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Novel Nanostructured Materials for Solar Fuel Production and Advanced Rechargeable Batteries

Cunyu Zhao
University of Wisconsin-Milwaukee

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NOVEL NANOSTRUCTURED MATERIALS FOR SOLAR FUEL PRODUCTION AND ADVANCED RECHARGEABLE BATTERIES

by

Cunyu Zhao

A Dissertation Submitted in Partial Fulfillment of the Requirements for the Degree of Doctor of Philosophy in Engineering at The University of Wisconsin-Milwaukee

December 2014
ABSTRACT

NOVEL NANOSTRUCTURED MATERIALS FOR SOLAR FUEL PRODUCTION AND ADVANCED RECHARGEABLE BATTERIES

by

Cunyu Zhao

The University of Wisconsin-Milwaukee, 2014
Under the Supervision of Professor Ying Li

Non-renewable fossil fuels are the major sources to meet the energy, electricity and transportation demands of today's world. The over consumption of fossil fuels will lead to the increasing energy crisis and disastrous effects such as air pollution, global warming etc.

The primary greenhouse gas is CO$_2$ mainly emits from the combustion of fossil fuels. Photocatalytic reduction of CO$_2$ using sunlight as the energy input is a promising way to reduce CO$_2$ level in the atmosphere and in the meantime produce alternative fuels such as CO, methane, methanol, etc. Among the various photocatalyst materials reported, nanomaterial TiO$_2$ is the most widely studied due to its suitable band positions, high chemical stability, non-toxic nature, and low cost. However, the energy conversion efficiency using TiO$_2$ for CO$_2$ photoreduction is still low, mainly due to the reasons of (1) high probability of recombination of photo-induced electron-hole pairs, (2) fast backward reaction of hydrogen and oxygen to form water, and (3) limited ability of visible light utilization.

Another efficient way to decrease CO$_2$ emissions is to reduce fossil fuels consumption. The invention of hybrid electric vehicles (HEVs) and electric vehicles (EVs) are great promise of replacing traditional gasoline driven automobiles. As one of the
new generation high energy density batteries, lithium-sulfur (Li-S) battery is very attractive because sulfur has a high theoretical capacity of 1,675 mA h g\(^{-1}\). However, the practical realization of Li-S batteries is limited by several problems: (1) poor electrical conductivity of sulfur (2) dissolution of the lithium polysulfide intermediates into the electrolyte, and (3) large volume expansion of sulfur during cycling.

One objective of this study is to demonstrate high-efficiency photocatalysts using innovative hybrid nanostructures that consist of Ce doped TiO\(_2\) dispersed on mesoporous silica (SBA-15) or noble-metal nanoparticles Ag supported on TiO\(_2\) or MgAl-LDOs (layered double oxides) grafted on TiO\(_2\) (TiO\(_2\)-MgAl LDOs). The use of Ce doping could result in smaller TiO\(_2\) nanocrystals and facilitate electron-hole separation, while SBA-15 provides good dispersion of TiO\(_2\) and a strong interaction between TiO\(_2\) and the substrate. And Ag species on TiO\(_2\) facilitate electron trapping and transport to the catalyst surface, and thus, can potentially enhance multi-electron transfer processes. TiO\(_2\)-MgAl LDOs is favorable for CO\(_2\) species adsorption on the photocatalyst, therefore, compensating the weakened CO\(_2\) adsorption ability at higher temperature in the presence of H\(_2\)O vapor. The other objective of this study is to find alternative materials as anode for Li-ion batteries and demonstrate high-performance Li-S battery electrodes using hybrid nanomaterials consist of sulfur infiltrated porous microsphere carbon (PMC). Carbon/TiO\(_2\) was found to be promising as anode alternative to replace graphite materials to avoid safety issues for Li-ion batteries. Cathodes made of sulfur infiltrated in such a multi-modal porous carbon framework provide advantageous properties that guarantee the superior electrochemical performance.
Ce-doped TiO$_2$ on SBA-15, Ag deposited TiO$_2$ (Ag/TiO$_2$) and MgAl-LDOs grafted on TiO$_2$ (TiO$_2$-MgAl LDOs) were synthesized and characterized for applications in CO$_2$ photoreduction with H$_2$O. Ce-doped TiO$_2$ were synthesized using sol-gel method and SBA-15 was then added to the sol to prepare Ce-doped TiO$_2$ on SBA-15 nanocomposites. Modification of TiO$_2$ with Ce significantly stabilized the TiO$_2$ anatase phase and increased the specific surface area, which contributed to an improvement of CO production from CO$_2$ reduction. Dispersing Ce-TiO$_2$ nanoparticles on the mesoporous SBA-15 support further enhanced both CO and CH$_4$ production. The superior catalytic activity may be related to the partially embedded Ce-TiO$_2$ nanoparticles in the ordered 1-D pores in SBA-15 that form synergies between the different components of the catalysts and enhance the diffusion and adsorption of CO$_2$. Ag/TiO$_2$ nanocomposites were synthesized by spray pyrolysis technique. This work has demonstrated the feasibility of syngas (H$_2$ and CO) production from a gas mixture of CO$_2$, H$_2$O and CH$_3$OH through a photocatalytic reduction process on Ag/TiO$_2$ nanocomposite catalysts under solar irradiation. The material property analysis and photocatalytic activity results showed that the ultrasonic spray pyrolysis method is much superior to conventional wet impregnation process with the advantages of smaller Ag nanoparticles, a better Ag dispersion on TiO$_2$, and a higher fraction of metallic Ag species, which facilitate charge transfer and improve photocatalytic activity. TiO$_2$-MgAl LDOs were synthesized by hydrothermal and coprecipitation method. As the MgAl LDOs concentration increases, TiO$_2$ crystal size was increased. MgAl LDOs grafting on TiO$_2$ cuboids may help improve the adsorption ability of CO$_2$ onto TiO$_2$ which improves the photocatalytic activity of CO$_2$ reduction.
Our work also entails the synthesis and characterization of carbon coated TiO$_2$ for the application of Li-ion batteries and sulfur infiltrated porous microsphere carbon (PMC/S) for the application of Li-S batteries. Carbon decorated on commercial TiO$_2$ nanoparticles (P25 and P90) composites with optimized carbon concentration and structure were fabricated by a facile process employing carbonization method. The electrochemical performance of C-P90 was superior to C-P25 because of its higher specific surface area and larger anatase fraction that can accommodate more lithium ions. 1.9% carbon was found to form an optimized carbon layer on TiO$_2$ that can improve the electronic conductivity. The PMC was synthesized by spray pyrolysis method. Then PMC/S was fabricated via a liquid phase infiltration. The novel-structured porous carbon microspheres possess a controllable multi-modal pore size distribution, i.e., a combination of interconnected micropores, mesopores and macropores, which is beneficial for Li-S batteries electrochemical performance.

Future work includes further improvement of PMC/S composites to inhibit shuttle effect and improve the electrode performance including the cyclability and rate capability.
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<td>1D</td>
<td>one-dimensional</td>
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<tr>
<td>3D</td>
<td>three-dimensional</td>
</tr>
<tr>
<td>BET</td>
<td>Brunauer-Emmett-Teller method</td>
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<tr>
<td>BJH</td>
<td>Barrett-Joyner-Halenda method</td>
</tr>
<tr>
<td>CB</td>
<td>conduction band</td>
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<td>cyclic voltammogram</td>
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<tr>
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<td>electron-hole</td>
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<td>flame ionization detector</td>
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<td>greenhouse gas</td>
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<td>hybrid electric vehicles</td>
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<tr>
<td>HRTEM</td>
<td>high resolution transmission electron microscopy</td>
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<tr>
<td>IR</td>
<td>infrared</td>
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<td>LDHs</td>
<td>layered double hydroxides</td>
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<td>layered double oxides</td>
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<tr>
<td>LIBs</td>
<td>lithium-ion batteries</td>
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<td>Abbreviation</td>
<td>Full Form</td>
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<td>---------------------------</td>
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<tr>
<td>Li-S</td>
<td>lithium-sulfur</td>
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<tr>
<td>NPs</td>
<td>nanoparticles</td>
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<td>PMC</td>
<td>porous microsphere carbon</td>
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<tr>
<td>PSD</td>
<td>pore size distribution</td>
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<tr>
<td>Rct</td>
<td>interfacial resistance</td>
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<tr>
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<td>solid-electrolyte interface</td>
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CHAPTER 1 INTRODUCTION AND RESEARCH OBJECTIVES

1.1 Introduction

Energy supply is essential to economic development everywhere in the world, and current pattern of global energy development greatly relies on fossil fuels (coal, natural gas and oil). The future global economy is likely to consume ever more such non-renewable fossil fuels, especially with the rising energy demand of developing countries. As a result, this is causing crisis that our energy resources are starting to run out, with devastating consequences for the global economy development. The combustion of fossil fuels for energy and transportation (see Fig. 1.1, main activities that consume fossil fuels and emits CO$_2$) emits CO$_2$ directly to the atmosphere, which is the primary greenhouse gas (GHG). The energy crisis is further exacerbated by the tremendous risk of climate change associated with the use of fossil fuels.

![Overview of Greenhouse Gases](image1.png)

![Sources of Greenhouse Gas Emissions](image2.png)

**Figure 1.1** Greenhouse gases (GHGs) emissions and the primary sources of greenhouse gas emissions in the United States in 2011.¹
CO₂ emissions is likely to increase as the fossil fuels consumption increases. The huge amount of CO₂ existing in the atmosphere is expected to cause continued global warming which has substantial impacts on the environment, human health and economy. How to solve this environmental concern has become an emergency. Although strategies have tried to bury CO₂ in the deep ground or ocean, potential risk is not fully evaluated and catastrophe may occur if overload of CO₂ pumped into deep ground. Photocatalytic reduction of CO₂ using sunlight as the energy input is a promising way to reduce CO₂ level in the atmosphere and in the meantime produce alternative fuels such as CO, methane, methanol, etc. Semiconductor photocatalysts such as WO₃, ZrO₂, Ga₂O₃, and TiO₂ have been studied for such applications, and among them, TiO₂ has been considered the most appropriate photocatalyst due to its high photosensitivity, non-toxic nature, low cost, and easy availability. However, the photoefficiency of CO₂ reduction on TiO₂ is usually very low, mainly due to the fast recombination of photo-excited electron-hole (e-h) pairs and the wide band gap of TiO₂ (3.2 eV for anatase) that does not allow the utilization of visible light. Innovations are needed overcome these limitations to enhance the CO₂ conversion to alternative fuels.

Photocatalytic reduction of CO₂ to alternative fuels is a promising way to reduce CO₂ level in the long run, which does not mean we could keep using the non-renewable fossil fuels. It is essential to find alternative energy to replace the fossil fuels especially for transport energy, as transportation consists almost 1/3 of the total energy consumption (see Fig. 1.1). And it is expected that the energy use in the transportation sector will continue to grow at rates that are considerably larger than other sectors of the U.S. economy (see Fig. 1.2). Fig. 1.3 shows that petroleum is the main fossil fuels used for the energy
requirement in the transportation. Is it possible to replace the petroleum by other energy system? Scientists have proposed to apply lithium-ion batteries (LIBs) in the hybrid electric vehicles (HEVs) or electric vehicles (EVs) to lower or avoid the petroleum consumption. The HEVs or EVs demand for safe, light, low-cost, high-energy density, and long-lasting rechargeable batteries. Current lithium-ion batteries (LIBs) using transition metal compounds as cathode materials operate on the basis of topotactic intercalation reactions, which inherently limits its storage capacity to around 300 mAh g\(^{-1}\) for any perspective system. As a result, the current LIBs cannot provide a suitably long driving distance for EVs.\(^{15-18}\) As one of the new generation high energy density batteries, lithium-sulfur (Li-S) battery is very attractive because sulfur has a high theoretical capacity of 1,675 mA h g\(^{-1}\), nearly five times higher than that of existing transition metal oxides and phosphates materials.\(^{19-22}\)

![Delivered Energy Consumption](image)

**Figure 1.2** Energy consumption for different sectors of US economy.\(^{23}\)
1.2 Solar fuel production using CO$_2$ as the feedstock

Photocatalytic hydrogen production from water using semiconductor catalysts is a promising way to convert solar energy to clean and renewable hydrogen fuel energy. For efficient water splitting, the conduction band (CB) level of the semiconductor should be more negative than water reduction level (to produce H$_2$) while the valence band (VB) level should be more positive than water oxidation level (to produce O$_2$), as shown in Fig.
Among the various photocatalyst materials reported, TiO$_2$ is the most widely studied due to its suitable band positions, high chemical stability, non-toxic nature, low cost, and easy availability.

Figure 1.5 Mechanism of photocatalytic hydrogen production and CO$_2$ reduction with water on TiO$_2$ semiconductor and their reduction potentials at pH 7 vs. NHE.

Besides hydrogen production from photocatalytic water splitting, another promising solar fuel technology is to photocatalytically reduce carbon dioxide with water to generate gaseous or liquid fuels such as CO, methane, methanol, etc. Fig. 1.5 shows that these CO$_2$ reduction reactions are theoretically feasible on TiO$_2$. However, photocatalytic CO$_2$ reduction is believed to be more challenging than simply water splitting, because CO$_2$ is thermodynamically very stable, the CO$_2$/CO reduction potential is even more negative than that of H$_2$O/H$_2$, and the multi-electron processes (i.e., six electron reduction to produce methanol and eight-electron reduction to produce methane) have very low efficiency. Hence, many studies have also been conducted to modify the TiO$_2$ catalysts to enhance the
CO$_2$ photoreduction rate, including metal loading, nonmetal doping, and usage of a high surface area substrate, etc.

1.3 Lithium Secondary Batteries

Lithium ion batteries (LIBs) have been dominating the power sources of portable electronic devices for decades. The critical performance indicators for LIBs are capacity, cyclability and rate capability, which are strongly dependent on the active electrode materials. Graphitic carbons are most commonly used as conventional commercial anode materials owing to its low cost, relatively high energy density and very low flat working potential. However, graphitic carbon anode materials suffer the disadvantages of severe safety issues because of its low operating voltage (~0.1 V vs Li/Li$^+$), poor life cycle performance due to solid-electrolyte interface (SEI) film and the formation of dendritic lithium on the surface of graphitic carbons anode. Thus, it is essential to make improvements such as safety, cost, high efficiency and long cycle life of the anode materials for LIBs.

As one of the new generation high energy density batteries, lithium-sulfur (Li-S) battery is very attractive because sulfur has a high theoretical capacity of 1,675 mA h g$^{-1}$, nearly five times higher than that of existing transition metal oxides and phosphates materials.$^{19-22}$ Moreover, sulfur is light, cheap, and abundant. However, the practical realization of Li-S batteries is limited by several problems: (1) poor electrical conductivity of sulfur (only 5×10$^{-30}$ S/cm at 25 °C),$^{29-30}$ (2) dissolution of the lithium polysulfide (Li$_2$S$_x$, 4≤x≤8) intermediates into the electrolyte,$^{31-32}$ and (3) large volume expansion of sulfur (~80%) during cycling.$^{33-34}$ Consequently, Li-S batteries suffer from rapid capacity decay and low Coulombic efficiency.
1.4 Summary and Literature Review

Aimed at solving the problems of TiO$_2$-based photocatalysts for CO$_2$ photoreduction, i.e. low photoefficiency of CO$_2$ reduction on TiO$_2$, the fast recombination of photo-excited electron-hole (e-h) pairs and the wide band gap of TiO$_2$, modification of TiO$_2$ needs to be studied in this work. During the past decade, lots of work about CO$_2$ photoreduction has been done but the mechanism still maintains opaque. Ceria and silver will be used to modify TiO$_2$ to study the unclear mechanism of CO$_2$ photoreduction.

Rare earth element modified TiO$_2$ have not been studied for CO$_2$ photoreduction, although they have demonstrated enhanced photocatalytic oxidation ability than bare TiO$_2$. Cerium (Ce) is one of the four most abundant rare earth elements, and composites of Ce-TiO$_2$ have shown enhanced photocatalytic activity for water splitting and degradation of organic compounds. The Ce$^{3+}$/Ce$^{4+}$ redox couple may facilitate charge transfer. Besides modifying the properties and nanostructures of TiO$_2$ itself, many approaches have been used to immobilize TiO$_2$ nanoparticles on mesoporous substrates such as molecular sieve 5A, amorphous silica, and SBA-15, and the direct benefits include larger surface area and better dispersion of the nano-sized catalysts. It has been reported that Cu-TiO$_2$, Ru-TiO$_2$, and TiO$_2$ supported on amorphous silica showed higher CO$_2$ photoreduction rates than the catalysts without supports. The Si-OH group on SBA-15 was found to enhance the interaction of CO$_2$ and H$_2$O on the catalyst surface based on the results of an IR study. Experimental results also demonstrated an enhancement of CO$_2$ photoreduction to methanol on a 2%Cu-TiO$_2$/SBA-15 sample compared with 2%Cu-TiO$_2$, and the promotional effect was likely attributed to the synergistic effect between the metal species, TiO$_2$ and the support.
As for Ag/TiO\textsubscript{2} photocatalyst, photocatalytic H\textsubscript{2} production and CO\textsubscript{2} reduction using water as the reductant are mostly being investigated separately in the literature, very few publications report on the simultaneous H\textsubscript{2} production and CO\textsubscript{2} reduction in one experimental system. Many literature publications on CO\textsubscript{2} photoreduction either did not detect H\textsubscript{2} production or simply did not measure H\textsubscript{2} production; only a few studies reported concurrent production of H\textsubscript{2} and C1 or C2 fuels but the H\textsubscript{2} yield was at the similar level or even less than the hydrocarbon fuels.\textsuperscript{6, 48-49} In addition, most of the experiments were carried out in an aqueous system where many other reaction products such as methanol, formic acid, and formaldehyde were detected and the product selectivity was not controlled.\textsuperscript{50-51} Hence, it is desirable to be able to control the H\textsubscript{2}: CO ratio in the products and to eliminate the unwanted reaction intermediates or products so that a high purity syngas can be produced. In this regard, a gas-phase reaction system may be more appropriate than an aqueous system, because in our previous work of photocatalytic CO\textsubscript{2} reduction with water vapor,\textsuperscript{5, 52} CO was the main reaction product with formation of minor CH\textsubscript{4} and no other products were detected. H\textsubscript{2} production was not detected using bare TiO\textsubscript{2} or Cu/TiO\textsubscript{2} in our previous work,\textsuperscript{5, 52} but the addition of noble metals such as Pt and Ag to TiO\textsubscript{2} has been demonstrated for effective H\textsubscript{2} production in the literature.\textsuperscript{53-55} Therefore, the first objective of this work was to explore the feasibility of syngas production from CO\textsubscript{2} and water vapor through a photocatalytic process using a silver-modified TiO\textsubscript{2} (Ag/TiO\textsubscript{2}) nanoparticle catalyst. Ag was selected because it is much less expensive than Pt and Ag/TiO\textsubscript{2} has shown effectiveness in H\textsubscript{2} production\textsuperscript{53} and CO\textsubscript{2} photoreduction\textsuperscript{48} in separate studies.

Our research work on CO\textsubscript{2} photoreduction was not only focused on tackling the
major obstacles, but also on other factors that hinder the photocatalytic activity but are rarely studied in the open access literature, such as the weakened CO₂ adsorption on TiO₂ in the presence of H₂O vapor at the solid-gas interface and limited desorption of reaction products or intermediates from the catalyst surface. In order to improve the CO₂ photoreduction conversion efficiency to hydrocarbon fuels, it is highly desirable to enhance the CO₂ adsorption on the photocatalysts. We have reported works about MgO-TiO₂ hybrid adsorbent-photocatalyst prepared by different methods. MgO is chosen as CO₂ adsorbent because of its good CO₂ adsorption ability that is boosted in the presence of H₂O vapor. We found that MgO-TiO₂ hybrid catalysts possessed a much more stable catalytic performance than pristine TiO₂, which may be due to the easier desorption of reaction intermediates and the enhanced CO₂ adsorption by MgO.

However, the CO₂ adsorption capacity for MgO was not high enough comparing with hydrotalcite-derived mixed oxides based on our previous work and other groups’ reported literature. As sorption materials, activated MgAl-LDOs have recently been found to have a high sorption capacity for CO₂. Since the flue gas from a coal burner is with a high temperature around 400-600 °C and the real flue gas emitted to the atmosphere is with a temperature around 110-160 °C, it is essential to adsorb CO₂ at high temperatures instead of cooling it to room temperature for CO₂ photocatalytic reduction application to be more feasible. In particular, MgAl-LDOs can capture much more CO₂ than MgO at higher temperature range 150-450 °C. Therefore, we aim to design a novel hybrid composite incorporating an adsorbent with strong CO₂ adsorption ability and a TiO₂ photocatalyst. Most literature reported that nano-size TiO₂ were used for many purposes such as air purification, degradation of organic matters, heavy metal removal and photocatalytic
CO\(_2\) reduction to alternative fuels because nano-size TiO\(_2\) possess high surface area which guarantee its high photocatalytic activity. However, more and more papers recently pointed out that nano-sized TiO\(_2\) material could be possibly dangerous to human health since the ultra-small size particles could penetrate into human body from skin and nose especially when it is applied for air purification or photocatalysts for gaseous phase CO\(_2\) conversion. There are also problems when nano-sized TiO\(_2\) is used in liquid phase. The nano-sized TiO\(_2\) particles make it extremely difficult to separate from solutions to recover. Therefore, much attention is focused on the possibility of micron size TiO\(_2\) application as photocatalysts.

