄²⁻ and CO₃²⁻, on the phosphate adsorption capacity of Zr0.62-MT anions at two concentrations (5 mM and 10 mM). Among the four anions, CO₃²⁻ has strongest effect on the phosphate adsorption, in the presence of 5 mM and 10 mM CO₃²⁻, the phosphate adsorption capacity of Zr0.62-MT reduced by 7% and 44%, respectively. Conversely, the presence of Cl⁻, NO₃⁻, SO₄²⁻, slightly facilitates phosphate adsorption, even at very high concentrations compared with that of phosphate (30 times as 5 mM and 60 times as 10 mM). Thus, Zr0.62-MT
could remove phosphate effectively when competing anions existed, which is beneficial to potential industrial applications.

![Figure 4-9 Effect of coexisting anions on the phosphorus adsorption by Zr0.62-MT with phosphorus concentration at 5 mg P/L, adsorbent dose at 1 g/L, pH 7, contact time 24 h](image)

The effect of ionic strength was also studied by adding NaCl into 5 mg P/L solution to the NaCl concentration at 8.5, 85 and 600 mM. These values simulate salinities in freshwater, brackish water and seawater, respectively. As demonstrated in Figure 4-10, the phosphate adsorption increased with the increase of the solution ionic strength. Similar observations have been reported in literature for the phosphate adsorption on ZrO₂ based absorbent [59, 61]. If phosphate form outer-sphere surface complexes through electrostatic forces, the adsorption of phosphate would decrease with the increase of ionic strength. In this study, the adsorption of phosphate would either not change or increase with the increase of ionic strength, suggesting that phosphate is removed from forming inner-sphere complexes at the absorbent surface [16, 59].
Figure 4-10 Effect of ionic strength on phosphorus adsorption by Zr0.62-MT, with phosphorus concentration at 5 mg P/L, adsorbent dose at 1 g/L, pH 7, contact time 24 h

Figure 4-11 Effect of humic acid concentration on phosphorus adsorption by Zr0.62-MT, with phosphorus concentration at 5 mg P/L, adsorbent dose at 1 g/L, pH 7, contact time 24 h
Phoslock (CSIRO, Australia), a commercial lanthanum ion-exchanged bentonite clay, is effective for phosphate removal from water and among the main commercial restoration products in Europe. As Dithmer et al. reported, the presence of dissolved organic carbon (DOC) interferences with the binding capacity of Phoslock. Since DOC can possibly form complexes with La$^{3+}$, DOC has a negatively influence on phosphate removal by Phoslock [152]. In water containing DOC, Phoslock is capable to bind less phosphate than theoretical value. To study the influence of DOC on phosphate removal on zirconium modified clays, humic acid (HA) was added into 5 mg P/L solution to concentration from 1-10 mg/L. As demonstrated in Figure 4-11, with increase of HA concentration, the phosphate adsorption capacity of Zr0.62-MT almost retained the same. Thus, this model adsorption study indicates that DOC has barely influence on zirconium modified clay adsorption capacity. It can be concluded that unlike lanthanum-based adsorbent, zirconium modified clays are favorable for phosphate removal in water containing DOC.

4.4 Conclusion

In summary, zirconium modified clays have been prepared and studied for phosphate adsorptive removal from aqueous solution. Comparing three clays, zirconium modified MT (2:1), VT (2:1) and KT (1:1) exhibit different structure and surface properties, and thus performance differently toward phosphate adsorption. MT with 2:1 layer structure and highly expansibility served best as substrate of amorphous ZrO$_2$. The adsorption kinetics data of phosphate on zirconium modified clays could be well described by the pseudo-second-order model, indicating that the adsorption was through chemisorption. The experimental equilibrium data of phosphate adsorption on modified clays were fitted better by Langmuir isotherm model than the Freundlich, implying monolayer adsorption. Based on the Langmuir isotherm model, the maximum phosphate adsorption capacity for Zr2.48-MT was determined as 15.5 mg P/g. The different behaviors of
three clays were observed, yet the understanding of the mechanism is still not enough. Future work can be done on interaction between zirconium ions and different clays. The phosphate adsorption capacity is pH dependent, which is favorable in acidic and neutral pH range. Co-existing anions like Cl\(^-\), NO\(_3^-\), SO\(_4^{2-}\) have no significant influence on phosphate removal, but the presence of CO\(_3^{2-}\) decreases the adsorption capacity at high concentration. Unlike lanthanum-based phosphate restoration products, DOC has no significant effect on phosphate adsorption capacity on zirconium modified clays, the obtained clays can be studied for practical water treatment.
CHAPTER 5  Summary of research work and proposed future work

5.1 Summary of research work

Nanostructured metal oxide materials have offered new opportunities to design novel materials in applications of water treatment and LIBs. The objective of this study is to demonstrate high performance TiO$_2$ based materials as photocatalyst and anode material, and ZrO$_2$ based absorbents.