Targeting on making improvements such as safety, cost, high efficiency and long cycle life of the anode materials for LIBs, it is essential to find alternatives to replace the commercial graphite materials. The critical performance indicators for LIBs are capacity, cyclability and rate capability, which are strongly dependent on the active electrode materials. Graphitic carbons are most commonly used as conventional commercial anode materials owing to its low cost, relatively high energy density and very low flat working potential. However, graphitic carbon anode materials suffer the disadvantages of severe safety issues because of its low operating voltage (~0.1 V vs Li/Li\(^+\)), poor life cycle performance due to solid-electrolyte interface (SEI) film and the formation of dendritic lithium on the surface of graphitic carbons anode. Researchers in the scientific and industrial communities have investigated different materials such as silicon, tin (II) based oxide, and TiO\(_2\) as LIBs anode to replace graphite materials in order to eliminate the safety issue and meanwhile improve LIBs performance. Different from tin (II) based oxides and silicon, TiO\(_2\) material is structural stable during lithium insertion and extraction which means a very low volume change (<4% for Li\(_x\)TiO\(_2\), 0≤x≤1) for electrode. This advantage
guarantees its good stability during charge/discharge process at high current density rate. Moreover, its high working voltage (> 1.5 V vs Li/Li⁺) avoids reactions with organic electrolyte and short circuit caused by electrode expansion during cycling that make it safer than graphite anode material. However, TiO₂ anodes practical capacity is only 168 mAh g⁻¹ because of the poor electronic and ionic conductivity in its bulk form, limiting TiO₂ anode practical applications. These problems could be tackled via modification with carbon coating on TiO₂.

While the new generation of Li-S batteries system offers promising alternative for the energy requirements, it also presents challenges as mentioned above. To address the above challenges, various cathode materials for Li-S batteries have been explored to enhance electrical conductivity, reduce the active sulfur loss by preventing polysulfides from dissolving in the electrolyte, and accommodate the sulfur volume expansion. In 2009, Nazar and co-workers reported a pioneering work in the Li-S cathode material by incorporating sulfur into a highly ordered mesoporous carbon (CMK-3) and coating sulfur-CMK-3 with polyethylene glycol (PEG).¹⁵ Those materials exhibited good stability over 20 cycles tested and had high initial discharge capacities at 1,320 mA h g⁻¹ and 1,005 mA h g⁻¹ at 0.1 C rate (1 C = 1,675 mA/g) with and without PEG modification, respectively. Nazar’s work inspired the investigation of many carbon materials as sulfur substrates such as porous carbon frameworks,⁵⁹-⁶⁰ carbon nanotubes,³⁴ ⁶¹-⁶² hollow carbon nanofibers,¹⁶ graphene,⁶³-⁶⁴ and conductive polymers.¹⁵ ⁶⁵-⁶⁶ Cui et al.⁶⁵ reported a poly (3,4-ethylenedioxythiophene):poly (styrene sulfonate) (PEDOT:PSS) coated CMK-3/sulfur that could achieve 96-98% Coulombic efficiency.
Most of the above mentioned literature reports carbon materials having one-dimensional (1D) channel/pore structure or two-dimensional (2D) layered architectures with micropores (< 2 nm) or mesopores (6-8 nm). Three-dimensional (3D) hierarchical porous structures such as bulk carbon or nanoplates with a large surface area, good electrolyte permeability, and good electrochemical performance have also been designed in supercapacitors,\textsuperscript{67-68} lithium-ion batteries,\textsuperscript{69-70} and lithium-sulfur batteries.\textsuperscript{71} Regardless of the number of dimensions, pore structure and sizes are important factors that determine the electrochemical performance. It has been reported that mesopores can enhance the initial discharge capacity of carbon-sulfur composites and micropores can effectively restrain the shuttling effect.\textsuperscript{72-74} A 3D carbon framework can usually be designed with larger pores (e.g., macropores) that are in favor of ion transport and electrolyte permeation from various directions into the carbon porous structure.
1.5 Research Objectives of Outline of the Dissertation

One of the objectives of the research is to demonstrate novel TiO$_2$-based photocatalysts with hybrid composites structures and enhanced activity for CO$_2$ reduction and H$_2$ production and contribute more information about the unclear reaction mechanism. Another objective is to incorporate carbon and sulfur with novel structure to solve the problems that limit lithium-sulfur batteries technology. In order to achieve the research objectives, the research work is divided into five major tasks, which are summarized below.

Task 1. Nanostructured Ce-TiO$_2$/SBA-15 Composites for CO$_2$ Photoreduction (Chapter 2)

The objective of this task is to synthesize Ce-TiO$_2$/SBA-15 materials with high dispersion of TiO$_2$ and Ce could help facilitating electron-hole separation to decrease e-h pair recombination rate to enhance the activity. Three subtasks will be carried out:

- Ce-TiO$_2$ composites will be synthesized by sol-gel method.
- SBA-15 will be synthesized according to well-known method to be used as a support for Ce-TiO$_2$. Ce-TiO$_2$/aSiO$_2$ (amorphous silica) will be synthesized to compare with Ce-TiO$_2$/SBA-15.
- 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 CO$_2$ photoreduction.

Task 2. Ag/TiO$_2$ Nanocomposite Photocatalysts For Simultaneous H$_2$ Production And CO$_2$ Reduction (Chapter 3)
The objectives of this task are to fabricate and characterize Ag/TiO$_2$ nanocomposites via ultrasonic spray pyrolysis for simultaneous H$_2$ production and CO$_2$ reduction to produce syngas (CO+H$_2$) and to study the mechanism for the photocatalytic reaction. Three subtasks will be carried out:

- Ag/TiO$_2$ nanocomposites will be fabricated using ultrasonic spray pyrolysis and wet impregnation method.
- 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), and UV-vis diffuse reflectance spectra. And the nanomaterials will be applied to both water splitting and CO$_2$ photoreduction.
- The CO$_2$/H$_2$O/CH$_3$OH ratio will be adjusted for the photocatalytic reaction to check if we can produce syngas with tunable CO/H$_2$ ratio.

Task 3. Hybrid MgAl LDO-TiO$_2$ Composites for CO$_2$ Photoreduction (Chapter 4)

The objectives of this task are to fabricate and characterize MgAl layered double oxides grafted TiO$_2$ rods composites and apply the materials to CO$_2$ photoreduction. The mechanism of CO$_2$ adsorption-desorption ability provided by MgAl LDO on TiO$_2$ will be studied. Three subtasks will be carried out:

- MgAl LDO-TiO$_2$ with different MgAl LDO concentration will be fabricated by coprecipitation method. The morphology of MgAl LDO will be controlled by adjusting the Mg, Al amount.
- The resulting 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), thermogravimetric analysis (TGA) and UV-vis diffuse reflectance spectra. Finally, the nanomaterials will be applied to CO₂ photoreduction.

**Task 4. Carbon-TiO₂ Nanocomposites Prepared With Simple Method For Lithium-Ion Batteries (LIBs) (Chapter 5)**

The objectives of this task are to fabricate and characterize carbon-TiO₂ hybrid nanocomposites and apply the materials to LIBs. The mechanism of carbon formation on TiO₂ will be discussed and the electrochemical performance of C-TiO₂ hybrid materials will be studied. Four subtasks will be carried out:

- C-TiO₂ with different carbon concentration will be synthesized by a simple carbonization process. The carbon content will be controlled by adjusting the sucrose amount.

- The resulting 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), and thermogravimetric analysis (TGA). And the nanomaterials will be used as anode materials for LIBs.

- Electrodes and half-coin cell will be prepared and their performance will be investigated.

- The performance of C-TiO₂ nanomaterials as anode will be optimized.

**Task 5. Hierarchical Sulfur-Infiltrated Porous Microspheres Carbon For High Energy Lithium-Sulfur Batteries (Chapter 6)**

The objectives of this task are to fabricate and characterize hierarchical structure sulfur-infiltrated porous microsphere carbon and apply the materials to lithium-sulfur
batteries cathode. The mechanism of the porous structure for constraining intermediates polysulfide will be studied. Four subtasks will be carried out:

- Porous microsphere carbon with different pore size distribution will be fabricated by adjusting different silica nanoparticle template size or mixing ratio.
- 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), and thermogravimetric analysis (TGA). And the nanomaterials will be used as cathode materials for lithium-sulfur batteries.
- Electrodes and half-coin cell will be prepared and their performance will be investigated.
- The performance of PMC/S nanomaterials as cathode will be optimized.
CHAPTER 2 CHARACTERIZATION TECHNIQUES

X-ray Diffraction: X-ray diffraction is the primary method for determining the phase and the crystalline structure.

The crystal structures of the powder catalysts were identified by X-ray diffraction (XRD) (Scintag XDS 2000) using Cu Kα irradiation at 45 kV and a diffracted beam monochromator operated at 40 mA in the setting 2θ range at a scan rate of 1°/min. The crystal size of different crystal phases for TiO₂ was calculated by the Scherrer equation. The fractional phase content, \( W_A, W_B, \) and \( W_R \), for anatase, brookite, and rutile, respectively, are mathematically defined in Equations (1), (2) and (3).

\[
W_A = \frac{0.886 \times A_A}{(0.886 \times A_A + A_R + 2.721 \times A_B)} \tag{1}
\]

\[
W_B = \frac{2.721 \times A_B}{(0.886 \times A_A + A_R + 2.721 \times A_B)} \tag{2}
\]

\[
W_R = \frac{A_R}{(0.886 \times A_A + A_R + 2.721 \times A_B)} \tag{3}
\]

where \( A_A, A_B \) and \( A_R \) represent the integrated intensity of the anatase (1 0 1) peak (\( 2\theta = 25.28^\circ \)), the brookite (1 2 1) peak (\( 2\theta = 30.81^\circ \)), and the rutile (1 1 0) peak (\( 2\theta = 27.45^\circ \)), respectively. Because the brookite (1 2 0) (\( 2\theta = 35.3^\circ \)) and brookite (1 1 1) (\( 2\theta = 25.69^\circ \)) peaks overlap with the anatase (1 0 1) peak, \( A_A \) and \( A_B \) were calculated by the following method. Using the single isolated brookite (1 2 1) peak as a reference, the anatase (1 0 1), brookite (1 2 0) and brookite (1 1 1) overlapped peaks were deconvoluted by 0.9 and 0.8 intensity ratio for \( I_{(1 2 1)}^{(\text{anatase})}/I_{(1 2 1)}^{(\text{brookite})} \) and \( I_{(1 1 1)}^{(\text{brookite})}/I_{(1 2 0)}^{(\text{brookite})} \), respectively, with the same full width at half maximum (FWHM) of brookite (1 2 1).

BET Analysis: The Brunauer, Emmett and Teller (BET) technique is the most common method for determining the surface area of powders and porous materials. Nitrogen gas is
generally employed as the probe molecule and is exposed to a solid under investigation at liquid nitrogen conditions (i.e. 77 K).

The Brunauer-Emmett-Teller (BET) specific surface area and pore size of the catalysts were measured by nitrogen adsorption-desorption isotherms using a Micrometrics ASAP 2020 Surface Area and Porosity Analyzer.

For Ce-TiO$_2$/SiO$_2$ and C/TiO$_2$ materials: Before each adsorption measurement, approximate 0.10 g sample was degassed at 100 °C for 18 hours. The BET surface area was determined by a multipoint BET method using the adsorption data in a relative pressure (P/P$_0$) range 0.05-0.3. Pore size distribution was determined via Barret-Joyner-Halender (BJH) method using desorption isotherm. The N$_2$ adsorption volume at the relative pressure of 0.998 was used to determine the total pore volume and average pore size.

For PMC and PMC/S materials: Before each adsorption measurement, 0.10 g PMC sample was degassed at 300 °C for 4 h. It is not possible to follow the conventional protocol to degas the PMC/S samples above 155 °C before BET measurements owing to the volatility of the sulfur, and thus no pretreatment was done for PMC/S. The pore size distribution plot was derived from the adsorption branch of the isotherm based on the density functional theory (DFT).

**XPS Technique:** X-ray photoelectron spectroscopy (XPS) is a surface-sensitive quantitative spectroscopic technique that measures the elemental composition at the parts per thousand range, empirical formula, chemical state and electronic state of the elements that exist within a material. XPS spectra are obtained by irradiating a material with a beam of X-rays while simultaneously measuring the kinetic energy and number of electrons that escape from the top 0 to 10 nm of the material being analyzed. XPS requires high vacuum
(P $\sim 10^{-8}$ millibar) or ultra-high vacuum (UHV; P < $10^{-9}$ millibar) conditions, although a current area of development is ambient-pressure XPS, in which samples are analyzed at pressures of a few tens of millibar.

X-ray photoelectron spectroscopy (XPS) (Perkin-Elmer PHI 5100) was used to examine the valance states of Ce in the Ce/TiO$_2$ samples before and after CO$_2$ photoreduction with water vapor. A thin layer of powder samples were loaded on a silicon substrate and subject to XPS analysis. Subsequently, the same samples were exposed to photoillumination in the presence of CO$_2$ and water vapor for 4 h and then subject to XPS analysis again.

**SEM Characterization Technique:** A scanning electron microscope (SEM) is a type of electron microscope that produces images of a sample by scanning it with a focused beam of electrons. The electrons interact with atoms in the sample, producing various signals that can be detected and that contain information about the sample's surface topography and composition. The electron beam is generally scanned in a raster scan pattern, and the beam’s position is combined with the detected signal to produce an image. SEM can achieve resolution better than 1 nanometer.

Scanning electron microscopy (SEM) equipped with energy-dispersive X-ray (EDX) spectroscopy (Hitachi S4800) was used to analyze the catalyst morphology and surface elemental composition.

**TEM Characterization Technique:** Transmission electron microscopy (TEM) is a microscopy technique in which a beam of electrons is transmitted through an ultra-thin specimen, interacting with the specimen as it passes through. An image is formed from the interaction of the electrons transmitted through the specimen; the image is magnified
and focused onto an imaging device, such as a fluorescent screen, on a layer of photographic film, or to be detected by a sensor such as a CCD camera.

The particle size and crystal lattice of TiO$_2$, the particle size of Ag and the pore structure of SBA-15 were analyzed by transmission electron microscopy (TEM) (Hitachi H9000NAR) and high resolution TEM (HR-TEM).

**TGA Analysis:** Thermogravimetric analysis or thermal gravimetric analysis (TGA) is a method of thermal analysis in which changes in physical and chemical properties of materials are measured as a function of increasing temperature (with constant heating rate), or as a function of time (with constant temperature and/or constant mass loss). TGA is commonly used to determine selected characteristics of materials that exhibit either mass loss or gain due to decomposition, oxidation, or loss of volatiles (such as moisture).

The thermal stability of electrode materials were carried out on a thermal gravimetric analyzer (TGA-DAT-2960 SDT).

*For C/TiO$_2$: TGA were carried out at a heating rate of 20 °C min$^{-1}$ from 25 to 1000 °C in air.*

*For PMC/S: TGA were carried out at a heating rate of 20 °C min$^{-1}$ from 25 to 500 °C in argon.*

**UV-vis reflectance spectra:** Ultraviolet–visible spectroscopy or ultraviolet-visible spectrophotometry (UV-Vis or UV/Vis) refers to absorption spectroscopy or reflectance spectroscopy in the ultraviolet-visible spectral region. This means it uses light in the visible and adjacent (near-UV and near-infrared) ranges. The absorption or reflectance in the visible range directly affects the perceived color of the chemicals involved. In this region of the electromagnetic spectrum, molecules undergo electronic transitions.
The UV-vis reflectance spectra were recorded with a UV-vis spectrophotometer (Ocean Optics) using BaSO$_4$ as the background. The reflectance was converted to F(R) using the Kubelka-Munk (K-M) function \[ F(R) = \frac{(1-R)^2}{2R} \], and the band gap energy was obtained from the plot of \([F(R)E_{ph}]^{1/2}\) against the photon energy $E_{ph}$. 
CHAPTER 3  TIO₂-BASED PHOTOCATALYSTS FOR SOLAR FUEL PRODUCTION

3.1 Nanostructured Ce-TiO₂/SBA-15 Composites for CO₂ Photoreduction

According to the literature review results in Chapter 1, we hypothesize that this unique charge separation capacity with the assistance of Ce species could improve CO₂ photoreduction with water on TiO₂ photocatalysts. SBA-15 as support for Ce-TiO₂ will provide sufficient surface area and may have Si-OH group that is beneficial to CO₂ photoreduction.

3.2 Materials and Methods

The Ce-TiO₂ nanoparticles were synthesized by a sol-gel method. In a typical synthesis process, 10 mL titanium butoxide (Ti(OBu)₄, 99%) was ultrasonically dispersed in 40 mL absolute ethyl alcohol for 10 min (Solution A). A certain amount of Ce(NO₃)₃·6H₂O (99.5%) was dissolved in 10 mL H₂O, 10 mL absolute ethyl alcohol and 2 mL 62% nitric acid (Solution B). Then Solution A was added dropwise to Solution B (forming Solution C) with vigorous stirring for 3 h at room temperature. The obtained transparent sol was further aged for 6 h at room temperature, dried at 70 °C for 36 h and finally calcined in a muffle furnace at 500 °C for 2 h. Samples with molar ratios of Ce to Ti at 0.01, 0.03, 0.08 and 0.12 were prepared. For comparison, pristine TiO₂ was also prepared without adding the Ce precursor.

SBA-15 was prepared according to a well-established procedure reported by Zhao et. al. 4 g of Pluronic P123 (Aldrich) was dissolved in 125 g of 2 M HCl at 35 °C with stirring. Tetraethyl orthosilicate (TEOS, Aldrich) was then added into the solution after
P123 was dissolved. The resultant solution was stirred for 20 h at 35 °C, after which the mixture was aged at 80 °C for 24 h in a sealed Teflon bottle. The solid product was recovered by filtration and air-dried at room-temperature overnight. SBA-15 was obtained by calcination of the solid product at 500 °C for 6 h.

Ce-TiO$_2$ was loaded on SBA-15 by adding SBA-15 particles to Solution C in the sol-gel process of preparing Ce-TiO$_2$ as previously described. The same aging, drying, and calcination procedure was applied. Ce-TiO$_2$/SBA-15 composites with a molar ratio of Ce:Ti:Si at 0.03:1:1, 0.03:1:2, and 0.03:1:4 were prepared. For comparison, Ce-TiO$_2$ loaded on amorphous silica was also prepared following the same procedure, and the amorphous mesoporous silica particles were prepared according to the process reported in our previous study.$^5$ The prepared samples are denoted as $xCe_yTi$, $xCe_yTi_zSi$, and $xCe_yTi_z$aSi, where Ce represents cerium, Ti represents titania, Si represents SBA-15, aSi presents amorphous silica, and the numbers of x, y, and z indicate the molar ratio of Ce, Ti, and Si. For example, 0.03Ce_1Ti_4Si represents Ce-TiO$_2$/SBA-15 sample with a Ce:Ti:Si molar ratio at 0.03:1:4.

The Brunauer-Emmett-Teller (BET) specific surface area and pore size of the catalysts were measured by nitrogen adsorption-desorption isotherms using a Micrometrics ASAP 2020 Surface Area and Porosity Analyzer. The crystal structures of the powder catalysts were identified by X-ray diffraction (XRD) (Scintag XDS 2000) using Cu Kα irradiation at 45 kV and a diffracted beam monochromator operated at 40 mA in the 2θ range from 20° to 60° at a scan rate of 1°/min. The fractional phase contents of anatase, brookite, and rutile TiO$_2$ were calculated by the method reported in previous work.$^{11}$ The Scherrer equation was applied to calculate the crystal size of TiO$_2$. The UV–vis diffuse
reflectance spectra were recorded by a UV–vis spectrophotometer (Ocean Optic) using BaSO$_4$ as the background. Scanning electron microscopy (SEM) (Hitachi S4800) was used to obtain the catalysts surface morphology. The particle size and crystal lattice of TiO$_2$ and the pore structure of SBA-15 were analyzed by transmission electron microscopy (TEM) (Hitachi H9000NAR) and high resolution TEM (HR-TEM).

The photocatalytic activity of CO$_2$ reduction with water vapor was investigated in an experimental system as shown in Fig. 3.1. Compressed CO$_2$ (99.999%, Praxair) regulated by a mass flow controller (at a flow rate of 4 mL/min) passed through a deionized water bubbler to introduce CO$_2$ and water vapor mixture (volume fraction of H$_2$O ≈ 2.3%) into a photoreactor that has stainless steel walls and a quartz window. For each test, 200 mg powder catalyst was evenly dispersed on a glass-fiber filter and placed in the photoreactor facing the quartz window. A 450 W Xe lamp (Oriel) was used as the irradiation source with a light intensity around 400 mW/cm$^2$ (UV-vis region), measured by a spectroradiometer (International Light Technologies ILT950). Circulated cooling water was applied to absorb the infrared portion of the Xe lamp irradiation. The photoreactor was operating at a continuous-flow mode and the effluent gas sample was analyzed every 30 min by a gas chromatograph (GC, Agilent 7890A) equipped with an automated gas sampling valve, a thermal conductivity detector (TCD) and flame ionization detector (FID).
Figure 3.1 Experimental setup for photocatalytic CO₂ reduction with water vapor

3.3 Results and Discussion

Fig. 3.2 shows the XRD patterns of pristine TiO₂ and Ce-TiO₂ nanoparticles with different Ce:Ti molar ratios. The calculated fractional phase contents and crystal sizes of TiO₂ are summarized in Table 3.1. Pristine TiO₂ exhibited prominent characteristic peaks for anatase (JCPDS No. 21-1272) and rutile (JCPDS No. 21-1276), as well as a very weak peak for brookite (JCPDS No. 29-1360). It had the highest rutile phase fraction, 53.7%. With the addition of 0.01% Ce, the fraction of rutile phase remarkably decreased to 4.8% with significant increases in anatase and brookite phases. Further increasing the Ce:Ti ratio to 0.03 and 0.08 resulted in the disappearance of the rutile TiO₂. This result is in agreement with the literature reports that even a low concentration of ceria can stabilize the anatase phase and inhibit transformation of anatase to rutile. The stabilization mechanism was attributed to the preferential nucleation of ceria on the oxygen vacancies of anatase TiO₂. In addition, no obvious peaks corresponding to cubic Ce or cerium oxides were observed in the XRD patterns of Ce-TiO₂, probably due to the low concentration of Ce and the very small particle size of ceria, as well as its well dispersion on the TiO₂ surface.
The diffraction peaks of Ce-TiO$_2$ are broadened compared with pristine TiO$_2$, suggesting smaller crystal size of TiO$_2$ due to Ce addition. As summarized in Table 3.1, the average crystal sizes of pristine TiO$_2$ were 17 nm for anatase and 9.5 nm for brookite, and they both decreased with the Ce concentration. It has been reported that dopant in TiO$_2$ favors the formation of smaller crystals.$^{11,80}$ The smaller Ce-TiO$_2$ size in this work is an indirect evidence that Ce was partially doped in TiO$_2$ lattice. This also contributes to the reason that no Ce or CeO$_2$ peaks were observed in the XRD patterns. Table 3.1 also shows the BET specific surface area and pore size data. The pristine TiO$_2$ had the lowest surface area at 25.2 m$^2$/g. The incorporation of Ce in TiO$_2$ dramatically increased the surface area to 136.7 m$^2$/g for the sample of 0.03Ce$_1$Ti. This is likely due to the smaller particle size of 0.03Ce$_1$Ti as previously mentioned.

**Table 3.1** Fractional phase content, crystal size, BET specific surface area and pore size of TiO$_2$, SBA-15, Ce-TiO$_2$, Ce-TiO$_2$/SBA-15 and Ce-TiO$_2$/aSiO$_2$ samples

<table>
<thead>
<tr>
<th>Samples</th>
<th>Phase Content (%)</th>
<th>Crystal Size (nm)</th>
<th>S$_{\text{BET}}$ (m$^2$/g)</th>
<th>Pore Size (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A</td>
<td>B</td>
<td>R</td>
<td>A</td>
</tr>
<tr>
<td>TiO$_2$</td>
<td>34.5</td>
<td>11.7</td>
<td>53.7</td>
<td>17</td>
</tr>
<tr>
<td>0.01Ce$_1$Ti</td>
<td>66.3</td>
<td>28.9</td>
<td>4.8</td>
<td>7.6</td>
</tr>
<tr>
<td>0.03Ce$_1$Ti</td>
<td>71.5</td>
<td>28.5</td>
<td>0</td>
<td>5.7</td>
</tr>
<tr>
<td>0.08Ce$_1$Ti</td>
<td>100</td>
<td>0</td>
<td>0</td>
<td>3.6</td>
</tr>
<tr>
<td>SBA-15</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>0.03Ce$_1$Ti$_1$Si</td>
<td>100</td>
<td>0</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>0.03Ce$_1$Ti$_2$Si</td>
<td>100</td>
<td>0</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>0.03Ce$_1$Ti$_4$Si</td>
<td>100</td>
<td>0</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>0.03Ce$_1$Ti$_4$aSi</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
Figure 3.2 XRD patterns of pristine TiO$_2$ and Ce-TiO$_2$ samples with different Ce/Ti molar ratios: (a) pristine TiO$_2$; (b) 0.01Ce:1Ti; (c) 0.03Ce:1Ti; and (d) 0.08Ce:1Ti. (A: Anatase; B: Brookite; R: Rutile)

Figure 3.3 XRD patterns of Ce-TiO$_2$/SBA-15 samples with different Ti:Si molar ratios: (a) 0.03Ce:1Ti:1Si; (b) 0.03Ce:1Ti:2Si; and (c) 0.03Ce:1Ti:4Si.
**Fig. 3.3** shows the XRD patterns of Ce-TiO$_2$/SBA-15 samples. Only anatase TiO$_2$ diffraction peaks were observed and the noisy background was due to the silica substrate. The specific surface area was significantly increased due to the presence of SBA-15. It is interesting to find that the 0.03Ce$_{0.1}$Ti$_{0.4}$Si sample had a slightly smaller surface area than that of 0.03Ce$_{0.1}$Ti$_{0.2}$Si, considering more SBA-15 was in the 0.03Ce$_{0.1}$Ti$_{0.4}$Si sample. It may be because that the Ce-TiO$_2$ particles were better dispersed on the 0.03Ce$_{0.1}$Ti$_{0.4}$Si sample and partially blocked the pores of SBA-15, resulting in a smaller surface area.

N$_2$ adsorption-desorption was applied to explore the textual property of Ce-TiO$_2$ and Ce-TiO$_2$/SBA-15 catalysts. As shown in **Fig. 3.4**, compared to bare TiO$_2$, Ce-TiO$_2$ samples showed the typical type IV adsorption-desorption isotherms with a relatively wide H$_1$ hysteresis loop, an indicative of mesoporous structure. This mesopore may be resulted from the inter-space of aggregated nanoparticles. With Ce concentration increasing in Ce-TiO$_2$, the specific surface area is also increasing. It may be because that the Ce addition can help decrease particle size which leads to a higher specific surface area. Ce-TiO$_2$ samples showed the similar narrow BJH pore-size distribution with average pore diameter about 4-6 nm.

**Figure 3.4** N$_2$ adsorption-desorption isotherms (a) and BJH pore size distributions (b) of TiO$_2$ and Ce/TiO$_2$ samples.
Figure 3.5 N2 adsorption-desorption isotherms (a) and BJH pore size distributions (b) of SBA-15 and Ce-TiO2/SBA-15 samples.

Fig. 3.5 shows the N2 adsorption-desorption isotherms and BJH pore size distribution of SBA-15 and Ce-TiO2/SBA-15 samples. SBA-15 and Ce-TiO2/SBA-15 sample exhibits the typical type IV adsorption-desorption isotherm with a large H2 hysteresis loops. However, the H2 hysteresis loop in the P/P0 on pure SBA-15 ranges from 0.55 to 0.80, while that of Ce-TiO2/SBA-15 is from 0.45 to 0.70, which suggest that Ce-TiO2/SBA-15 samples have large uniform mesopores without changing the pore frame of SBA-15. The different P/P0 positions of the inflection points between SBA-15 and Ce-TiO2/SBA-15 indicate their different pore widths in the mesopore rage. As evidenced in Fig. 3.5b, SBA-15 shows a narrow pore size distribution with the diameter of 5.3 nm. In contrast, Ce-TiO2/SBA-15 photocatalysts display a lower pore size distribution around 4.1-4.7 nm. This narrow is probably resulted from the interaction between Ce-TiO2 and SBA-15 that may block some pore openings. Interestingly, 0.03Ce_1Ti_4Si displays two pore size distribution at 3.6 nm and 5.5 nm. This could be resulted from the Ce-TiO2 particles filling into SBA-15 pores and also the pores between the disordered Ce-TiO2 particles on the surface of SBA-15.
Figure 3.6 SEM images of the 0.03Ce_1Ti_4Si sample.

Detailed morphological information of the Ce-TiO₂/SBA-15 sample (0.03Ce_1Ti_4Si) was examined by SEM and TEM. The SEM images in Fig. 3.6 demonstrate micrometer size particles with irregular shapes mainly in rods and spheres, which are assigned to SBA-15 particles. The TiO₂ nanoparticles are not discernible in the SEM images but are clearly shown in the TEM images in Fig. 3.7. Fig. 3.7a-b demonstrates the morphology of bare SBA-15 samples, where one-dimensional channels of ordered pores are clearly identified. The pore opening is around 8 nm, consistent with the pore size analysis data shown in Fig. 3.5. Fig. 3.7c demonstrates that for the Ce-TiO₂/SBA-15 sample, the TiO₂ nanoparticles are dispersed on the surface, at the edge, and possibly in the pores of the SBA-15 support. The fraction of TiO₂ nanoparticles embedded in the pores of SBA-15 may be very small, but it may increase as the SBA-15 loading increases. These embedded particles may block a portion of the pores, which explains why the measured specific surface area of the 0.03Ce_1Ti_4Si sample is smaller than that of the 0.03Ce_1Ti_2Si sample (see Table 3.1). Aggregates of TiO₂ nanoparticles on the outer surface of SBA-15 were also identified, as shown in Fig. 3.7d. The TiO₂ crystal size is around 5 nm, as shown in the HR-TEM image in Fig. 3.7e, which matches the calculated crystal size listed in Table 3.1 for 3%Ce-TiO₂ sample. Clear lattice fringes of nanocrystal
with an interplanar space of 0.350 nm were observed, which corresponds to the (101) plane of anatase TiO$_2$.\textsuperscript{81}

**Figure 3.7** TEM and HRTEM images of SBA-15 (a,b) and 0.03Ce$_1$Ti$_4$Si$_2$ (c,d,e) samples.

Diffuse reflectance UV-vis spectra were recorded to explore the influence of Ce and
SBA-15 addition on the optical property of TiO\(_2\). As shown in Fig. 3.8a, pristine TiO\(_2\) absorbs light at wavelength shorter than 400 nm, corresponding to a band gap about 3.0 eV. Incorporation of Ce in TiO\(_2\) resulted in a red-shift in the absorption edge, extending to the visible light region in the range of 400 ~ 500 nm. The red-shift was enhanced at an increased Ce concentration, but the enhancement was not significant when the Ce concentration exceeded 3%. Fig. 3.8b shows the UV-vis spectra of Ce-TiO\(_2\)/SBA-15 samples. In contrast to the red-shift effect brought by the Ce doping, the incorporation of SBA-15 as the support resulted in a blue-shift of the absorption spectra, which is signified at a higher SBA-15 loading. This result is in agreement with the literature that a blue shift of light absorption edge of TiO\(_2\)/SBA-15 has been observed compared with pure anatase TiO\(_2\) or P25,\(^{82-84}\) probably due to the quantum size effect of smaller TiO\(_2\) nanocrystals in the SBA-15 matrix.\(^{85}\)

![Figure 3.8](image)

**Figure 3.8** UV–vis diffuse reflectance spectra of TiO\(_2\) and Ce-TiO\(_2\) with different Ce:Ti molar ratios (a) and Ce-TiO\(_2\)/SBA-15 with different Ti:Si molar ratios (b).

The results of XPS for the 0.03Ce_1Ti sample are shown in Fig. 3.9. The energy interval between approximately 882 eV and 903 eV was attributed by Ce 3d 5/2, while the Ce 3d 3/2 was at the 905 eV to 918 eV interval. To perform quantitative analysis, the
background was first subtracted using Shirley approximation function in Augerscan software, and then the 3d 3/2 and 3d 5/2 peak structures were fitted. There were in total 10 peaks fitted for the XPS spectrum between 870 and 920 eV, where Ce$^{4+}$ (CeO$_2$) has six peaks (#5 – 10) for the total 3d lines due to strong hybridization of the oxygen 2p valence band with the Ce 4f orbital, while Ce$^{3+}$ (Ce$_2$O$_3$) has four peaks (#1 – 4). Relative ratio/concentrations of Ce$^{3+}$/Ce$^{4+}$ were calculated based on the summation of the corresponding peak areas, which gives a Ce$^{3+}$/Ce$^{4+}$ ratio equal to 55.3%/44.7% before photoreaction and 49.5%/50.5% after photoreaction. The same analysis was done for the 0.08Ce_1Ti sample (XPS spectra not shown here), and the Ce$^{3+}$/Ce$^{4+}$ ratio was 51.8%/48.2% before photoreaction and 52.7%/47.2% after photoreaction. These results indicate that the Ce$^{3+}$/Ce$^{4+}$ ratios were similar for the different Ce/TiO$_2$ samples and almost did not change during the CO$_2$ photoreduction process.

![XPS spectra of Ce 3d](image)

**Figure 3.9** XPS spectra of Ce 3d of the 0.03Ce_1Ti sample before (a) and after (b) CO$_2$ photocatalytic reduction with water vapor.

In this study, CO and CH$_4$ were identified as the main products of CO$_2$ photoreduction with H$_2$O on Ce–TiO$_2$ and Ce–TiO$_2$/SBA-15 nanocomposites. This was in line with our previous published work$^5$, $^52$, $^87$ that CO and CH$_4$ are the main products of CO$_2$ photoreduction with water vapor. **Fig. 3.10** shows the production rates of CO and CH$_4$ on
selected photocatalysts as a function of photo-illumination time. For all catalysts, the production rates gradually decreased after reaching the maximum values, indicating a gradual deactivation of the catalytic performance. Similar trend in deactivation has been observed in our previous studies\textsuperscript{5, 88} and in the literature reports.\textsuperscript{89-90} Possible reasons are surface coverage of reaction intermediates or re-oxidation of the reaction products.\textsuperscript{89} To better compare the activities of the different photocatalysts, accumulative yields during the 4 h photo-illumination were calculated by integrating the production rate with time. The comparison results are presented as follows.

\textbf{Figure 3.10} Production rate of (a) CO and (b) CH\textsubscript{4} over TiO\textsubscript{2}, 0.03Ce\textsubscript{1}Ti, 0.03Ce\textsubscript{1}Ti\textsubscript{4}Si and 0.03Ce\textsubscript{1}Ti\textsubscript{4}aSi catalysts under UV-vis illumination.
Figure 3.11 Product yields of CO and CH₄ production over TiO₂ and Ce-TiO₂ catalysts under UV-vis illumination for 4 hr.

Fig 3.11 shows the accumulative yields of CO and CH₄ production during a 4 h photo-illumination period using pristine TiO₂ and Ce-TiO₂ samples with different Ce:Ti ratios. Pristine TiO₂ exhibited very low photocatalytic activity, with a CO yield at 0.25 μmol g⁻¹ and a CH₄ yield at 0.05 μmol g⁻¹. Ce-modification in TiO₂ significantly influenced CO production. At 1% and 3% Ce concentration, the CO yield reached approximately 1.0 μmol g⁻¹, four times as high as the pristine TiO₂. However, further increasing the Ce concentration to 8% resulted in a lower CO yield, and at 12% Ce concentration, the CO yield was even lower than that of pristine TiO₂. The CH₄ yields of all the TiO₂ and Ce-TiO₂ samples remained in the same order and at least one magnitude lower than CO yields, indicating that CH₄ yield was not affected by the Ce addition.

The enhanced CO production due to low concentration Ce-modification on TiO₂ can be attributed to the following reasons. First, Ce-dopant decreased the crystal size of TiO₂ and increased the surface area, as shown in Table 3.1, thus enhancing the catalytic
activity. Second, Ce-dopant inhibited the transformation of anatase to rutile phase and promoted the growth of both anatase and brookite phases, as shown in Table 3.1. It is well known that anatase TiO$_2$ is photocatalytically more active than rutile, but a mixture of the two phases can be advantageous. The above results indicate that a binary anatase-brookite with dominating anatase phase at low Ce concentration could promote CO$_2$ photoreduction activity.

Another important reason for the enhanced activity may be related to the Ce$^{3+}$/Ce$^{4+}$ redox couple on TiO$_2$ that facilitates electron-hole separation. It is well known that Ce$^{4+}$ ions can trap photo-excited electrons at the CeO$_2$/TiO$_2$ interface through reaction (1).\textsuperscript{7, 36, 38, 91} The Ce$^{3+}$ ions can then react with gas-phase O$_2$, if any (reaction (2)) to regenerate Ce$^{4+}$, or react with CO$_2$ (reaction (3)) in the case of this study, to form surface adsorbed CO$_2^-$ that can be further reduced to CO (reaction (4)). The formation of CO$_2^-$ species as CO$_2$ reduction intermediates has been confirmed by our previous in situ DRIFTS study using Cu/TiO$_2$ catalysts.\textsuperscript{88} Hence, the presence of Ce$^{4+}$ enhanced electron trapping and transfer to CO$_2$.

\begin{align*}
  e^- + Ce^{4+} & \rightarrow Ce^{3+} \quad (1) \\
  Ce^{3+} + O_2 & \rightarrow Ce^{4+} + O_2^- \quad (2) \\
  Ce^{3+} + CO_2 & \rightarrow Ce^{4+} + CO_2^- \quad (3) \\
  CO_2^- + e^- & \rightarrow CO + O_2^- \quad (4) \\
  Ce^{3+} + h^+ & \rightarrow Ce^{4+} \quad (5)
\end{align*}

On the other hand, it is possible that Ce$^{3+}$ ions can be directly oxidized back to Ce$^{4+}$ by photo-excited holes, according to reaction (5),\textsuperscript{7} which may diminish the promoting
effect of electron transfer as described above. However, the pre-existing $\text{Ce}^{3+}$ ions on the $\text{TiO}_2$, in the case of Ce-TiO$_2$ samples prepared in this work, can provide abundant $\text{Ce}^{3+}$ sites to promote reaction (3). Furthermore, the hole trapping effect of $\text{Ce}^{3+}$ can also prevent holes from re-oxidizing the reaction product (i.e., CO) back to CO$_2$. Therefore, the $\text{Ce}^{3+}/\text{Ce}^{4+}$ redox couples are advantageous over $\text{Ce}^{4+}$ or $\text{Ce}^{3+}$ alone to promote charge separation and enhance CO$_2$ photoreduction. The XPS result that the $\text{Ce}^{3+}/\text{Ce}^{4+}$ ratio almost did not change during the photo-reaction was another evidence for the abovementioned $\text{Ce}^{3+}/\text{Ce}^{4+}$ redox reactions that remained at a dynamic equilibrium.

In this set of experiments, a fixed amount of 200 mg Ce-TiO$_2$/SBA-15 was used for each test, while the Ce concentration relative to TiO$_2$ remained the same at 3% and the Ti:Si ratio varied from 1 to 4. Compared with the bare Ce-TiO$_2$ sample, dispersing the Ce-TiO$_2$ nanoparticles on the SBA-15 matrix resulted in a significant enhancement in the yields of CO and CH$_4$ production, as shown in Fig. 3.12. The enhancement seemed to be proportional to the loading of SBA-15 used. Among the three Ce-TiO$_2$/SBA-15 samples tested, the sample with the highest SBA-15 loading, 0.03Ce$_1$Ti$_4$Si, showed the highest activity for both CO and CH$_4$ production, reaching 7.5 and 7.9 μmol g$^{-1}$, respectively. Compared with the 0.03Ce$_1$Ti sample, the 0.03Ce$_1$Ti$_4$Si sample demonstrated 8-fold enhancement in CO production and 115-fold enhancement in CH$_4$ production.
Figure 3.12 Product yields of CO and CH$_4$ production over Ce-TiO$_2$/SBA-15 and Ce-TiO$_2$/aSiO$_2$ catalysts under UV-vis illumination for 4 hr.

This superior activity of SBA-15 supported Ce-TiO$_2$ catalysts is believed to be mainly attributed to the higher surface area and better dispersion of the Ce-TiO$_2$ nanoparticles, as verified by the BET and TEM analyses. Moreover, SBA-15 as the support may enhance the stability of the TiO$_2$ anatase phase and prevent the grain growth of TiO$_2$ nanocrystals.$^{92}$ This mechanism was also supported by our XRD results that 100% anatase TiO$_2$ was present in Ce-TiO$_2$/SBA-15 samples and supported by our UV-vis DRS results that smaller TiO$_2$ nanoparticles were present due to the SBA-15 confinement effect leading to a blue-shift in light absorption. In addition, mesoporous silica is also a well-known CO$_2$ adsorbent that may increase the localized concentration of CO$_2$ near the surface of TiO$_2$, and thus improve CO$_2$ adsorption on TiO$_2$ and subsequent reduction. In a recent study by Yang et al.$^{47}$ an enhanced photocatalytic CO$_2$ reduction was observed on a 2%Cu-TiO$_2$/SBA-15 sample (45 wt% TiO$_2$ loading) compared with 2%Cu-TiO$_2$; however,
the enhancement due to the SBA-15 support was only 20%. In our work, the enhancement factor was much more significant on Ce-TiO$_2$/SBA-15, suggesting the superior effect of Ce addition. Furthermore, the dramatic 115-fold enhancement in CH$_4$ production observed in this study is very intriguing. Because eight protons are needed for generation of one CH$_4$ molecule in comparison with two electrons for CO generation, the surface Si-OH groups or active OH sites due to the presence of SBA-15 may be more readily available for CO$_2$ reduction and conversion to CH$_4$. Investigations on this high CH$_4$ selectivity as well as the potential synergies between the Ce species, TiO$_2$ and the SBA-15 support are in our future research plan.

Another important finding in this study was that Ce-TiO$_2$ dispersed on SBA-15 was much more active than that dispersed on amorphous mesoporous silica. As shown in Fig. 3.12, the CO yield on the amorphous silica supported sample, 0.03Ce$_1$Ti$_4$Si$_4$ was very small, even lower than the catalyst without silica support. The CH$_4$ yield of 0.03Ce$_1$Ti$_4$Si$_4$ was 1.5 μmol g$^{-1}$, higher than the catalyst without silica support but still much lower than the catalyst supported on SBA-15. The overall CO$_2$ reduction rate to CO and CH$_4$ was more than 10 times higher on 0.03Ce$_1$Ti$_4$Si than on 0.03Ce$_1$Ti$_4$Si. Since the specific surface area of the 0.03Ce$_1$Ti$_4$Si sample was only about twice as much as that of the 0.03Ce$_1$Ti$_4$Si sample (Table 3.1), the much higher activity of the SBA-15 supported sample should result from other factors besides the surface area effect.