1. Pt/rGO-TiO$_2$ photocatalyst for photodegradation of MO

By incorporating Pt nanoparticles and rGO with 3DOM TiO$_2$, a new effective photocatalyst was facilely synthesized. The composite material was thoroughly characterized by various techniques, including SEM, TEM, XRD, Raman, DRS and the photocatalytic activity was tested under visible light. The excellent charge-separation property and high adsorption capacity of rGO increased the charge carrier lifetime and affinity to organic molecules. The introduction of Pt nanoparticles increased spectral response to visible light through surface plasmon resonance and suppressed charge recombination. The synergism between rGO and Pt nanoparticles results in the novel ternary photocatalyst effective to remove MO under visible light, the photodegradation rate toward MO of resulting material is 6 times higher than 3DOM TiO$_2$. Moreover, the photocatalysis process is stable and reusable, making it a promising photocatalyst for practical water treatment.

2. Carbon coated TiO$_2$@Fe$_2$O$_3$ nanocomposite for LIBs

TiO$_2$@Fe$_2$O$_3$ was synthesized through hydrolysis of FeCl$_3$ in solution, and the composite was carbon coated using glucose as carbon source. The carbon coated TiO$_2$@Fe$_2$O$_3$ material was applied as electrode mater in LIBs and showed good electrochemical performance. The composite exhibits a large reversible capacity about 570 mAh g$^{-1}$, which is about four times of the reversible capacity of 3DOM TiO$_2$. In addition to the high reversible capacity, the obtained material also
exhibits good cycle performance and superior rate capacity. This rationally designed composite benefits from both good stability of TiO₂, high capacity of Fe₂O₃, and good electron conductivity of carbon.

3. Adsorption removal of phosphors by zirconium modified clays

In summary, zirconium modified clays have been prepared and studied for phosphate adsorptive removal from aqueous solution. Comparing three clays, zirconium modified MT (2:1), VT (2:1) and KT (1:1) exhibit different structure and surface properties, and thus performance differently toward phosphate adsorption. The adsorption kinetics data of phosphate on zirconium modified clays could be well described by the pseudo-second-order model, indicating that the adsorption was through chemisorption. The experimental equilibrium data of phosphate adsorption on modified clays were fitted better by Langmuir isotherm model than the Freundlich, implying monolayer adsorption. Based on the Langmuir isotherm model, the maximum phosphate adsorption capacity for Zr2.48-MT was determined as 15.5 mg P/g.

5.2 Future work

In this thesis, three different novel nanomaterial have been demonstrated in applications of water treatment and LIBs through rational design of metal oxides. However, the synthesis process/methods are still far from satisfaction to meet the practical applications. Therefore, there is a plenty of room for discovering the new synthesis methods and optimizing existing procedures towards scale-up phase. Some recommendations for future works are summarized as following:

- 3DOM TiO₂ possess unique optimal and electrochemical properties, however, the synthesis process needs involving preparation and removal of sacrificial template, which increases cost and time towards preparation process. More works can be done to
balance cost and effort in production process and performance of the TiO$_2$ based materials.

- Three clays have been studied as support of ZrO$_2$ for adsorbents to remove phosphate in aqueous solution. The different behaviors of three clays were observed, yet the understanding of the mechanism is still not enough. Future work can be done on interaction between zirconium ions and different clays. And the obtained clays can be studied for practical water treatment.
References

1. OECD Environmental Outlook to 2050. 2012.

2. Climate change 2014: Impacts, Adaptation, and Vulnerability Intergovernmental panel on climate change. 2014, IPCC.


126. Xia, H., et al., Hierarchical TiO2-B nanowire@α-Fe2O3 nanothorn core-branch arrays as superior electrodes for lithium-ion microbatteries. Nano Research, 2014. 7(12): p. 1797-1808.


Appendix: Supporting information for Chapter 2

Bragg reflection peak estimation

\[
\lambda_{\text{max}} = \frac{2D\sqrt{2}}{\sqrt{3}} \sqrt{n_{\text{eff}}^2 - \sin^2 \theta}
\]

\(\lambda_{\text{max}}\) (nm) is stopband position for 1\(^{st}\) order Bragg diffraction

\(n_{\text{eff}}\) is the average refractive index of photonic crystal.

D is pore size of 3DOM TiO\(_2\)

Assume incident angle of incident light with respect to surface normal, \(\theta=0^\circ\).

\[
n_{\text{eff}} = \sqrt{f n_{\text{TiO}_2}^2 + (1 - f) n_{\text{air}}^2}
\]

f is TiO\(_2\) phase volume percentage, generally taken as 0.26

\(n_{\text{TiO}_2}\) is refractive index of TiO\(_2\), for anatase taken as 2.5

\(n_{\text{air}}\) is refractive index of air, taken as 1.

Figure S2-1 SEM image of PMMA template

PMMA (330 nm) spheres are uniform in size and effectively packed into face-centered-cubic (FCC) structure. Each colloidal particle is connected with 6 surrounding particles through 6 bridges on the two-dimensional plane.
Figure S2-2 Elemental mapping of the Pt-TiO$_2$ composite. (a) Ti, Pt and O element mapping images, (b)-(d) Ti, Pt and O dispersion, respectively, (e) SEM image of Pt-TiO$_2$, (f) EDS element spectrum of Pt-TiO$_2$.