The possible reason is that ordered pores of SBA-15 are pores accessible to TiO$_2$ precursors during the preparation process, leading to a better dispersion of TiO$_2$ nanoparticles. Amorphous silica with random arrays of pores and shapes in some cases are
poor supports for functional agents because not all the volume of irregular pores is accessible to the incorporated species.\textsuperscript{93}

### 3.4 Conclusion

A novel Ce-TiO\textsubscript{2}/SBA-15 nanocomposite was synthesized for the first time in the literature and tested as a photocatalyst for converting CO\textsubscript{2} and H\textsubscript{2}O to fuels such as CO and CH\textsubscript{4} under photo-illumination. Modification of TiO\textsubscript{2} with Ce significantly stabilized the TiO\textsubscript{2} anatase phase and increased the specific surface area, which contributed to an improvement of CO production from CO\textsubscript{2} reduction. Dispersing Ce-TiO\textsubscript{2} nanoparticles on the mesoporous SBA-15 support further enhanced both CO and CH\textsubscript{4} production. Particularly the CH\textsubscript{4} production was enhanced by up to 115 times compared with unsupported Ce-TiO\textsubscript{2}. The superior catalytic activity may be related to the partially embedded Ce-TiO\textsubscript{2} nanoparticles in the ordered 1-D pores in SBA-15 that form synergies between the different components of the catalysts and enhance the diffusion and adsorption of CO\textsubscript{2}. This mechanism also correlates well the results that using SBA-15 as the support led to more than 10 times higher activity in CO\textsubscript{2} photoreduction than using amorphous silica as the support.
CHAPTER 4 AG/TIO\textsubscript{2} NANOCOMPOSITE PHOTOCATALYSTS FOR SIMULTANEOUS H\textsubscript{2} PRODUCTION AND CO\textsubscript{2} REDUCTION

4.1 Introduction

As shown in Fig. 1.5, both photocatalytic H\textsubscript{2} production and CO\textsubscript{2} reduction using water as the reductant processes could occur concurrently and compete with each other for photo-induced electrons. However, the benefit of having both reactions in one system is the possibility of producing syngas, a mixture of H\textsubscript{2} and CO, which is a valuable feedstock in producing synthetic petroleum through Fischer-Tropsch process. According to the literature review, the first objective of this work was to explore the feasibility of syngas production from CO\textsubscript{2} and water vapor through a photocatalytic process using a silver-modified TiO\textsubscript{2} (Ag/TiO\textsubscript{2}) nanoparticle catalyst. Ag was selected because it is much less expensive than Pt and Ag/TiO\textsubscript{2} has shown effectiveness in H\textsubscript{2} production\textsuperscript{53} and CO\textsubscript{2} photoreduction\textsuperscript{48} in separate studies.

Another unique feature of this work is the method of preparing the Ag/TiO\textsubscript{2} catalyst using a simple ultrasonic spray pyrolysis (USP) process. Conventional preparation methods for Ag/TiO\textsubscript{2} catalysts include sol-gel\textsuperscript{48, 53, 94} and wet impregnation,\textsuperscript{95} which are batch processes and require several after-treatment steps. USP offers several advantages over conventional material processing techniques, including a narrow particle size distribution and a high purity of the product powder.\textsuperscript{96-98}

Among the limited number of publications, no studies have been investigated on USP synthesized Ag/TiO\textsubscript{2} catalysts for simultaneous H\textsubscript{2} production and CO\textsubscript{2} reduction. Therefore, the second objective of this work was to explore the potential advantages of
USP synthesis of Ag/TiO$_2$ catalysts over conventional wet-impregnation method with regard to material property and catalytic activity.

4.2 Materials and Methods

The apparatus for ultrasonic spray pyrolysis process and the formation mechanism of Ag/TiO$_2$ nanoparticle composites are shown in Fig. 4.1. The spraying solution was prepared by mixing aqueous AgNO$_3$ solution and TiO$_2$ (Degussa P25) nanoparticles. The TiO$_2$ concentration was fixed at 16 mg/ml and the amount of AgNO$_3$ was varied so that the nominal weight percentage of Ag to TiO$_2$ in the catalysts was in the range of 1%-8%. The ultrasonic nebulizer (Sonaer) has a frequency of 2.4 MHz and produces droplets with a volumetric mean diameter of 1.8 mm. The generated mists containing AgNO$_3$/TiO$_2$ were carried out by N$_2$ and passed through a diffusion dryer composed of silica gels. The particles then passed through a cylindrical quartz reactor that was placed inside a tube furnace (Thermo Fisher) with a set temperature of 550 $^\circ$C. The spray-pyrolyzed particles were collected on a glass fiber filter placed inside a stainless-steel filter holder. As a control, Ag modified TiO$_2$ samples with the same nominal Ag weight percentages were also synthesized by a benchmark wet-impregnation method by mixing and stirring desired amount of TiO$_2$ (P25) in AgNO$_3$ aqueous solution at 80 $^\circ$C overnight and finally calcining the sample in air environment at 400 $^\circ$C for 4 h. To have a comparable basis for material property, the spray pyrolysis samples were also calcined in air at 400 $^\circ$C for 4 h as a final step. The samples were denoted as m%Ag/TiO$_2$-SP or m%Ag/TiO$_2$-WI, where m represents the nominal Ag weight percentage, SP represents materials prepared by spray pyrolysis method, and WI represents materials prepared by wet impregnation method.
The Brunauere-Emmete-Teller (BET) specific surface area of the catalysts was measured by nitrogen adsorption using a surface area and porosity analyzer (Micrometrics ASAP2020). The crystal structures of the catalysts were identified by X-ray diffraction (XRD) (Scintag XDS 2000) using Cu Ka irradiation at 45 kV and a diffracted beam monochromator operated at 40 mA in the 2θ range from 20° to 80° at a scan rate of 2°/min. The UV-vis reflectance spectra were recorded with a UV-vis spectrophotometer (Ocean Optics) using BaSO₄ as the background. The reflectance was converted to F(R) using the Kubelka-Munk (K-M) function \[ F(R) = \frac{(1-R)^2}{2R} \], and the band gap energy was obtained from the plot of \[ F(R)E_{ph}^{1/2} \] against the photon energy \( E_{ph} \).\(^8\) Scanning electron microscopy (SEM) equipped with energy-dispersive X-ray (EDX) spectroscopy (Hitachi S4800) was used to analyze the catalyst morphology and surface elemental composition. Transmission electron microscopy (TEM, Hitachi H9000NAR) and high-resolution TEM were used to examine the particle size and dispersion of Ag nanoparticles on TiO₂ as well as the lattice information.

The schematic of the photocatalytic reaction system is illustrated in Fig. 4.2. Compressed N₂ (99.998%, Praxair) or CO₂ (99.999%, Praxair) was used as the carrier gas.
for measuring the catalytic activity of H$_2$ production or CO$_2$ reduction, respectively. The carrier gas regulated by a mass flow controller passed through a bubbler containing 5 vol% methanol in de-ionized water to generate a water-methanol vapor mixture. Methanol was expected to serve as the hole scavenger and thus enhance the rate of reduction by electrons.$^{87, 99-100}$ Estimated from the saturation vapor pressure-temperature profile,$^{101}$ the volume fraction of H$_2$O and CH$_3$OH in the gas mixture at 20 °C was calculated to be 2.2% and 0.6%, respectively, balanced with the carrier gas (N$_2$ or CO$_2$). The gas mixture was then introduced into a photoreactor. 100 mg powder catalysts were dispersed on a glass fiber filter and placed at the bottom of the reactor. A 150 W Solar Simulator (Oriel) was used as the photo-excitation source. Before photo-illumination, the reactor was purged with the carrier gas (N$_2$ or CO$_2$) at a flow rate of 120 ml/min for 1 h and then at 4 ml/min for 1 h. During the illumination, the carrier gas flow rate maintained at 4 ml/min and the sample of effluent gas was taken every 30 min by an automated valve to a gas chromatograph (GC, Agilent 7890A). The GC was equipped with a thermal conductivity detector (TCD) and flame ionization detector (FID).

![Figure 4.2](image.png)  
**Figure 4.2** Experimental system for photocatalytic H$_2$ production and/or CO$_2$ reduction.
4.3 Results and Discussion

Fig. 4.3 compares the XRD patterns of Ag/TiO$_2$-SP and Ag/TiO$_2$-WI samples. All patterns indicate a mixture of TiO$_2$ anatase and rutile phases, which agrees with the composition of P25 (approximately 80% anatase and 20% rutile). At a lower Ag concentration (i.e., 2%), no diffraction lines of Ag species were observed, suggesting that Ag is well dispersed or the particle size is very small. At a higher Ag concentration (i.e., 8%), the SP sample showed peaks at $2\theta = 38.2^\circ$, $44.4^\circ$, $64.5^\circ$ and $77.4^\circ$ that correspond to metallic silver.$^{53,102-104}$ This indicates that Ag ions are thermally reduced during the spray pyrolysis process.$^{105}$ In contrast, for the 8% Ag/TiO$_2$-WI sample, neither metallic silver or silver oxides were observed. The result implies that silver oxides could be the major Ag species on WI samples as some literature publications reported the presence of Ag$_2$O in a Ag/TiO$_2$ composite prepared by impregnation$^{95}$ without detecting Ag$_2$O diffraction peaks in the XRD patterns.

Figure 4.3 XRD patterns of (a) 2% Ag/TiO$_2$-SP, (b) 2% Ag/TiO$_2$-WI, (c) 8% Ag/TiO$_2$-SP and (d) 8% Ag/TiO$_2$-WI.
Table 4.1 Comparison of BET and EDX analysis results for 2% Ag/TiO$_2$ samples prepared by SP and WI methods.

<table>
<thead>
<tr>
<th>Sample</th>
<th>BET Analysis</th>
<th>EDX Elemental Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Specific Surface Area (m$^2$/g)</td>
<td>Pore Volume (cm$^3$/g)</td>
</tr>
<tr>
<td>2% Ag/TiO$_2$-SP</td>
<td>54.2</td>
<td>0.34</td>
</tr>
<tr>
<td>2% Ag/TiO$_2$-WI</td>
<td>43.8</td>
<td>0.27</td>
</tr>
</tbody>
</table>

The textural properties of the Ag/TiO$_2$ samples were characterized by BET, SEM/EDX, and TEM/HRTEM analysis. Table 4.1 compares the results of BET analysis for the SP and WI samples with the same nominal Ag concentration (i.e., 2%). The SP sample had a larger specific surface area (54.2 m$^2$/g) compared with the WI sample (43.8 m$^2$/g). The pore volume and average pore size for the SP sample were also larger than those of the WI sample. The results suggest that the spray pyrolysis process may lessen the degree of agglomeration of the TiO$_2$ nanoparticles through the solvent evaporation process, resulting in a larger pore size and a larger surface area. The pore size in the range of 15-18 nm indicate a mesoporous structure for both samples, which is confirmed by the SEM images shown in Fig. 4.4. The SP samples (Fig. 4.4a) are mesoporous spheres composed of the primary nanoparticles while the WI samples (Fig. 4.4b) are irregular-shape agglomerates of the primary nanoparticles. TEM and HRTEM images for the SP and WI samples are shown in Fig. 4.5. It is clear that Ag nanoparticles in the size range of 1-2 nm are well dispersed on crystalline TiO$_2$ particles (~20 nm, agreeing with P25 primary particle size) for both SP and WI samples. However, some “big” Ag nanoparticles (~8 nm, see Fig.
4.5d) and aggregates of Ag nanoparticles (see Fig. 4.5e) were found on the WI sample, indicating that the Ag dispersion on the WI sample was worse than on the SP sample.

Figure 4.4 SEM images of 2% Ag/TiO$_2$-SP (a) and 2% Ag/TiO$_2$-WI (b).
Figure 4.5 TEM images of (a) and (b) 2% Ag/TiO$_2$-SP; (d) and (e) 2% Ag/TiO$_2$-WI and HRTEM images of (c) 2% Ag/TiO$_2$-SP; (f) 2% Ag/TiO$_2$-WI

The EDX elemental analysis data for the 2% Ag/TiO$_2$ samples are listed in Table 4.1. A major difference between the SP and WI samples is the measured Ag concentration, although both of them had the same nominal Ag concentration (i.e., 2 wt%) calculated from the precursor composition in the starting solution. For the WI sample, the measured Ag concentration (2.2 wt%) was very close to the nominal concentration, while for the SP sample, the measured Ag concentration was only 1.2 wt%. A possible reason for the lower Ag concentration on SP samples could be that a portion of the Ag nanoparticles formed during the spray pyrolysis escaped from the TiO$_2$ surface along with the solvent evaporation process and these Ag nanoparticles were not collected on the filter due to their extremely small size.
**Fig. 4.6** shows the results of band gap analysis derived from the UV-vis diffuse reflectance spectra. Pure TiO\(_2\) P25 has a band gap of 3.20 eV. With incorporation of Ag species, the band gap slightly decreased to 3.0-3.1 eV for SP samples and 2.9-3.0 eV for WI samples. However, the change in band gap is not an apparent function of Ag concentration. The shift of band gap is probably due to the interaction of Ag ions with TiO\(_2\) or formation of impurity band inside the TiO\(_2\) band gap.\(^{95,106-107}\) A broad peak around 2.35 eV (or 500-550 nm in absorption spectra) was observed for both SP and WI samples with higher Ag concentration (>4 wt\%). This unique peak is an indication of the localized surface plasmon resonance (SPR) of Ag nanoparticles,\(^{95,108}\) which again confirms the presence of metallic Ag on TiO\(_2\).

![Diffuse reflectance spectra](image)

**Figure 4.6** Diffuse reflectance spectra of TiO\(_2\), Ag/TiO\(_2\) (a) prepared by ultrasonic spray pyrolysis (b) prepared by wet impregnation.

The experiments of photocatalytic hydrogen production were carried out by using N\(_2\) as the carrier gas to bubble through the methanol/water mixture. Since the reactor was running in a continuous-flow mode, the H\(_2\) production rate gradually increased and reached a steady value after approximately 4 h of photo-illumination (the curves of H\(_2\) production...
rate with illumination time are not shown here). **Fig. 4.7** summarizes the steady-state H₂ production rate for SP and WI samples with different Ag concentrations. For bare TiO₂, the H₂ production rate was very small for both SP and WI samples. The incorporation of Ag species significantly enhanced the H₂ production rate. For Ag/TiO₂ samples with Ag concentration greater than 1%, the SP samples were much more active than the WI samples and the optimal Ag concentration appeared to be 2%. For 2% Ag/TiO₂-SP, the H₂ production rate was 1130 mmol/g/h, 6 times as much as that of 2% Ag/TiO₂-WI (180 mmol/g/h), and 60 times as much as that of bare TiO₂-SP (19 mmol/g/h). Ag concentration greater than 2% resulted in a lower H₂ production rate, probably because too high a metal concentration on TiO₂ leads to the formation of electron-hole recombination centers.

**Figure 4.7** Rates of photocatalytic H₂ production for Ag/TiO₂-SP and Ag/TiO₂-WI samples at different Ag concentration (the reaction gas was a mixture of 2.2% H₂O and 0.6% CH₃OH balanced with N₂).
The H$_2$ production rate of the 2% Ag/TiO$_2$-SP was compared with the literature data. Park et al. \textsuperscript{53} reported that the H$_2$ production rate was 8560 mmol/g/h by a Ag/TiO$_2$(rutile) powder catalyst suspended in a methanol/water (1:1 v/v) solution under 365 nm UV irradiation at an intensity of 36 W/cm$^2$. The relatively high yield was attributed to the extremely high light intensity. Alenzi N. et al. \textsuperscript{100} reported the average H$_2$ production rate of Ag/TiO$_2$ (anatase) film was 148 mmol/h/g in a methanol/water (1:1 v/v) solution system (4.7 mmol/h/g if without Ag) under UV irradiation at 10 mW/cm$^2$. Onsuratoom et al. \textsuperscript{99} reported the highest H$_2$ production rate was 4 cm$^3$/g/h or 178 mmol/g/h on an optimized Ag/TiO$_2$-ZrO$_2$ photocatalyst (Ag = 0.5 wt%) in a methanol/water (1:1 v/v) aqueous system under UV light at 2.3 mW/cm$^2$. In this work, the H$_2$ production rate of 2% Ag/TiO$_2$-SP was 1130 mmol/g/h, with a comparable light intensity of 12.5 mW/cm$^2$ (for 200 < $\lambda$ < 400 nm) and a much lower methanol/water ratio (0.6:2.2 v/v). Comparisons with the literature data indicate that the 2% Ag/TiO$_2$-SP catalyst we synthesized is a very effective photocatalyst toward H$_2$ production.

The experiments of photocatalytic CO$_2$ reduction were carried out by using CO$_2$ as the carrier gas to bubble through the methanol/water mixture. \textbf{Fig. 4.8} shows the photocatalytic activity of CO$_2$ reduction with the same sets of catalysts used in the H$_2$ production tests. CO was found to be the major product with minor CH$_4$ production. Like the trend observed for H$_2$ production, the incorporation of Ag on TiO$_2$ significantly increased the CO$_2$ reduction rate and the SP samples generally had higher activity than WI samples. The production rates of both CO and CH$_4$ were also affected by the Ag concentration. As shown in \textbf{Fig. 4.8a} 2% Ag/TiO$_2$-SP demonstrates the highest CO production rate at 87 mmol/h/g, 50% more than that of 2% Ag/TiO$_2$-WI (57 mmol/h/g). As
shown in Fig. 4.8b, the production rates of CH$_4$ are one order of magnitude lower than those of CO. Ag/TiO$_2$-SP samples with a silver concentration in the range of 2-8% had the highest and very close CH$_4$ production rates at approximately 10 mmol/h/g, more than twice as much as the average rates of WI samples in the range of 2-8% Ag. The production rate of CO was generally one order of magnitude higher than that of CH$_4$, as shown in Fig. 4.8. The selectivity of CO and CH$_4$ from CO$_2$ photoreduction, however, has not been well understood in the literature. It has been reported in our previous work $^5$ that pure TiO$_2$ favors CO production from CO$_2$ photoreduction with water vapor, while the selectivity of CH$_4$ increases if copper species are deposited on TiO$_2$ surface. The results in this work on Ag-deposited TiO$_2$ demonstrate the same trend that Ag species promote CH$_4$ production rate. However, CO is the major product (>90% selectivity) in both this work and our previous work. $^5$ The reasons that the production rate of CH$_4$ is much smaller than CO may be: (1) it is more difficult to proceed an eight-electron transfer to generate CH$_4$ compared with a two-electron process for CO, and (2) CH$_4$ may be mainly derived from further reduction of CO and thus CH$_4$ formation rate is limited by that of CO. The deposited metal species on TiO$_2$ facilitate electron trapping and transport to the catalyst surface, and thus, can potentially enhance multi-electron transfer processes, leading to a higher rate of CH$_4$. 
Figure 4.8 Rates of photocatalytic CO₂ reduction to CO (a) and CH₄ (b) for Ag/TiO₂-SP and Ag/TiO₂-WI samples at different Ag concentration (the reaction gas was a mixture of 97.2% CO₂, 2.2% H₂O and 0.6% CH₃OH).

The CO₂ reduction results again confirm that SP samples are advantageous over WI samples and 2% Ag/TiO₂-SP seems to have the highest overall activity in terms of both H₂ production and CO₂ reduction. By correlating to the material property data, the higher activities of SP samples compared with WI samples are mainly ascribed to the larger
specific surface area (Table 4.1) and better dispersion of the Ag nanoparticles (Fig. 4.5). It is well known that the Schottky barrier at the metal-semiconductor (Ag-TiO$_2$) interface facilitates electron transfer from the semiconductor to the metal, and consequently enhancing electron-hole separation and improving photocatalytic activity. It should be noted that the actual amount of Ag on SP samples was only approximately half of that on WI samples (Table 4.1). Hence, it is reasonable to also compare the activity of 2% Ag/TiO$_2$-SP with 1% Ag/TiO$_2$-WI where the two samples have similar actual Ag concentration (~1%). Again, the 2% Ag/TiO$_2$-SP sample is much more active than 1% Ag/TiO$_2$-WI with regard to both H$_2$ production and CO$_2$ reduction.

**Table 4.2** Production rates and product selectivity of the 2% Ag/TiO$_2$-SP sample under simulated solar illumination.

<table>
<thead>
<tr>
<th>Reactant gas components and their volume fractions in percentage</th>
<th>Production Rate (μmol/h/g)</th>
<th>Selectivity (%)</th>
<th>H$_2$/CO ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>H$_2$</td>
<td>CO</td>
<td>CH$_4$</td>
</tr>
<tr>
<td>Case1: H$_2$O/CH$_3$OH=2.2/0.6 $^a$</td>
<td>1130</td>
<td>18</td>
<td>0.8</td>
</tr>
<tr>
<td>Case2: CO$_2$/H$_2$O/CH$_3$OH=10/2.2/0.6 $^a$</td>
<td>439</td>
<td>43</td>
<td>0.3</td>
</tr>
<tr>
<td>Case3: CO$_2$/H$_2$O/CH$_3$OH=50/2.2/0.6 $^a$</td>
<td>321</td>
<td>78</td>
<td>0.7</td>
</tr>
<tr>
<td>Case4: CO$_2$/H$_2$O/CH$_3$OH=97.2/2.2/0.6 $^a$</td>
<td>220</td>
<td>103</td>
<td>12</td>
</tr>
<tr>
<td>Case5: CO$_2$/H$_2$O/CH$_3$OH=96.6/2.2/1.2 $^a$</td>
<td>429</td>
<td>100</td>
<td>2</td>
</tr>
</tbody>
</table>

$^a$ The reactant was balanced with N$_2$.

As described in the Experimental Section that it is not accurate to measure both H$_2$ and CO in a single photo-activity experiment because of the very low sensitivity of H$_2$ measurement when He was used as the GC carrier gas to measure CO. Hence, separate experiments were conducted (by using N$_2$ as the GC carrier gas) to measure the potential H$_2$ production by the same catalyst 2% Ag/TiO$_2$-SP using a reaction gas mixture of CO$_2$/H$_2$O/CH$_3$OH. By combining the two sets of experiments of H$_2$ and CO measurements,
the feasibility of syngas (H₂ + CO) production can be evaluated. The CO₂ partial pressure in the reaction gas was varied from 0% to nearly 100% to examine the potential influence on H₂/CO ratio in the products. The experimental results are summarized in Table 4.2, and related chemical reactions involved in the photocatalytic syngas production are listed in Reactions (1)-(6).

\[
TiO_2 \xrightarrow{hv} e_{cb}^+ + h_{ib}^+ \tag{1}
\]

\[
2H_2O + 4h^+ \rightarrow 4H^+ + O_2 \tag{2}
\]

\[
CH_3OH + h^+ \rightarrow CH_3O \bullet + H^+ \xrightarrow{\text{multi-step-decomp}} CO + nH^+ \tag{3}
\]

\[
2H^+ + 2e^- \rightarrow H_2 \tag{4}
\]

\[
CO_2 + 2H^+ + 2e^- \rightarrow CO + H_2O \tag{5}
\]

\[
CO_2 + 8H^+ + 8e^- \rightarrow CH_4 + 2H_2O \tag{6}
\]

As shown in Table 4.2, for Case 1 when H₂O + CH₃OH was the reactant gas in the absence of CO₂, the majority of final products is H₂ (rate = 1130 mmol/h/g; selectivity = 98.3%), with minor CO (rate = 18 mmol/h/g; selectivity = 1.6%) and negligible CH₄. The result indicates that direct decomposition of CH₃OH to H₂ and CO is not likely because the CO selectivity was far less than the stoichiometry where CO/H₂ should be 1:2 for complete decomposition. The minor CO produced may be from multi-step decomposition process where protons are released in each step while photo-induced holes are scavenged (Reaction (3)). In Cases 2 to 4, CO₂ + H₂O+ CH₃OH mixture were used as the reactant gas where the CO₂ partial pressure varied at three levels (10%, 50% and 97.2%), while the H₂O and CH₃OH partial pressure maintained constant at 2.2% and 0.6%, respectively. Compared to Case 1, the production rate of H₂ in the presence of CO₂ (Cases 2-4) decreased while that
of CO significantly increased and that of CH₄ remained negligible. The result clearly indicates that H₂ production (Reaction (4)) and CO₂ reduction (Reactions (5 and 6)) are competing with each other for consumption of electrons and protons, in agreement with the scheme plotted in Fig. 1.5. This competition mechanism is also supported by the result that a higher CO₂ partial pressure resulted in a higher CO production rate but a lower H₂ production rate (Cases 2-4). Accordingly, the H₂/CO ratio in the products varied from about 10 to 2, indicating that the ratio is controllable by varying the CO₂ partial pressure or the CO₂/H₂O ratio in the reactants. Another way to control the H₂/CO ratio is to vary the concentration of hole scavenger, CH₃OH, in the reaction gas. By doubling the CH₃OH volume fraction from 0.6% to 1.2% while maintaining the same volume fractions of CO₂ and H₂O (comparing Cases 4 and 5), the H₂/CO ratio also doubled from 2.1 to 4.3, due to the reason that more protons are made available by CH₃OH to favor H₂ production.

It should be noted that the produced CO was mainly derived from CO₂ (Reaction (5)) and not from CH₃OH (Reaction (3)), which is supported by the following evidences. First, in the absence of CO₂, the selectivity of CO production is only 1.6% (Case 1), indicating that generation of CO from direct decomposition of CH₃OH is insignificant. Second, in the presence of CO₂, the selectivity of CO production proportionally increased with the CO₂ concentration in the reaction gas (Cases 2-4), indicating that CO production is directly correlated with CO₂. Third, at the same CO₂ concentration, increasing the CH₃OH concentration enhanced H₂ production rate but the CO production rate remained at the same level (Cases 4 and 5). This suggests that CO production rate is limited by CO₂ and not by CH₃OH. If CO were derived from CH₃OH, an increased CH₃OH concentration could have led to a higher CO production rate. All the above experimental evidences
strongly indicate that CO is indeed a product from CO\textsubscript{2} and not from the hole scavenger, CH\textsubscript{3}OH. The production of H\textsubscript{2}, however, could come from both H\textsubscript{2}O and CH\textsubscript{3}OH.

4.4 Conclusion

This work has demonstrated the feasibility of syngas (H\textsubscript{2} and CO) production from a gas mixture of CO\textsubscript{2}, H\textsubscript{2}O and CH\textsubscript{3}OH through a photocatalytic reduction process on Ag/TiO\textsubscript{2} nanocomposite catalysts under solar irradiation. Although the H\textsubscript{2} production and CO\textsubscript{2} reduction processes compete for electrons and protons in the photocatalytic process, the final product selectivity (H\textsubscript{2}/CO ratio as low as 2, with negligible CH\textsubscript{4}) is in the appropriate range for subsequent Fischer-Tropsch process for liquid hydrocarbon fuel production. The selectivity is tunable by varying the reaction gas composition and the H\textsubscript{2}/CO ratio in the range from 2 to 10 has been achieved. In addition, the material property analysis and photocatalytic activity results showed that the ultrasonic spray pyrolysis method is much superior to conventional wet impregnation process with the advantages of smaller Ag nanoparticles, a better Ag dispersion on TiO\textsubscript{2}, and a higher fraction of metallic Ag species, which facilitate charge transfer and improve photocatalytic activity. The rates of photocatalytic conversion in this work were among the highest ones reported in the literature.
CHAPTER 5  SYNTHESIS OF NOVEL MGAL LAYERED DOUBLE OXIDES GRAFTED TiO\textsubscript{2} CUBOIDS AND THEIR PHOTOCATALYTIC ACTIVITY ON CO\textsubscript{2} REDUCTION WITH WATER VAPOR

5.1 Introduction

As mentioned in Chapter 1, photocatalytic CO\textsubscript{2} reduction with water using TiO\textsubscript{2} as the photocatalyst typically has low energy conversion efficiency\textsuperscript{109-112}. This is because of the major obstacles including the fast recombination rate of photo-generated electron-hole (e-h) pairs, the wide band gap of TiO\textsubscript{2} (3.2 eV for anatase), and the fast backward reactions\textsuperscript{11-12}. Approaches such as loading noble metal or metal oxides and doping with non-metal elements have been applied to improve the CO\textsubscript{2} photoreduction activity of TiO\textsubscript{2}\textsuperscript{113-115}. In addition to these well-known challenges, there are other factors that hinder the photocatalytic activity but are rarely studied in the literature, such as the weakened CO\textsubscript{2} adsorption on TiO\textsubscript{2} at the solid-gas interface in the presence of water vapor and limited desorption of reaction products or intermediates from the catalyst surface\textsuperscript{116-117}. As a result, to enhance the CO\textsubscript{2} adsorption on the photocatalyst, a prior step to photoreduction, is important to improve the CO\textsubscript{2} photoreduction efficiency to hydrocarbon fuels.

Our previous studies have reported MgO-TiO\textsubscript{2} composites as hybrid adsorbent/photocatalyst materials for enhanced CO\textsubscript{2} photoreduction\textsuperscript{116-117}. MgO was chosen as the CO\textsubscript{2} adsorbent because of its good CO\textsubscript{2} adsorption capability that is boosted in the presence of H\textsubscript{2}O vapor\textsuperscript{118-120}. We found that MgO-TiO\textsubscript{2} possessed much higher activity and more stable performance than pristine TiO\textsubscript{2}, particularly at a medium...
temperature range (around 150 ºC), which may be due to the easier desorption of reaction intermediates and the enhanced CO₂ adsorption by MgO at a higher temperature. We also reported that the concentration and dispersion of MgO on the MgO/TiO₂ composite strongly influenced the CO₂ photoreduction activity and 5% MgO was the optimum loading on TiO₂ surface. However, since both MgO and TiO₂ were in the form of nanoparticles in the MgO/TiO₂ composites, it was difficult to distinguish the two components by microscopic analysis and to correlate the catalytic activity with materials morphology or structure. In addition, MgO has been reported to have a relatively low kinetics in CO₂ adsorption.¹²¹-¹²² Hence, other medium temperature CO₂ sorbents with different morphologies and faster adsorption kinetics may be of greater interest to serve as the adsorbent component of the hybrid material for enhanced CO₂ photoreduction.

Layered double hydroxides (LDHs), also known as hydrotalcite-like compounds, and their post-calcination product, layered double oxides (LDOs) have been investigated as CO₂ sorbent.¹²³-¹²⁵ LDHs can be chemically expressed by a general formula $M_{1-x}^{2+}M_x^{3+}(OH)_2A_{n} \cdot nH_2O$, where $M^{2+}$ and $M^{3+}$ present metal ions and $A$ stands for exchangeable anions (Cl⁻, NO₃⁻, CO₃²⁻, SO₄²⁻).¹²⁶-¹²⁹ The characteristics of LDHs provide the LDOs properties with a large number of Brønsted basic sites, and thus LDOs is considered as a promising candidate for CO₂ adsorption,¹³⁰ and activated MgAl-LDOs are found to have a high sorption capacity for CO₂.¹²³,¹²⁵,¹³¹ It is reported that MgAl-LDOs can capture much more CO₂ than MgO at the medium temperature range 150-450 ºC.¹³²-¹³³ Hence, MgAl-LDOs could be a good candidate in replacement of MgO to combine with TiO₂ photocatalyst to improve CO₂ photoreduction operated at a medium temperature (~150 ºC).
The objective of this work is to design a novel hybrid adsorbent/photocatalyst material by grafting MgAl-LDOs (as the CO₂ adsorbent component) onto the surface of micrometer size TiO₂ cuboids (as the photocatalyst component) and to investigate the catalytic activity of CO₂ photoreduction in correlation with the materials properties. To obtain such a hybrid material structure, it is desirable to have micrometer size TiO₂ for MgAl-LDOs to be grafted on because MgAl-LDOs are reported to be micrometer or sub-micrometer size platelets. The platelet-shape MgAl-LDOs can be distinct from the micrometer-size TiO₂ cuboids, and thus, the morphology and concentration of the two components can be easily manipulated.

5.2 Materials and Methods

The micron size TiO₂ were prepared via a hydrothermal method that has been described in the literature. Typically, 1.0 g TiO₂ (Aeroxide P90) was dispersed in 60 ml 10 M NaOH aqueous solution with 1:1 H₂O/ethanol volume ratio. The solution was kept stirring for 30 min and then transferred to a Teflon-lined autoclave. The TiO₂ sample obtained after 16 h hydrothermal at 180 ˚C was washed with 0.1 M HCl aqueous solution until pH equal to 3-4 and then washed with deionized (DI) water until pH 7. MgAl-LDH grafted TiO₂ cuboids were prepared by a coprecipitation method. The as-prepared TiO₂ was re-dispersed in 30 ml 3.0 M urea solution to form solution A. Mg(NO₃)₂•6H₂O and Al(NO₃)₃•9H₂O were dissolved in 30 ml DI water with [Mg²⁺]+[Al³⁺]=0.15 M, n(Mg)/n(Al)=2:1 to obtain solution B. Solution B was dropwise added into solution A under stirring. The mixture was transferred to a round bottom flask in an oil bath. The flask was equipped with a water condenser with cooling water. The temperature for the mixture was set to be ~95 ˚C and was kept under continuous magnetic stirring for 12 hours. The
obtained precipitate was washed with DI water until pH 7 and vacuum dried at 80 °C overnight to form MgAl-LDH grafted TiO₂. Finally, the powder was calcined at 400 °C for 3 hours to form MgAl-LDO grafted TiO₂. To find the optimum MgAl-LDO concentration in the composites to obtain the best photocatalytic CO₂ reduction activity, the samples with different mass ratio of MgAl-LDO to TiO₂ were prepared. The samples were denoted as x% MgAl-LDO/TiO₂, in which x represents the nominal weight percentage of MgAl-LDO in the sample. To obtain pristine TiO₂ cuboids as a control, the MgAl-LDO/TiO₂ sample was washed with 60 °C 1 M HCl aqueous solution to remove the MgAl-LDO component.

The crystal structures of the MgAl-LDO/TiO₂ samples were identified by X-ray diffraction (XRD, Scintag XDS 2000) using Cu Kα irradiation at 45 kV, and a diffracted beam monochromator at 40 mA. The optical properties were examined by UV-vis diffuse reflectance spectroscopy using a UV-vis spectrometer (Ocean Optics) with BaSO₄ as the background. Scanning electron microscopy (SEM) (Hitachi S4800) was used to investigate the catalyst morphology. The dispersion of elements (Mg, Al, Ti, O) on MgAl-LDO/TiO₂ was analyzed by X-ray elemental mapping. Samples (15 mg) were examined under an air flow rate of 100 ml/min at a scan rate of 10 °C min⁻¹ between room temperature and 600 °C.

The photocatalytic reduction of CO₂ with H₂O vapor was conducted using a home-made quartz tube photoreactor operating in a continuous flow mode, as shown in Fig. 5.1. For each test, 100 mg catalyst was used and evenly dispersed onto a rectangular glass-fiber filter that was placed alongside the wall of the quartz tube and facing the UV light illumination. Cylinder gas CO₂ (99.999%, Praxair) continuously passed through a DI water bubbler to bring a gas mixture of CO₂ + H₂O (with 2.3 vol.% H₂O) into the photoreactor.
A high gas flow rate was used initially to purge out air inside the reactor and then the flow rate was lowered and maintained at 2.0 ml min\(^{-1}\) during photoreaction. A home-made 100 W UV lamp was used as the UV light source, and the light intensity is about 10 mW cm\(^{-2}\) in the UV range (\(\lambda < 390\) nm). To reach and maintain a reaction temperature at 150 °C, the optimum temperature found in our previous work for MgO–TiO\(_2\) catalysts, a 250 W infrared lamp was used to heat up the photoreactor and the temperature can be adjusted by varying the distance between the IR lamp and the reactor. The gaseous products in the reactor effluent were continuously analyzed with a 15 min interval using a gas chromatograph (GC, Agilent 7890A) equipped with an automated gas valve and a thermal conductivity detector (TCD) and flame ionization detector (FID).

![Experimental setup for photocatalytic CO\(_2\) reduction with water vapor](image)

**Figure 5.1** Experimental setup for photocatalytic CO\(_2\) reduction with water vapor

### 5.3 Results and Discussions

The crystal structures of TiO\(_2\) cuboids, MgAl-LDH, MgAl-LDOs, MgAl-LDH/TiO\(_2\) and MgAl-LDO/TiO\(_2\) composites were all characterized by XRD. **Fig. 5.2** shows the XRD patterns for uncalcined MgAl-LDH and 10% MgAl-LDH/TiO\(_2\) samples.
The diffraction peaks in both patterns were indexed to MgAl-LDH\textsuperscript{123, 127, 134} and weaker peaks were found for the MgAl-LDH/TiO\textsubscript{2} sample. No TiO\textsubscript{2} characteristic peaks were observed on the XRD patterns indicating that TiO\textsubscript{2} crystal phase was not formed before calcination. **Fig. 5.3** shows the XRD patterns of the samples after calcination, i.e., TiO\textsubscript{2} cuboids, MgAl-LDOs, and MgAl-LDO/TiO\textsubscript{2} composites. For TiO\textsubscript{2} cuboids, all the diffraction peaks were attributed to the anatase phase (JCPDS No. 21-1272).\textsuperscript{110, 112, 135} For the MgAl-LDOs sample, all the diffraction peaks were indexed to MgO.\textsuperscript{12, 19, 32} For MgAl-LDO/TiO\textsubscript{2} samples, as the MgAl-LDO concentration increases from 8 to 12 wt\% (or TiO\textsubscript{2} concentration decreases from 53 to 34 wt\%), both TiO\textsubscript{2} anatase diffraction peaks and MgO diffraction peaks can be observed while MgO peaks intensity increases. The Scherrer equation was applied to calculate the crystal size of TiO\textsubscript{2}. The diffraction peaks of TiO\textsubscript{2} anatase in MgAl-LDO/TiO\textsubscript{2} composites are narrowed compared with TiO\textsubscript{2} cuboids, suggesting bigger crystal size of TiO\textsubscript{2} due to MgAl-LDH addition. The average crystal size of TiO\textsubscript{2} anatase in TiO\textsubscript{2} cuboids was 15 nm. By comparison, the average crystal sizes of TiO\textsubscript{2} anatase in MgAl-LDO/TiO\textsubscript{2} composites were 26 nm, 32 nm and 40 nm for 8% MgAl-LDO/TiO\textsubscript{2}, 10% MgAl-LDO/TiO\textsubscript{2} and 12% MgAl-LDO/TiO\textsubscript{2}, respectively. Obviously, the crystal size of TiO\textsubscript{2} anatase in the MgAl-LDO/TiO\textsubscript{2} composites increases as the MgAl-LDO content increases or TiO\textsubscript{2} content decreases. The possible reason is highly crystalline LDH phase was formed as MgAl concentration increased that inhibits TiO\textsubscript{2} crystalline growth. This result was similar with MgAl-LDO/CeO\textsubscript{2} result reported by Valente et.al.\textsuperscript{136} that CeO\textsubscript{2} crystal size decreased as MgAl-LDO increased.
Figure 5.2 XRD patterns of the uncalcined samples: (a) MgAl-LDH and (b) 10% MgAl-LDH/TiO$_2$ (uncalcined samples; * represents MgAl-LDH characteristic peaks)
Figure 5.3 XRD patterns of the calcined TiO\(_2\), MgAl-LDO, and MgAl-LDO/TiO\(_2\) with different TiO\(_2\) concentrations: (a) TiO\(_2\) cuboids; (b) 8\% MgAl-LDO/TiO\(_2\); (c) 10\% MgAl-LDO/TiO\(_2\); (d) 12\% MgAl-LDO/TiO\(_2\); and (e) MgAl-LDO.

Fig. 5.4 shows the SEM images of MgAl-LDOs and TiO\(_2\) cuboids. MgAl-LDOs consists of platelets around 2 \(\mu\)m in size. The length of the TiO\(_2\) cuboids is in micrometer size range with a high aspect ratio. Fig. 5.5 shows the SEM images of MgAl-LDO/TiO\(_2\) samples with different compositions. The low magnification SEM images in Fig. 5.5 a-c show that the three samples were composed of MgAl-LDOs grafted on micrometer size TiO\(_2\) cuboids. The platelet shape of MgAl-LDOs on the composites after calcination was almost the same as the as-prepared uncalcined composites MgAl-LDHs. In most of literature reported, MgAl-LDHs platelets morphology cannot be maintained after calcination.\(^{129, 137}\) MgAl-LDOs platelets with a highly ordered layered array are proved
beneficial for the application as adsorbent and catalyst.\textsuperscript{138-139} The high magnification SEM images in Fig. 5.5 d-f show both the size and the coverage of MgAl-LDOs grafted on TiO\textsubscript{2} cuboids increased as the MgAl-LDO ratio increased. The distributions of Mg, Al, Ti and O elements in the MgAl-LDO/TiO\textsubscript{2} composites were analyzed by X-ray elemental mapping, and the results are shown in Fig. 5.6. The elemental mapping images demonstrated that the cuboids skeleton were mainly composed of Ti element and the grafted platelets were composed of Mg and Al elements that were evenly distributed on the cuboids surface.

**Figure 5.4** SEM images of (a) MgAl-LDOs and (b) TiO\textsubscript{2} cuboids
Figure 5.5 SEM images of the calcined MgAl-LDO/TiO$_2$ composites: (a,d) 8% MgAl-LDO/TiO$_2$, (b,e) 10% MgAl-LDO/TiO$_2$, and (c,f) 12% MgAl-LDO/TiO$_2$.

Figure 5.6 X-ray elemental mapping images of the MgAl-LDO/TiO$_2$ composites: (a) 8% MgAl-LDO/TiO$_2$, (b) 10% MgAl-LDO/TiO$_2$, and (c) 12% MgAl-LDO/TiO$_2$.