First order kinetic model

Table S2-1 Parameters of first order kinetic model for degradation of MO

<table>
<thead>
<tr>
<th>Catalysts</th>
<th>Rate constant, k $\times 10^3$, min$^{-1}$</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>TiO$_2$</td>
<td>y=0.0021x+0.0037</td>
<td>2.1</td>
</tr>
<tr>
<td>Pt-TiO$_2$</td>
<td>y=0.0032x+0.0291</td>
<td>3.2</td>
</tr>
<tr>
<td>rGO-TiO$_2$</td>
<td>y=0.0062x+0.0706</td>
<td>6.2</td>
</tr>
<tr>
<td>Pt/rGO-TiO$_2$</td>
<td>y=0.0129x+0.0762</td>
<td>12.9</td>
</tr>
</tbody>
</table>
Curriculum Vitae

Jingwan Huo

EDUCATION

• University of Wisconsin-Milwaukee, Milwaukee, Wisconsin, USA
  Ph.D. Major in Mechanical Engineering, Minor in Civil Engineering 2018.5
  Selected Courses: Intermediate Fluid Mechanics; Transport in Porous Media; Thermal Radiation and Conduction; Electron Microscopy Laboratory (SEM, TEM).

• University of Wisconsin-Milwaukee, Milwaukee, Wisconsin, USA
  M.S. Major in Mechanical Engineering 2012.12
  Selected Courses: Introduction to Water and Sewage Treatment; Freshwater Engineering; Water Quality Assessment; Transition Metal/organometallic.

• Huazhong University of Science and Technology, Wuhan, Hubei, China
  B.S. Major in Chemical Engineering 2009.9

TECHNICAL SKILLS:

• Material background: Polymer, metal oxide, liquid, ceramic films, graphene, composite
• Synthesis processes: Atomic Layer Deposition (ALD); Solution processing (polymerization, hydrothermal, sol-gel, anodization)
• Characterization: Gas chromatography (GC, Agilent 7890A); X-ray diffractometer (XRD); Scanning electron microscopy (SEM)/Energy-dispersive X-ray Spectroscopy (EDS); Transmission electron microscopy (TEM); Raman spectrum; Inductively coupled plasma (ICP); UV-Vis Spectrometer; Scanning Mobility Particle Sizer Spectrometer (SMPS); High-performance Liquid Chromatography-Mass Spectrometer (HPLC-MS).
• Computer based skills: PTC Cero; LabView; Microsoft Office software.

RESEARCH EXPERIENCES:

Research Assistant  University of Wisconsin-Milwaukee, Milwaukee, USA 2011.1 - Present
  • Real time monitoring nanoparticle emissions from Al2O3 ALD process from Savannah S100 ALD system using SMPS. Optimization of operation conditions such as pulsing & purging time and temperature to minimize nanoparticles emissions and improve utilization of reaction precursors.
  • Designed and synthesized novel TiO2-based hybrid nanomaterials as photo-catalyst for dye photodegradation and anti-fouling membrane under visible-light illumination.
  • Research on economic friendly inexpensive phosphorous adsorbent using clay and metal oxide- modified clay.
  • Explored hybrid nanostructures alternative materials as anode for Lithium-ion batteries. By growing Fe2O3 on porous TiO2, attained good cyclability and rate capability for Li-ion batteries.

Teaching Assistant  University of Wisconsin-Milwaukee, Milwaukee, USA 2011.9 – 2017.5
  • Experienced instructor in fundamental engineering labs, including Engineering Fundamentals I and II lab; How things works: Understanding Technology lab; and
Mechanical Engineering Experimentation lab. Students and instructors of classes were satisfied about the course and lab instruction.

- Hands-on helping student understanding basic engineering disciplines, data collection and analysis, sensor calibration, and software including Microsoft office, Matlab, PTC Cero, and LabView. Guide student through experiment design & planning and complete industrial projects from companies such as Master Lock.

**Chemical Analyst**  Oil Crops Research Institute, Chinese Academy of Agricultural Science, Wuhan, China 2010.6 – 2010.12

- In charge of determination the composition of various fatty acids in rapeseeds, through esterification and extraction fatty acids and quantifying with Gas Chromatography (GC);
- Explored the method to determine molecular structure of fatty acids in rapeseeds using HPLC-MS;
- Be familiar with operations of GC and HPLC-MS.

**Research Assistant**  Huazhong University of Science and Technology, Wuhan, China 2009.2 – 2010.6

- Modified multi-walls carbon nanotubes (MWCNs) by inorganic-coating (MgO and SiO₂) through sol-gel method. Improved heat conductivity of epoxy for ~200% by integrating inorganic-coated MWCNs into composites.
- Designed and research on SiO₂/polyacrylate paint nano-hybrid for abrasion-resistant paint.
- Characterization of inorganic-polymer hybrids using FTIR (especially Temperature-Dependant FTIR), DSC, TG, SEM, rheometer, abrasion test.

**SELECTED PUBLICATIONS:**