Diffuse reflectance UV-vis spectra were recorded to explore the influence of MgAl-LDOs addition on the optical property of TiO$_2$. As shown in Fig. 5.7, TiO$_2$ cuboids absorbs light at wavelength shorter than 400 nm, corresponding to a band gap of about 3.0 eV. MgAl-LDOs show no adsorption ability of light at the wavelength range between 300-600 nm. Incorporation of MgAl-LDOs in TiO$_2$ cuboids leads to a blue-shift in the adsorption
Photocatalytic activity tests were carried out under UV light irradiation at 50 °C for 4 h and subsequently at 150 °C for another 8 h, and the results are shown in Fig. 5.8. CO was found to be the major product with a minor CH₄ concentration that was one or two orders of magnitude lower than CO. Thus, only CO production results were shown in Fig. 5.8 to compare the activities of the different catalysts. Fig. 5.8a shows bare TiO₂ cuboids had a CO production rate around 1.0 μmol g⁻¹ h⁻¹ at 50 °C while pure MgAl-LDO had no activity for CO production. MgAl-LDOs addition on TiO₂ cuboids had no significant improvement for CO production rate. The possible reason is MgAl platelets partially or fully covered TiO₂ cuboids which blocks CO₂ contact with TiO₂. Moreover, MgAl-LDOs has weak adsorption ability of CO₂ at mild room temperature that also leads to poor CO₂ contacting with TiO₂.¹³⁰
For the proceeding 8 hours test under UV light at 150 °C, CO production rate for TiO₂ cuboids was almost the same as the first 4 hours at 50 °C. Pure MgAl-LDOs still showed no activity of CO₂ reduction. Interestingly, 10% MgAl-LDO/TiO₂ exhibited an obvious enhancement of CO production rate with a maximum 4.8 μmol g⁻¹ h⁻¹, 2.5 times higher than that under 50 °C. For MgAl-LDO/TiO₂ samples, the activity of CO₂ photoreduction was in the order of 10% MgAl-LDO/TiO₂ > 8% MgAl-LDO/TiO₂ > 12% MgAl-LDO/TiO₂. The optimum concentration of MgAl-LDOs on the MgAl-LDO/TiO₂ was 10%. 12% MgAl-LDO/TiO₂ performed the lowest CO production rate amongst the MgAl-LDO/TiO₂ composites. Even though, it showed a similar CO production rate as TiO₂ cuboids under 150 °C. This indicates that MgAl-LDOs grafting on TiO₂ helps improve the photoactivity of CO₂ reduction. It is possibly because MgAl-LDOs absorbs more CO₂ at higher temperature (150 °C) onto the composites which creates more opportunity for CO₂ to be photoreduced to CO. On the other hand, as MgAl-LDOs concentration decreased to 8%, CO production rate is between that of 12% TiO₂-MgAl-LDOs and 10% TiO₂-MgAl-LDOs but still much lower than that of 10% TiO₂-MgAl-LDOs. This may be explained in terms of increased crystal size of the anatase TiO₂ nanoparticles. Another reason could possibly be that MgAl-LDOs platelets covered the majority parts of TiO₂ cuboids surface either blocked the CO₂ and H₂O vapor contacting with TiO₂ or electrons transferring from TiO₂ to CO₂ and H₂O molecules.

To better understand the carbon source of CO, further photocatalytic activity tests were carried out by changing the CO₂+H₂O vapor atmosphere to Helium (He) + H₂O vapor atmosphere. As shown in Fig. 5.8b, the CO production rate were compared on 10% MgAl-LDO/TiO₂ under both conditions. Under UV irradiation, there was no CO production in
He atmosphere, indicating CO$_3^{2-}$ in MgAl LDOs structure was stable under mild room temperature. Compared with CO$_2$ atmosphere, CO was produced on 10% MgAl-LDO/TiO$_2$ in the CO$_2$ under UV irradiation, which came from CO$_2$ photoreduction on this composite. After introducing IR irradiation (150 °C), CO production was shown in He atmosphere with a maximum rate of 1.3 μmol g$^{-1}$ h$^{-1}$. The experiments was carried out several times to rule out other impurities that might produce CO. The results indicated that most of CO was attributed to CO$_2$ photoreduction and a small portion of CO was from CO$_3^{2-}$ in MgAl LDOs.

![Graph](image1)

**Figure 5.8** The rate of CO production from CO$_2$ photoreduction under UV light irradiation at 50 °C for 4 h and subsequently at 150 °C for 8 h using (a) TiO$_2$ cuboids, MgAl-LDO and three MgAl-LDO/TiO$_2$ composites in CO$_2$ + H$_2$O vapor atmosphere, and (b) 10% MgAl-LDO/TiO$_2$

### 5.4 Conclusion

This work has demonstrated that the successful synthesis for novel hybrid structure of MgAl-LDOs platelets grafted on TiO$_2$ cuboids with high length to width ratio. MgAl-LDOs addition has no impact on the optical propertied of TiO$_2$. The morphology of TiO$_2$-MgAl-LDOs showed that the surface coverage on TiO$_2$ on MgAl-LDOs platelets could be controlled by tuning the MgAl-LDOs to TiO$_2$ ratio. As the MgAl-LDOs concentration increases, TiO$_2$ crystal size was increased. MgAl-LDOs grafting on TiO$_2$ cuboids may help
improve the adsorption ability of CO\textsubscript{2} onto TiO\textsubscript{2} which improves the photocatalytic activity of CO\textsubscript{2} reduction. The optimum MgAl-LDOs concentration was found to be 10\% which performed the highest CO production rate. The proposed reason for this is: at higher MgAl-LDOs concentration, MgAl-LDOs platelets covered the majority parts of TiO\textsubscript{2} cuboids surface either blocked the CO\textsubscript{2} and H\textsubscript{2}O vapor contacting with TiO\textsubscript{2} or electrons transferring from TiO\textsubscript{2} to CO\textsubscript{2} and H\textsubscript{2}O molecules; Moreover, the crystal size increase of TiO\textsubscript{2} anatase may also lead to a poor CO\textsubscript{2} photoreduction activity. At lower MgAl-LDOs concentration, less CO\textsubscript{2} and H\textsubscript{2}O amount may be adsorbed onto the composites which also leads to lower CO production rate.
CHAPTER 6 FACILE SYNTHESIS OF CARBON-TIO$_2$ NANOCOMPOSITES WITH ENHANCED REVERSIBLE CAPACITY AND CYCLIC PERFORMANCE AS ANODES FOR LITHIUM-ION BATTERIES

6.1 Introduction

As mentioned in Chapter 1, Graphitic carbons are most commonly used as conventional commercial anode materials owing to its low cost, relatively high energy density and very low flat working potential. However, graphitic carbon anode materials suffer the disadvantages of severe safety issues because of its low operating voltage (~0.1 V vs Li/Li$^+$), poor life cycle performance due to solid-electrolyte interface (SEI) film and the formation of dendritic lithium on the surface of graphitic carbons anode.$^{141-144}$

Thus, it is essential to make improvements such as safety, cost, high efficiency and long cycle life of the anode materials for LIBs. Researchers in the scientific and industrial communities have investigated different materials such as silicon, tin (II) based oxide$^{145-146}$ and TiO$_2$ as LIBs anode to replace graphite materials in order to eliminate the safety issue and meanwhile improve LIBs performance. Different from tin (II) based oxides and silicon, TiO$_2$ material is structural stable during lithium insertion and extraction which means a very low volume change ($<4\%$ for Li$_x$TiO$_2$, $0\leq x \leq 1$) for electrode.$^{147-148}$ This advantage guarantees its good stability during charge/discharge process at high current density rate. Moreover, its high working voltage ($> 1.5$ V vs Li/Li$^+$) avoids reactions with organic electrolyte and short circuit caused by electrode expansion during cycling that
make it safer than graphite anode material.\textsuperscript{149-153} TiO\textsubscript{2} offers a theoretical capacity of 335 mAh g\textsuperscript{-1} based on the complete reduction of Ti\textsuperscript{4+} to Ti\textsuperscript{3+}, which is comparable to that of conventional graphite anode materials.\textsuperscript{154-156} Being abundant, low cost and environmentally benign,\textsuperscript{109, 111, 143, 156} TiO\textsubscript{2} has been considered as a promising candidate as LIBs anode material.

It is well known that TiO\textsubscript{2} exists in three different polymorphs in nature: anatase, brookite and rutile, among which, it was reported that brookite and rutile phase TiO\textsubscript{2} can only accommodate limited small amount of Li\textsuperscript{+} ions, while anatase phase has superior accommodation capacity of lithium insertion.\textsuperscript{157-158} However, TiO\textsubscript{2} anodes practical capacity is only 168 mAh g\textsuperscript{-1} because of the poor electronic and ionic conductivity in its bulk form, limiting TiO\textsubscript{2} anode practical applications.\textsuperscript{143-144, 151, 154, 156} Two general approaches are widely used to improve the electronic and ionic conductivity associated with Li\textsuperscript{+} ions in the active materials of the anode. One approach is tailoring the bulk state TiO\textsubscript{2} to nanoscale dimensions to reduce Li\textsuperscript{+} ions diffusion length and increase the energy storage capacity.\textsuperscript{143, 159} The other approach is blending TiO\textsubscript{2} with conductive material such as conductive carbon to improve its electronic conductivity.\textsuperscript{144, 160} Various materials based on the above two aspects consideration have been investigated and exhibited relatively superior performance as anode for LIBs, including 3D TiO\textsubscript{2}-mesoporous carbon nanocomposite,\textsuperscript{143} TiO\textsubscript{2}-C composites nanotubes,\textsuperscript{144} ordered mesoporous C-TiO\textsubscript{2},\textsuperscript{147} TiO\textsubscript{2}@C composites nanofibers,\textsuperscript{160} etc. TiO\textsubscript{2} in these composites are commonly synthesized via hydrothermal, template and other post-treatment procedures. Specifically, hydrothermal method usually requires a relatively long reaction time at high temperature and pressure condition. The template method for TiO\textsubscript{2} synthesis takes further steps to
remove the template, which makes the method unfavorable for practical application. Furthermore, none of the electrode materials exhibited capacities higher than 300 mA h g\(^{-1}\) between the potentials of 1.0 V to 3.0 V.

Herein, we report carbon decorated TiO\(_2\) nanocomposites (C-TiO\(_2\)) synthesized via a facile carbonization process as an excellent candidate for anode materials in LIBs. The fabrication process is the first time reported for carbon decorated directly on commercial TiO\(_2\) materials under relative mild temperature condition, with special focus on the carbon growth control on TiO\(_2\) and application as LIBs anode material. Conductive carbon material directly decorated on TiO\(_2\) particles without any post-treatment processes. In this trial, the objective is to generate a synthesis process for C-TiO\(_2\) used as anode material for LIBs which is cost-effective and easy to be applied in commercial system. The starting materials used for preparation are commercial TiO\(_2\) products (P25 and P90), which are easily available and inexpensive. Sucrose is also used to form the carbon on TiO\(_2\) being carbonized with very little amount of sulfuric acid. The preparation process is quite simple and easy to be scaled up. This nanocomposites structure offers advantages such as: (1) an improved Li\(^+\) ion transport due to the size of the individual TiO\(_2\) anatase nanoparticles, (2) an improvement in the electronic conductivity achieved via amorphous carbon coating. The battery performance results demonstrate that C-TiO\(_2\) nanocomposites exhibit significantly enhanced capacity and rate capability as compared to bare TiO\(_2\) nanoparticles when used as LIBs anode material.

### 6.2 Experimental Method and Characterization

C-TiO\(_2\) were prepared via three simple steps (Fig. 6.1). Firstly, sucrose in sulfuric acid solution was mixed with TiO\(_2\) (Evonik P25 or P90) with a designated C:TiO\(_2\) weight
ratio. After thoroughly mixing, the mixture was transferred to a round-bottom flask and heated up to 90 °C in an oil bath under vigorous stirring. The round-bottom flask was equipped with cooling water reflux condenser to condense the vapor and ensure a constant solution amount in the flask. Secondly, the obtained brown color precipitate after 24 hours sucrose dehydration was washed until its pH value to 7. The resultant mixture was dried at 80 °C for 12 hours. Finally, the as-dried composite was calcined in N₂ atmosphere at 450 °C for 3 hours. For comparison purpose, P25 and P90 were also treated following the same procedure of C-TiO₂ synthesis without sucrose and sulfuric acid addition. The corresponding composites were denoted as a%C-P25 and a%C-P90 (a% represents carbon weight percentage).

**Figure 6.1** Schematic diagram of decorating TiO₂ with carbon process in a round bottom flask connected to a cooling water (CW) condenser and formation of C-TiO₂ nanocomposites

The crystal structures of the products were identified by X-ray diffraction (XRD) (Scintag XDS 2000) using Cu Kα irradiation at 45 kV and a diffracted beam monochromator operated at 40 mA in the 2θ range from 20° to 80° at a scan rate of 2°.
Scanning electron microscopy (SEM) equipped with energy-dispersive X-ray (EDX) spectroscopy (Hitachi S4800) was used to analyze the nanocomposites morphology and surface elemental composition. Transmission electron microscopy (TEM, Hitachi H9000NAR) and high-resolution TEM (HRTEM, Hitachi H9000NAR) were used to examine the particle size and dispersion of amorphous carbon on TiO$_2$ as well as the lattice information. The Brunauer-Emmett-Teller (BET) specific surface area of the composites was measured by nitrogen adsorption at 77 K on a surface area and porosity analyzer (Micrometrics ASAP 2020). Before each adsorption measurement, approximate 0.10 g sample was degassed at 100 °C for 18 hours. The BET surface area was determined by a multipoint BET method using the adsorption data in a relative pressure ($P/P_0$) range 0.05-0.3. Pore size distribution was determined via Barret-Joyner-Halender (BJH) method using desorption isotherm. The N$_2$ adsorption volume at the relative pressure of 0.998 was used to determine the total pore volume and average pore size. The thermal stability of electrode materials were carried out on a thermalgravimetric analyzer (TGA-DAT-2960 SDT) at a heating rate of 20 °C min$^{-1}$ from 25 to 1000 °C in air.

80 wt.% active material (C-TiO$_2$ or TiO$_2$) powder with 10 wt.% carbon black and 10 wt.% Poly(vinylidene fluoride) (PVDF) binder was homogeneously mixed in N-methyl pyrolidinone (NMP). The slurry was well deposited by doctor blading onto a copper foil, and the as-coated copper foils were dried under vacuum at 100 °C overnight. Each electrode was carefully weighed before use and several electrodes were tested to ensure the reproducibility of the electrochemical behavior.

The working electrodes and Li metal foil counter electrode were assembled into cells by using Celgard 2400 as the separator and 1 M solution of LiPF$_6$ in ethylene
carbonate (EC) and dimethyl carbonate (DMC) (EC: DMC volume ratio 1:1) as the electrolyte. The cells were constructed in an Argon-filled glove box. The charge/discharge curves were measured using a LAND Cell tester (Wuhan, China) at a cutoff voltage of 1.0-3.0 V under current densities of 30, 60, 300 mA g\(^{-1}\). The specific capacity and current density were calculated based on TiO\(_2\) mass only. Cyclic voltammetric curves were recorded in a voltage range of 1.0-3.0 V at a scan rate of 0.2 mV s\(^{-1}\). The electrochemical impedance spectra results were recorded by applying an alternating current of 5 mV in a frequency range from 0.01 Hz to 100 kHz.

6.3 Results and Discussion

Fig. 6.2 shows the XRD patterns of TiO\(_2\) and C-TiO\(_2\) nanocomposites. The phase contents and crystal sizes of anatase and rutile were calculated and summarized in Table 6.1. P25 exhibited characteristic peaks for both anatase (JCPDS No. 21-1272) and rutile (JCPDS No. 21-1276) with 85.9% anatase and 14.1% rutile. The phase composition in P25 sample is consistent with the reported literature.\(^{161}\) The average crystal size for anatase and rutile in P25 is 18.5 and 21.3 nm, respectively, according to the Scherrer equation. In comparison, P90 exhibited prominent characteristic peaks for anatase (92.6%) and very small peaks for rutile (7.4%). The phase composition in P90 also agrees with the results reported.\(^{162-163}\) The crystal size of anatase and rutile in P90 are 11.7 and 15.9 nm, respectively, smaller than that in P25. There was no significant phase fraction change for P90 after carbon decoration, while its crystal size slightly increased. Similar phenomena were found on P25. The increased crystal size can be explained by the additional calcinations process for the carbon decorated samples.\(^{164}\) From XRD patterns of C-P90 and C-P25, no peak shift was observed compared with carbon free samples, indicating the
absence of any chemical bonding or reaction between the TiO\textsubscript{2} nanoparticles and carbon. Furthermore, there are no characteristic peaks of carbon detected indicating the formation of amorphous carbon.\textsuperscript{153}

**Table 6.1** Fractional phase content, crystal size and BET specific surface area of P25, P90, C-P90 and C-TiO\textsubscript{2} anatase samples

<table>
<thead>
<tr>
<th>Samples</th>
<th>Phase content (%)</th>
<th>Crystal size (nm)</th>
<th>S\textsubscript{BET} (m\textsuperscript{2}/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A</td>
<td>R</td>
<td>A</td>
</tr>
<tr>
<td>P25</td>
<td>85.9</td>
<td>14.1</td>
<td>18.5</td>
</tr>
<tr>
<td>P90</td>
<td>92.6</td>
<td>7.4</td>
<td>11.7</td>
</tr>
<tr>
<td>1.8% C-P25</td>
<td>87.6</td>
<td>12.4</td>
<td>18.1</td>
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<tr>
<td>1.9% C-P90</td>
<td>95.5</td>
<td>4.5</td>
<td>12.8</td>
</tr>
</tbody>
</table>

**Figure 6.2** XRD pattern of (a) P90 (b) P25 (c) 1.8\% C-P25 (d) 1.9\% C-P90 and (e) 9.5\% C-P90.
The textural properties and morphology of C-TiO₂ samples were characterized by BET, SEM, EDX-X-ray elemental mapping and HRTEM. Fig. 6.3 shows the N₂ adsorption-desorption isotherms of TiO₂ and C-TiO₂ samples. P25, P90, 1.8% C-P25 and 1.9% C-P90 samples exhibited mesoporosity in the form of typical type IV isotherms with a H3 hysteresis loop. According to IUPAC classification, mesoporous materials characterizing with the type H3 loop are commonly observed with aggregates of plate-like particles, leading to the slit-shaped pores.¹⁶⁵ Table 6.1 compares the BET surface areas for TiO₂ (P25, P90) and C-TiO₂ samples. Bare TiO₂ and 1.8%C-P25 displayed a comparable surface area, probably because a small amount of carbon were deposited on the surface of TiO₂. Notably, P90 and 1.9%C-P90 have a larger specific surface area than P25 and 1.8%C-P25, likely due to the smaller particle size of P90 as evidenced by SEM images described later in this paper. The surface area of the sample was slightly decreased for both carbon decorated TiO₂, which was reasonable due to the agglomeration during the carbonization and calcination process. It is notable that 1.9% C-P90 still possesses a higher specific surface area (96.8 m²/g) compared with 1.8% C-P25 (54.2 m²/g).
Figure 6.3 N\textsubscript{2} adsorption/desorption isotherms of P25, P90, 1.8% C-P25 and 1.9% C-P90.

SEM images for 1.9%C-P90 and 1.8%C-P25 shown in Fig. 6.4 confirmed that 1.9%C-P90 is composed of primary nanoparticles with a particle size around 20~30 nm, while 1.8% C-P25 is composed of bigger size nanoparticles around 40~50 nm. The smaller particles will tend to form higher specific surface area. This is in consistent with the BET results that 1.9% C-P90 exhibits a higher specific surface area than 1.8% C-P25 does. There are short slit-pores formed by the TiO\textsubscript{2} particles overlaying, which is in good agreement with the N\textsubscript{2} sorption isotherms results. The X-ray elemental mapping results in Fig. 6.5 suggest that the carbon species (red color) were well dispersed in the TiO\textsubscript{2} particles surface.
Figure 6.4 SEM Images of (a) 1.9% C-P90 and (b) 1.8% C-P25 (insets are different magnification images, respectively)
Figure 6.5 X-ray mapping images of 1.9% C-P90.

HRTEM was conducted on 1.9%C-P90 and 9.5%C-P90 to explore the location and thickness of carbon layers. Fig. 6.6 shows that clear lattice fringes of nanocrystal with an interplanar space of 0.35 nm were observed on 1.9% C-P90 and 9.5% C-P90, which correspond to the (101) plane of anatase TiO$_2$. The lattice fringes of rutile TiO$_2$ were not seen under the observed projections that could be explained by the small fraction amount rutile phase. Carbon existence can be evidenced by the sample color (inset picture in Fig. 6.6a). Fig. 6a exhibited a thin carbon layer on some particles of TiO$_2$ with a thickness of ~0.4 nm (pointed out with blue arrows). In Fig. 6.6a, some crystalline particles (circled with dashed red line) with no continuous layer can also be observed. It can be confirmed that this continuous edge is carbon layer on the proof of the appearance comparison of coated and uncoated crystalline particles. In contrast, Fig. 6.6b showed a thicker carbon
layer which was around 5 nm on 9.5% C-P90. As carbon concentration increases, it tends to form a layer on TiO$_2$ particles. The layer becomes thicker as the carbon content increases. No lattice fringes were observed for the carbon layer which indicated carbon species were composed of amorphous carbon.

Figure 6.6 HRTEM images of (a) 1.9% C-P90, and (b) 9.5% C-P90.

Figure 6.7 Thermogravimetry analysis (TGA) for P90 and C-P90 with various carbon concentration
The carbon content of C-P90 samples were confirmed using thermogravimetric analysis (TGA) (Fig. 6.7). The samples were heated from 25 °C to 800 °C at a rate of 10 °C min\(^{-1}\). There are two steps weight loss of C-P90 samples in the temperature range between 25 and 800 °C. The weight loss below 200 °C is due to the adsorbed water while weight loss between 200 °C and 800 °C is attributed to the decomposition of amorphous carbon. The weight loss was estimated 0.4, 1.9 and 9.5 wt% of the total weight for various C-P90 materials in the range of 200-800 °C. It is noted that the carbon concentration on TiO\(_2\) can be tuned via this simple preparation procedure.

Figure 6.8 Cycling performance of (a) pristine TiO\(_2\) (P25 and P90) and (b) C-TiO\(_2\) (1.8% C-P25 and 1.9% C-P90) nanocomposites at a current density of 30 mA g\(^{-1}\).
To investigate the effects of crystal phases and carbon deposition on lithium storage performances of TiO$_2$, two common commercial TiO$_2$ P25, TiO$_2$ P90, 1.8% C-P25 and 1.9% C-P90 s were assembled in cells and tested in the range of 1.0-3.0 V at a 30 mA g$^{-1}$ current density. **Fig. 6.8a** shows the electrical performance of pure TiO$_2$. Both P25 and P90 electrodes showed a fast capacity fading from 200 mAh/g to 50 mAh/g in the initial 30 cycles, possibly due to the intrinsically poor electronic conductivity of pristine TiO$_2$. Li$^+$ insertion/ extraction into/out of TiO$_2$ mechanism can be expressed as TiO$_2$ $+$x(Li$^+$+$e^-$)$\leftrightarrow$Li$_x$TiO$_2$, where x depends on the materials crystallography and microstructure. And x is limited to 0.5 for pure TiO$_2$ due to its poor electronic conductivity and Li$^+$ diffusivity.$^{166}$ Interestingly, depositing carbon (1.8%) on P25 induced a high initial discharge capacity of 260 mA h g$^{-1}$, but still suffers from the capacity fading problem, as shown in **Fig. 6.8b**. By contrast, 1.9% C-P90 exhibited an initial discharge capacity of 231 mA h g$^{-1}$ (0.7Li$^+$) and an excellent cyclic performance. The capacity was retained at 173 mA h g$^{-1}$ after 100 cycles (75% retention), much superior to pure TiO2 and 1.8%C-P25. The superiority of 1.9% C-P29 was probably due to two reasons. First, as the carbon concentration on both samples is almost the same, the drastic capacity decay of 1.8% C-P25 can be ascribed to the higher rutile phase fraction in the structure of TiO$_2$ which largely influence the electrochemical performance. It is reported that the diffusion coefficient of Li$^+$ ions in rutile TiO$_2$ (1.42 $\times$10$^{-13}$ cm$^2$ s$^{-1}$), which is smaller than that of anatase TiO$_2$ (1.81 $\times$10$^{-13}$ cm$^2$ s$^{-1}$).$^{167}$ This can be possibly explained that the channels of rutile TiO$_2$ are narrower than those of anatase TiO$_2$, which is proved by the superior electrochemical performance of 1.9% C-P90 due to its less rutile phase fraction and more anatase phase fraction. Second, the higher specific surface area of 1.9% C-P90 than that of 1.8% C-P25
helps to store more Li\textsuperscript{+} ions to a certain extent.\textsuperscript{168}

**Figure 6.9** Electrochemical performance of (a) Representative cyclic voltammograms (CV) of 1.9\% C-P90 sample at a scan rate of 0.2 mV s\textsuperscript{-1} between 1 V and 3 V; (b) Charge-discharge voltage profiles of the first 3 cycles for 1.9\% C-P90 at 30 mA g\textsuperscript{-1}; (c) performance of C-P90 nanocomposites at 30 mA g\textsuperscript{-1}; (d) Rate capability of C-P90 nanocomposites at difference rates;

**Fig. 6.9 a** shows representative cyclic voltammograms (CV) of 1.9\% C-P90. It is obvious that the sample demonstrates a CV curve pattern with a pair of redox peaks indicating the Li\textsuperscript{+} insertion (cathodic sweep, 1.51 V) and Li\textsuperscript{+} extraction (anodic sweep, 2.15 V) into and out of TiO\textsubscript{2} processes. The cathodic and anodic peaks from CV are in close correspondence with the voltage plateaus obtained from the galvanostatic cycling in **Fig. 6.9 b**. This result is consistent with the results reported in the literature.\textsuperscript{169} The anodic peak with almost the same current density as the cathodic peak proves that the Li\textsuperscript{+} ions
extraction takes place to the equal extent, indicating a nearly perfect Coulombic efficiency. Furthermore, the galvanostatic charge-discharge curve in Fig. 6.9 a shows no substantial change after 2nd cycles, showing the high reversibility of the Li\(^+\) insertion/extraction reactions.

Due to the superior electrochemical performance of C-P90, we further studied different C-P90 materials in order to optimize the electrochemical performance by tuning the carbon concentration. Fig. 6.9 c showed the cycling performance of various C-P90 at a current density of 30 mA g\(^{-1}\). 0.4% C-P90 exhibited a relatively lower cycling capacity, although its initial capacity is higher (255 mA h g\(^{-1}\)). The lower cycling capacity can be explained by that less carbon decoration on P90 is ineffective to provide electronic conductivity. As the carbon concentration increased from 1.9% to 9.5%, an even poor cycling performance was exhibited. Only 55 mA h g\(^{-1}\) capacity was retained after 100 cycles for 9.5% C-P90.

Fig. 6.9 d showed the cycling performance at different current density (30 mA g\(^{-1}\), 60 mA g\(^{-1}\), 300 mA g\(^{-1}\) and 30 mA g\(^{-1}\)). 1.9%C-P90 has superior rate capability comparing with 0.4%C-P90 or 9.5%C-P90 materials, which also proves the proposal reason that the optimized carbon concentration improves the electronic conductivity. 1.9%C-P90 exhibited the best cyclability and rate capability comparing with 0.4%C-P90 and 9.5%C-P90. The electrochemical performance that 1.9%C-P90 possesses is even comparable to some hierarchical carbon-TiO\(_2\) composites reported in some literature. As mentioned above, 1.9% C-P90 exhibited a high initial discharge capacity of 231 mA h g\(^{-1}\) at a current density of 30 mA g\(^{-1}\) and a capacity of 130 mA h g\(^{-1}\) at a current density of 300 mA g\(^{-1}\).
Electrochemical impedance experiments were applied to explore the effect of carbon decoration on TiO$_2$ (P90) nanoparticles on the interface impedance of C-TiO$_2$ composites. **Fig. 6.10** shows the Nyquist plots of the carbon decorated TiO$_2$ (P90) particles after 3 cycles charge-discharge process. The semicircle in the intermediate frequency is related to the charge transfer resistance at the electrode/electrolyte interface, while the sloping line at the low frequency end represents the Warburg impedance caused by a semi-infinite diffusion of Li$^+$ ion in the electrode.$^{170-172}$ It can be seen clearly that the interfacial resistance ($R_{ct}$) is much smaller for the C-P90 electrode than for the pristine P90 electrode, which indicates that carbon decoration could enable much easier charge transfer at the electrode and electrolyte interface. As a result, carbon decoration decreases the overall battery internal resistance. For various carbon concentration electrode C-P90, the
interfacial resistance is with this order $R_{ct}^{1.9\%\text{C-P90}} < R_{ct}^{0.4\%\text{C-P90}} < R_{ct}^{9.5\%\text{C-P90}}$. It shows that less carbon concentration (e.g. 0.4%) is unable to provide good conductivity, while more carbon integration (e.g. 9.5%) into TiO$_2$ can increase charge transfer resistance.\textsuperscript{173} This can explain that 1.9% C-P90 exhibits the best electrochemical performance due to the optimum carbon concentration.

Comparing with the reported results in the open literature, 1.9% C-P90 prepared with a simpler method, which exhibited a very promising electrochemical performance result. Shen, et al.\textsuperscript{143} reported a three dimensional coherent titania-mesoporous carbon nanocomposite, which was prepared by a template method. This material exhibited a reversible capacity of \(~150\text{ mA h g}^{-1}\) at a current density of 200 mA h g$^{-1}$. Zhen, et al.\textsuperscript{166} studied a TiO$_2$ on reduced graphene oxides composite, which was synthesized by a two-step hydrothermal method. This composites performed a 200 mA h g$^{-1}$ capacity at 1.2 C (1 C=167.5 mA g$^{-1}$). Liu, et al.\textsuperscript{174} prepared a sandwich-like TiO$_2$-carbon hollow sphere structure composite via two-step hydrothermal method, which exhibited a 200 mA h g$^{-1}$ capacity at 1 C. These reported materials were either synthesized by hydrothermal, template method or involved with organic precursors as carbon source. In comparison, our C-P90 sample is prepared by one-step carbonization at mild conditions with commercial materials. 1.9% C-P90 material exhibited a comparable performance mainly due to its unique carbon decoration nanostructure and anatase-rutile phase composition that facilitate efficient lithium insertion and extraction.

To illustrate the mechanism of carbon layer function on TiO$_2$ P90, Fig. 6.11 shows the scheme according to the characterization results. At lower carbon concentration (0.4%), carbon cannot form a uniform layer on TiO$_2$ P90 nanoparticles, which results in poor
conductivity. At higher carbon concentration (9.5%), it tends to form a thicker layer on TiO$_2$ P90 nanoparticles, which inhibits Li$^+$ insertion/extraction. By comparison, on the TiO$_2$ P90 with moderate carbon content materials, carbon layer provides conductivity for the composites and the thickness of which enables Li$^+$ insert into TiO$_2$.

**Figure 6.11** Schematic representation of the nanostructure and lithium ion insertion and electron transport of the C-P90 nanocomposites.

### 6.4 Conclusion

In conclusion, carbon decorated TiO$_2$ were synthesized using commercial TiO$_2$ product (P25 and P90) in large scale through a simple carbonization process. C-P90 possesses higher specific area and larger fraction of anatase comparing to C-P25 with the same carbon concentration, which guaranteed its superior electrochemical performance as anode for LIBs. Carbon concentration had been optimized to obtain the best battery performance for P90. 1.9% C-P90 showed a significant stable cyclability which was attributed to its unique carbon decoration nanostructure that facilitates efficient lithium insertion and extraction. The easy preparation together with the advantage of stable cyclability of C-P90 makes the composite a very promising anode material for LIBs.
CHAPTER 7 HIERARCHICAL SULFUR-INFILTRATED POROUS MICROSPHERES CARBON FOR HIGH ENERGY LITHIUM-SULFUR BATTERIES

7.1 Introduction

As mentioned in Chapter 1, scientists fabricated different carbon framework to constrain sulfur to obtain better performance cathode for lithium-sulfur batteries. Regardless of the various carbon frameworks, pore structure and sizes are important factors that determine the electrochemical performance. So far, only a few studies reported sulfur/carbon composites with a bi-modal or tri-modal pore structure as cathodes for Li-S batteries. Schuster et al. developed a spherical ordered bi-modal (3.1 nm and 6 nm) mesoporous carbon/sulfur nanocomposite, and they claimed that the bimodal pore distribution and nano-carbon dimensions were beneficial for more sulfur incorporation and improved electrolyte accessibility compared with single pore size structure, resulting in an initial discharge capacity at 1,200 mA h g⁻¹ and 730 mA h g⁻¹ after 100 cycles at 1 C rate. On the other hand, it has been reported that Li⁺ diffusion within an ordered porous sulfur/carbon framework may be inhibited or limited due to fully or partially filled carbon pores by sulfur. Moreover, for high-rate electrochemical performance, mesopores larger than 10 nm or even macropores are required for better electrolyte diffusion and fast ion transport.

However, no research has aimed to integrate micropores, mesopores and macropores in a single carbon framework with a controllable manner to achieve a satisfied electrochemical performance for Li-S batteries. In this work, we have designed and fabricated a novel 3D multi-modal porous microsphere carbon framework (PMC) that
integrates interconnected micropores, mesopores, and macropores with easily tunable pore size distribution. The pore distribution can be controlled by adjusting the silica nanoparticles (NPs) sizes (40 and 10 nm) and the ratio of different sized silica NPs. Unlike conventional wet-chemistry methods to prepare porous carbon, an aerosol-based spray pyrolysis method was used in this work to synthesize the multi-modal PMC (see Fig. 7.1).

![Figure 7.1 Schematic of experimental setup and process for synthesizing porous carbon/sulfur microspheres.](image)

### 7.2 Experimental Method and Characterization

Porous Microsphere Carbon (PMC): The PMC was synthesized through a spray pyrolysis method with the aid of silica templates. In brief, a mixture of 4.2 g sucrose and 20 ml 20 wt.% silica sol (Nissan Chemical) with 1:2 weight ratio of carbon/silica was first prepared. 0.25 ml 95% sulfuric acid was then added into the evenly mixed solution. The solution was then heated up to 90 °C in an oil bath under vigorous stirring. The round-
bottom flask was equipped with a cooling water reflux condenser to condense the vapor and ensure a constant solution amount in the flask. After reflux for 30 h for sucrose carbonization, the solution was then transferred to a Collison nebulizer (BGI Inc.) for spray pyrolysis through a cylindrical quartz tube placed inside a tube furnace. The tube furnace temperature was set at 600 °C during spray pyrolysis and compressed air was used as the carrier gas. Powder samples were collected on a glass fiber filter and then finally were calcined in N₂ at 800 °C for 4 hours. Carbon/silica (C/SiO₂) microspheres were thus obtained. The PMC was produced by removing the silica templates in a 10% HF solution. Silica sols with different particle sizes (mode at 40 nm or 10 nm, or a mixture of 40/10 nm at 1:1 volume ratio) were used as templates for PMC. The PMC materials using different silica NP sizes were denoted as PMC-40, PMC-10, and PMC-40:10, respectively.

Porous Microsphere Carbon loaded with Sulfur (PMC/S): Elemental sulfur (SigmaAldrich, 99.99%) was loaded onto the PMC via liquid phase infiltration by using a freshly made solution of sulfur in carbon desulfide (CS₂). The infiltration of sulfur was carried out through impregnation and drying process to obtain samples with designated loadings. After each cycle of impregnation and drying, the sample was annealed at 155°C for 1 hour and 200°C for another hour in the N₂ atmosphere. Typically, five such cycles were applied to reach the desired total loading of sulfur. The corresponding PMC/S materials using different size silica as templates were denoted as PMC/S-40, PMC/S-10, and PMC/S-40:10, respectively.

The thermal stability of electrode materials were carried out on a thermalgravimetric analyzer (TGA-DAT-2960 SDT) at a heating rate of 20 °C min⁻¹ from 25 to 500 °C in argon. Scanning electron microscopy (SEM) equipped with energy-
dispersive X-ray (EDX) spectroscopy (Hitachi S4800) was used to analyze the material morphology and elemental composition. To carry out x-ray elemental mapping on a cross section of the PMC/S, we used a special resin to embed the PMC/S sample and prepared SEM specimen according to the method reported by Holdorf et al. The Brunauer-Emmett-Teller (BET) specific surface area of the composites was measured by nitrogen adsorption at 77 K on a surface area and porosity analyzer (Micrometrics ASAP 2020). Before each adsorption measurement, 0.10 g PMC sample was degassed at 300 °C for 4 h. It is not possible to follow the conventional protocol to degas the PMC/S samples above 155 °C before BET measurements owing to the volatility of the sulfur, and thus no pretreatment was done for PMC/S. The pore size distribution plot was derived from the adsorption branch of the isotherm based on the density functional theory (DFT).

The active material powder was mixed with carbon black and Poly(vinylidene fluoride), PVDF, dissolved in n-methyl pyrrolidinone, NMP, 8 wt.%, in a weight ratio 75:15:10. The slurry was mixed to obtain a homogeneous black paste which was then coated on an aluminum foil. The as-coated aluminum foils were dried under vacuum at 60°C for 1 h. The working electrode and Li metal foil counter electrode were assembled into coin cells by using Celgard 2400 as the separator and 1 M LiTFSI in DOL/DME 50:50 was used as the electrolyte. The cells were constructed in an Argon-filled glove box. The charge/discharge curves were measured using an Arbin potentiostat at a cutoff voltage of 1.5-3.0 V under a current density of 1 C. The specific capacity and current density were calculated based on sulfur mass only. Cyclic voltammetric curves were recorded in a voltage range of 1.5-2.8 V at a scan rate of 0.1 mV s⁻¹. The electrochemical impedance spectra were recorded by applying an alternating current of 5 mV in a frequency range
from 1 Hz to 100 kHz.

7.3 Results and Discussion

Scanning electron microscopy (SEM) coupled with energy-dispersive X-ray (EDX) spectroscopy were employed to identify the morphology, pore structure, and the distribution of sulfur in the 3D multi-modal PMC/S. Fig. 7.2a shows C/SiO$_2$-40 microspheres where 40 nm silica NPs are clearly seen imbedded in a carbon matrix. Fig. 7.2b shows porous carbon microspheres (PMC-40) after 40 nm silica templates were etched out. Because silica NPs have a size range of 30 to 50 nm with a mode diameter of 40 nm, some size variation of the pores is discernible. EDX analysis was conducted, and no Si was present on the PMC-40. PMC-40 could be completely burned off, further confirming silica templates were completely removed. This also suggests that the carbon walls in between the silica NPs were porous, allowing HF etching solution to penetrate into the entire microsphere. Fig. 7.2c and its inset show the SEM image of PMC-10, which is composed of smaller pores (~10 nm). SEM images of PMC-40:10, prepared by using mixed 40 and 10 nm silica templates at 1:1 ratio, are exhibited in Fig. 7.2d-f. Figure 1d clearly shows a mixed larger pores and smaller pores on the surface of PMC-40:10, agreeing well with the pore sizes of the two silica NP templates. The inner structure of the porous framework can be identified by a broken microsphere of PMC-40:10 (Fig. 7.2e) and a magnified image (Fig. 7.2f). Because of the equal total volume of 40 and 10 nm pores, the number of 10 nm pores is far more than that of the 40 nm pores. Hence, one can see that the large pores are surrounded by many smaller pores, and the smaller pores themselves are connected by carbon walls that are microporous (see pore size analysis in a later section).
Figure 7.2 SEM Images of (a) C/SiO$_2$-40, (b) PMC-40, (c) PMC-10 (inset showing magnified surface pore structures), (d) PMC-40:10, (e) PMC-40:10 with exposed inner surface, and (f) magnified image of PMC-40:10 inner pore structure.

The interconnected pores, although at different sizes, ensure uniform sulfur loading, and a high sulfur concentration is achievable. Upon sulfur infiltration and infusion, the pores on PMC-40 were partially occupied by sulfur without changing the spherical morphology (Fig. 7.3a). Fig. 7.3b shows the corresponding SEM image of PMC/S-40:10. After sulfur infiltration, the larger pores on PMC/S-40:10 still remain open, while the smaller pores were fully or partially blocked. The SEM image of PMC/S-10 is shown in Fig. 7.3c, which suggests that sulfur fully covered the surface due to the small pore size not easily discernible from SEM images. EDX and X-ray elemental mapping (XREM) results (see Fig. 7.4) demonstrate that carbon and sulfur are evenly distributed on the microsphere. To confirm the distribution of sulfur within the porous carbon, the PMC/S-40 sample was cut in the middle and its cross section was subject to XREM (see Fig. 7.5). The results confirm that sulfur was also uniformly dispersed in the entire inner pores of the
carbon framework. The sulfur weight percentage in the PMC/S composites ranged from 60.5% to 63.9% for the three samples.

**Figure 7.3** SEM Images of (a) PMC/S-40, (b) PMC/S-40:10, and (c) PMC/S-10

**Figure 7.4** SEM image (a), and X-ray elemental mapping of PMC/S-40: (b) Carbon element, and (c) Sulfur element.

**Figure 7.5** SEM image of the cross-section of PMC/S-40 (a), and X-ray elemental mapping images of the cross section of PMC/S-40: (b) carbon element, (c) sulfur element.

**Fig. 7.6** shows the XRD patterns of the pristine sulfur and PMC/S composites. The sharp diffraction peaks centered at $2\theta = 23.4^\circ$ and $28.0^\circ$ in the XRD pattern of pristine
sulfur match well with orthorhomic phase sulfur (JCPDS no. 08-0247). The XRD patterns for PMC/S composites are quite different from that of pristine sulfur. Only one broad peak can be observed at about 23.0° represents the presence of partially graphitized carbon structure in PMC. Sulfur diffraction peaks completely disappear for all the PMC/S composites, which suggests that sulfur is highly dispersed inside the pores of the carbon microsphere.\(^{73}\)

![XRD patterns](image)

**Figure 7.6** XRD patterns of (a) PMC/S-40, (b) PMC/S-40:10, (c) PMC/S-10, and (d) pristine sulfur

| Table 7.1 summarizes the physical characteristics of PMC and PMC/S nanocomposites derived from Brunauer-Emmett-Teller (BET) measurements. T-plot method, the most commonly method to determine the properties of microporous materials, is used to indicate micropore surface area and micropore volume of the PMC and PMC/S composites. The PMC materials have high specific surface areas and large pore volumes. Significantly, PMC-10 has an extraordinary high specific surface area of 2,776 m\(^2\) g\(^{-1}\), |
followed by PMC-40:10 at 2,485 m² g⁻¹ and PMC-40 at 1,236 m² g⁻¹. After sulfur infiltration in the porous carbon, the specific surface area and pore volume decrease remarkably, indicating that the pores of PMC are partially filled.

### Table 7.1 BET specific surface area and pore volume of PMC and PMC/S nanocomposites

<table>
<thead>
<tr>
<th>Samples</th>
<th>BET specific surface area (m² g⁻¹)</th>
<th>Pore volume (cm³ g⁻¹)</th>
<th>Micropore surface area (m² g⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PMC-40</td>
<td>1236</td>
<td>3.1</td>
<td>173</td>
</tr>
<tr>
<td>PMC-40:10</td>
<td>2485</td>
<td>5.1</td>
<td>118</td>
</tr>
<tr>
<td>PMC-10</td>
<td>2776</td>
<td>4.3</td>
<td>67</td>
</tr>
<tr>
<td>PMC/S-40</td>
<td>242</td>
<td>1.1</td>
<td>14.9</td>
</tr>
<tr>
<td>PMC/S-40:10</td>
<td>134</td>
<td>0.4</td>
<td>6.6</td>
</tr>
<tr>
<td>PMC/S-10</td>
<td>82</td>
<td>0.2</td>
<td>1.9</td>
</tr>
</tbody>
</table>

The porous structure and pore size distribution of PMC and PMC/S samples were investigated by N₂ adsorption-desorption isotherms. The results in Fig. 7.7a and 7.7b show that both PMC and PMC/S materials had a typical IV-type isotherm, corresponding with a mesoporous structure. The sharp incremental of N₂ adsorption amount on the isotherms for PMC-40 and PMC-40:10 materials at the relative pressure p/p₀ = 0.93–1.0 indicates the existence of macroporosity.⁷⁴ PMC-10 exhibited no such characteristic sharp incremental N₂ adsorption at p/p₀ = 0.93-1 (i.e., absence of macropores) because only 10 nm template SiO₂ NPs were used. Compared with bare PMC in Fig. 7.7a, the quantity of adsorbed N₂ in Fig. 7.7b significantly decreased in PMC/S due to the sulfur infiltration into the porous carbon, in agreement with the BET measurements of specific surface area. The calculated pore size distribution (PSD) curves of PMC (Fig. 7.8a) and PMC/S (Fig. 7.8b) further reveal the feature of multi-modal porous structure. Table 7.2 summarizes the pore size
distribution obtained from Fig. 7.8. All PMC samples had micropores around 1.5 nm, likely originated within the carbon walls (due to the interspaces of carbon nanoparticles and/or formed by gas releasing during HF etching). The macropores represented as a weak shoulder on the PSD curve at the mode of 55 nm for PMC-40 and PMC-40:10 are likely due to the relatively wide size range of silica NPs that contain a portion of NPs greater than 40 nm. Besides the sharp characteristic peaks around 10 and 40 nm, mesopores at other sizes are also observed. For the PMC/S composites, the micropores disappeared, indicating that micropores were fully occupied by sulfur. There are some new peaks (e.g. 20 nm) formed on PMC/S-40 and PMC/S-40:10, which are likely due to partial occupation of the 40 nm pores by sulfur.

Figure 7.7 N₂ adsorption/desorption isotherms of (a) PMC and (b) PMC/S
Table 7.2 Summary of pore size distribution and modes for PMC and PMC/S nanocomposites

<table>
<thead>
<tr>
<th>Samples</th>
<th>Micropores (&lt; 2 nm)</th>
<th>Mesopores (2~50 nm)</th>
<th>Macropores (&gt; 50 nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PMC-40</td>
<td>1.5</td>
<td>8, 36(D)</td>
<td>55</td>
</tr>
<tr>
<td>PMC-10</td>
<td>1.5</td>
<td>4, 10(D)</td>
<td>No</td>
</tr>
<tr>
<td>PMC-40:10</td>
<td>1.5</td>
<td>6, 10(D), 30~44(D)</td>
<td>55</td>
</tr>
<tr>
<td>PMC/S-40</td>
<td>No</td>
<td>9, 14, 20, 32~44(D)</td>
<td>55</td>
</tr>
<tr>
<td>PMC/S-10</td>
<td>No</td>
<td>9(D)</td>
<td>No</td>
</tr>
<tr>
<td>PMC/S-40:10</td>
<td>No</td>
<td>9(D), 20, 28~40(D)</td>
<td>55</td>
</tr>
</tbody>
</table>

Note: "(D)" indicates dominating pore size mode.

Figure 7.8 Pore size distribution of (a) PMC and (b) PMC/S (insets showing magnified pore size distribution in the dashed rectangular area)

Fig. 7.9a shows the charge/discharge voltage profile for the initial cycle of PMC/S at a high current density rate of 1 C (1 C = 1,675 mA g\(^{-1}\)). Two voltage plateaus during the discharge process can be observed for all PMC/S materials. An impressive high initial discharge capacity of 1,278 mAh g\(^{-1}\) was achieved on PMC/S-40:10, corresponding with 76.3 % of the theoretical capacity of sulfur. In comparison, PMC/S-40 and PMC/S-10 exhibited an initial discharge capacity of 1,158 mAh g\(^{-1}\) and 940 mAh g\(^{-1}\), respectively, lower than that of PMC/S-40:10.
The rate capability of the PMC/S composites at various current densities from C/10 to 2C is shown in Fig. 7.9b. PMC/S-40:10 composite shows an excellent rate performance, with specific capacities of 1462, 1053, 937, 855, 778 mAh g\(^{-1}\) at C/10, C/5, C/2, 1C and 2C, respectively. PMC/S-40:10 exhibits the highest capability at all rates compared with PMC/S-40 and PMC/S-10. In addition, the superior rate capability of PMC/S-40:10 is also demonstrated by the easier recovery of capacity when the current density was changed from 2C to C/10 at the end of the test.

Cycle performance and Coulombic efficiency of the three samples and pristine sulfur were compared over 100 cycles at 1 C rate, as shown in Fig. 7.9c. The initial discharge capacity follows the order of PMC/S-40:10 > PMC/S-40 > PMC/S-10 > sulfur. The discharge capacities at the 100\(^{th}\) cycle were 904, 660, 680 and 419 mAh g\(^{-1}\) for PMC/S-40:10, PMC/S-40, PMC/S-10, and sulfur, respectively. Again, in agreement with the trend as seen in the rate capability performance, the PMC/S-40:10 had the best long term performance, with 70.7% capacity retention after 100 cycles compared with its initial capacity. For PMC/S-40:10, the initial Coulombic efficiency was 98.4% and slightly dropped to 90% after 40 cycles, but remained stable thereafter at around 90%. As comparison, PMC/S-40 and PMC/S-10 had a Coulombic efficiency of 88% and 87%, respectively, over 100 cycles. The results indicate that all the PMC/S samples exhibited superior cycle performance and mitigated shuttle effect compared with pristine sulfur.
Figure 7.9 Electrochemical characterization and battery performance results of PMC/S: (a) initial cycle charge-discharge profiles between 1.5 V and 3.0 V at a current density of 1 C rate, (b) comparison of the rate capability of PMC/S at current density from C/10 to 2 C, and (c) cycle performance and Coulombic efficiency of PMC/S and pristine sulfur samples over 100 cycles at 1 C rate.

The higher electrochemical performance of PMC/S-40:10 than those of PMC/S-40 and PMC/S-10 can be explained by the differences in their pore structures (see Table 7.2); i.e., PMC/S-40:10 has a unique combination of macropores, mesopores, and micropores. First, the macropores (~55 nm) and large mesopores (~40 nm) allow easy electrolyte diffusion and Li$^+$ transport, and in the meantime, accommodate sulfur volume expansion during cycling. Second, the small mesopores (~10 nm) bridge the larger pores (40~55 nm) and provide passages and easy access in between the larger pores, thus lower the resistance
in charge transfer and diffusion. Third, the very small mesopores (4~6 nm) and micropores (< 2 nm) enhance the contact between carbon and sulfur due to their extremely small size, and they help trap polysulfides and prevent active sulfur material from escaping the carbon framework to a certain extent. In short, the hierarchical structure of macropore-mesopore-micropore network with differential pore sizes provide easy access for sulfur infiltration, electrolyte diffusion, and ion transport through larger pores, while the smaller pores surrounding the larger pores to some extent prevent polysulfides from leaving the larger pores so that each microsphere acts as a good sulfur container.

Comparing PMC/S-40 with PMC/S-40:10, the lack of large population of small mesopores in the 10 nm range in PMC/S-40 may have caused difficulty for larger pores (40~55 nm) to “communicate” with each other (e.g., ion transport and electrolyte permeation between the large particles), as the larger pores are segregated by carbon walls that have only micropores. In addition, large size pores (like the 40~55 nm pores in PMC/S-40) cannot effectively trap lithium polysulfides. In another word, PMC/S-40:10 performed better than PMC/S-40 because the large number of 10 nm small mesopores in PMC/S-40:10 may have created easier passages for ion transport in between the 40~55 nm pores and at the same time mitigated the loss of active sulfur by confining polysulfides to some extent. Comparing PMC/S-10 with PMC/S-40:10, the lack of large pores (40~55 nm) and the presence of only small mesopores (< 10 nm) in PMC/S-10 may have caused infiltrated sulfur to block the small pores, making electrolyte permeation and ion transport more difficult in PMC/S-10. Micropores and small mesopores have advantages in constraining lithium polysulfides to inhibit the "shuttle effect", but too small micropores (<0.7 nm) are not accessible to guest molecules or electrolyte ions. These could
explain that PMC/S-10 with only small pores exhibited lower initial capacity compared with those of PMC/S-40 and PMC/S-40:10 that have a considerable amount of large pores. The results from electrochemical impedance spectroscopy (EIS), as shown in Fig. 7.10, also confirm that PMC/S-40:10 has the lowest impedance, with PMC/S-10 having the highest impedance. The EIS results agree with our above analysis that PMC/S-40:10 has a unique hierarchical pore structure that is in favor of charge transport and reducing ohmic resistance.

![Electrochemical impedance spectroscopy (EIS) curves of PMC/S-40, PMC/S-40:10, and PMC-10 before initial discharge.](image)

The electrochemical performance data obtained using our PMC/S-40:10 cathode material (1,278 mAh g⁻¹ initial capacity, and 904 mAh g⁻¹ after 100 cycles at 1 C rate with stable 90% Coulombic efficiency) are comparable to the results in the leading literature on Li-S battery with sulfur loaded on porous carbon. But our material synthesis method is
relatively simple and our raw materials used are cheaper with promising scale-up potential. Based on our results in this work and the above literature review, we believe the shuttle effect still exists to some extent in our PMC/S materials because of the open porous structure that allows free access of electrolyte to the sulfur. However, the shuttle effect was significantly mitigated compared with the pristine sulfur without the carbon framework. Our work demonstrated that multi-modal porous carbon with relatively wide pore size distributions is also an excellent substrate for Li-S cathodes. Among the three porous PMC/S samples, the one with the mixed pore sizes, PMC/S-40:10, has the highest capacity and Coulombic efficiency (reduced shuttle effect), although its surface area is less than that of PMC/S-10. This is an original finding in this work in terms of a new material design.

Magasinski et al.\textsuperscript{182} indicated in a Li-ion battery study that an aperiodic porous network would allow for the redirection of the ion traffic if the path of Li ions is blocked or impeded in a narrow channel. Similarly, the hierarchically interconnected porous network with wide PSD warrants our PMC/S-40:10 nanocomposite the similar ability of redirecting the ion traffic to maintain rapid charging capability.

7.4 \textbf{Summary and Conclusion}

In summary, this work proves a novel concept that a sulfur infiltration 3D multi-modal PMC with interconnected micropores, mesopores and macropores can serve as a promising cathode material for Li-S batteries. The developed hierarchical pore structure offers several advantages: (1) interconnected multi-modal porous carbon framework in close contact with sulfur assures superb electrical conductivity (or low impedance), fast ion transport, and excellent electrolyte permeation to active sulfur within the entire framework; (2) large pores facilitate sulfur infiltration and small pores help trap
polysulfides and inhibit sulfur loss to some extent; and (3) sufficient total pore volume to accommodate sulfur volume expansion during cycling. The unique structure of small mesopores (~10 nm) as bridging agents between larger pores (40~55 nm) is indispensible in providing the above three advantages, rendering the PMC/S-40:10 material the best electrochemical performance among the PMC/s samples tested. We have demonstrated that a discharge capacity of 1,278 mA h g\(^{-1}\) was achieved at 1 C current density and the capacity maintained at 904 mA h g\(^{-1}\) even after 100 cycles. This electrochemical performance is among the highest ones reported in the open literature. It is possible that further tuning the pore size distribution of PMC/S (by adjusting silica template sizes and their ratios) will lead to higher and more stable cycling performance. In addition, coating a conductive layer on the PMC/S microspheres may further prohibit the shuttle effect and enhance the Coulombic efficiency. The aerosol-based spray pyrolysis synthesis process is simple, inexpensive, and can be easily scaled up, providing promising opportunities to commercialize the PMC materials that could be applied in a variety of electrode materials.
CHAPTER 8 SUMMARY OF RESEARCH WORK AND PROPOSED FUTURE WORK

8.1 Summary of Research Work

1. *Nanostructured Ce-TiO$_2$/SBA-15 Composites for CO$_2$ Photoreduction*

Ce-TiO$_2$/SBA-15 were synthesized via sol-gel method to help facilitate e-h pairs separation and improve catalysts dispersion. Modification of TiO$_2$ with Ce significantly stabilized the TiO$_2$ anatase phase and increased the specific surface area, which contributed to an improvement of CO production from CO$_2$ reduction. Dispersing Ce-TiO$_2$ nanoparticles on the mesoporous SBA-15 support further enhanced both CO and CH$_4$ production. We proposed that the superior catalytic activity may be related to the partially embedded Ce–TiO$_2$ nanoparticles in the ordered 1-D pores in SBA-15 that form synergies between the different components of the catalysts and enhance the diffusion and adsorption of CO$_2$. This mechanism also correlates well the results that using SBA-15 as the support led to more than 10 times higher activity in CO$_2$ photoreduction than using amorphous silica as the support. Findings in this work demonstrate the feasibility of solar fuel production from CO$_2$ and H$_2$O using the prepared nanocomposite photocatalysts.

2. *Ag/TiO$_2$ Nanocomposite Photocatalysts For Simultaneous H$_2$ Production And CO$_2$ Reduction*

Ag/TiO$_2$ were fabricated via ultrasonic spray pyrolysis technique. The material property analysis and photocatalytic activity results showed that the ultrasonic spray pyrolysis method is much superior to conventional wet impregnation process with the advantages of smaller Ag nanoparticles, a better Ag dispersion on TiO$_2$, and a higher fraction of metallic Ag species, which facilitate charge transfer and improve photocatalytic
activity. The rates of photocatalytic conversion in this work were among the highest ones reported in the literature. This work has also demonstrated the feasibility of syngas (H₂ and CO) production from a gas mixture of CO₂, H₂O and CH₃OH through a photocatalytic reduction process on Ag/TiO₂ nanocomposite catalysts under solar irradiation. Although the H₂ production and CO₂ reduction processes compete for electrons and protons in the photocatalytic process, the final product selectivity (H₂/CO ratio as low as 2, with negligible CH₄) is in the appropriate range for subsequent Fischer-Tropsch process for liquid hydrocarbon fuel production. The selectivity is tunable by varying the reaction gas composition and the H₂/CO ratio in the range from 2 to 10 has been achieved.

3. Hybrid MgAl-LDO/TiO₂ Composites for CO₂ Photoreduction

MgAl-LDO/TiO₂ composites were prepared by a combination of hydrothermal and co-precipitation method. This work has demonstrated that the successful synthesis for novel hybrid structure of MgAl-LDOs platelets grafted on TiO₂ cuboids with high length to width ratio. MgAl-LDOs addition has no impact on the optical properties of TiO₂. The morphology of TiO₂-MgAl-LDOs showed that the surface coverage on TiO₂ on MgAl-LDOs platelets could be controlled by tuning the MgAl-LDOs to TiO₂ ratio. As the MgAl-LDOs concentration increases, TiO₂ crystal size was increased. MgAl-LDOs grafting on TiO₂ cuboids may help improve the adsorption ability of CO₂ onto TiO₂ which improves the photocatalytic activity of CO₂ reduction. The optimum MgAl-LDOs concentration was found to be 10% which performed the highest CO production rate. The proposed reason for this is: at higher MgAl-LDOs concentration, MgAl-LDOs platelets covered the majority parts of TiO₂ cuboids surface either blocked the CO₂ and H₂O vapor contacting with TiO₂ or electrons transferring from TiO₂ to CO₂ and H₂O molecules; Moreover, the
crystal size increase of TiO2 anatase may also lead to a poor CO2 photoreduction activity. At lower MgAl-LDOs concentration, less CO2 and H2O amount may be adsorbed onto the composites which also leads to lower CO production rate.

4. Carbon-TiO2 Nanocomposites Prepared With Simple Method For Lithium-Ion Batteries (LIBs)

Carbon decorated TiO2 were synthesized using commercial TiO2 product (P25 and P90) in large scale through a simple carbonization process. C-P90 possesses higher specific area and larger fraction of anatase comparing to C-P25 with the same carbon concentration, which guaranteed its superior electrochemical performance as anode for LIBs. Carbon concentration had been optimized to obtain the best battery performance for P90. 1.9% C-P90 showed a significant stable cyclablity which was attributed to its unique carbon decoration nanostructure that facilitates efficient lithium insertion and extraction. The easy preparation together with the advantage of stable cyclability of C-P90 makes the composite a very promising anode material for LIBs.

5. Hierarchical Sulfur-infiltrated porous microsphere carbon as cathode for Li-S batteries

PMC materials were synthesized via aerosol method. The aerosol-based spray pyrolysis synthesis process is simple, inexpensive, and can be easily scaled up, providing promising opportunities to commercialize the PMC materials that could be applied in a variety of electrode materials. This work proves a novel concept that a sulfur infiltration 3D multi-modal PMC with interconnected micropores, mesopores and macropores can serve as a promising cathode material for Li–S batteries. The developed hierarchical pore structure offers several advantages: (1) an interconnected multi-modal porous carbon framework in close contact with sulfur assures superb electrical conductivity (or low
impedance), fast ion transport, and excellent electrolyte permeation to active sulfur within the entire framework; (2) large pores facilitate sulfur infiltration and small pores help trap polysulfides and inhibit sulfur loss to some extent; and (3) sufficient total pore volume to accommodate sulfur volume expansion during cycling. The unique structure of small mesopores (~10 nm) as bridging agents between larger pores (40–55 nm) is indispensable in providing the above three advantages, giving the PMC/S-40:10 material the best electrochemical performance among the PMC/S samples tested. It is possible that further tuning the pore size distribution of PMC/S (by adjusting silica template sizes and their ratios) will lead to a higher and more stable cycling performance. In addition, coating a conductive layer on the PMC/S microspheres may further prohibit the shuttle effect and enhance the Coulombic efficiency.
8.2 Proposed Future Work

1. PMC/S Composites for Li-S Batteries

As demonstrated in Chapter 4, PMC/S with multi-modal pore size distribution materials have been synthesized using aerosol technique for Li-S battery. The further work will be carried out to improving the electrochemical performance of Li-S battery.

In 2009, Nazar and co-workers reported a pioneering work in the Li-S cathode material by incorporating sulfur into a highly ordered mesoporous carbon (CMK-3) and coating sulfur-CMK-3 with polyethylene glycol (PEG).\(^{15}\) Those materials exhibited good stability over 20 cycles tested and had high initial discharge capacities at 1,320 mA h g\(^{-1}\) and 1,005 mA h g\(^{-1}\) at 0.1 C rate (1 C = 1,675 mA/g) with and without PEG modification, respectively. Nazar’s work inspired the investigation of many carbon materials as sulfur substrates such as porous carbon frameworks,\(^{29, 56-60}\) carbon nanotubes,\(^{34, 61-62}\) hollow carbon nanofibers,\(^{16}\) graphene,\(^{63-64}\) and conductive polymers.\(^{15, 65-66}\) Cui et al.\(^{65}\) reported a poly (3,4-ethylenedioxythiophene):poly (styrene sulfonate) (PEDOT:PSS) coated CMK-3/sulfur that could achieve 96-98% Coulombic efficiency. Following this direction, PMC/S materials can be further modified such as coating with conductive polymer to confine sulfur more efficiently.

Wang et al.\(^{34}\) reported a sulfur-impregnated disordered carbon nanotubes cathode for lithium-sulfur batteries. They prepared this materials at 300-500 °C and they proposed that carbon and sulfur could obtain bonding function to avoid sulfur loss during charge-discharge process. Following this direction, we can also improve our PMC/S preparation procedure to obtain "bonded" carbon-sulfur materials.

The proposed future work for Li-S battery project is summarized as follows:
• To find a better way to modify the PMC/S materials.

• 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), and thermogravimetric analysis (TGA). And the nanomaterials will be used as cathode materials for lithium-sulfur batteries.

• Electrodes and half-coin cell will be prepared and their performance will be investigated.

• The performance of PMC/S nanomaterials as cathode will be optimized.
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CURRICULUM VITAE

CUNYU ZHAO

EDUCATION
Ph.D., Mechanical Engineering, University of WI-Milwaukee (Dec. 2014)
M.S., Chemical Engineering, Tianjin University, P.R. China (2008)
B.S., Chemical Engineering, Hefei University of Technology, P.R. China (2006)

PROFESSIONAL SKILLS
Gas chromatography (Agilent 7890A); Mass spectroscopy (Agilent 5975C); Physisorption apparatus (Micromeritics ASAP 2020); X-ray diffractometer (XRD); Scanning electron microscopy (SEM); Transmittng electron microscopy (TEM); Demonstrated experience with battery electrochemical characterization e.g. impedance spectroscopy (EIS), galvanostatic potentiometry, cycle life testing.
Microsoft Office software; PROII simulation software; Aspen Software; AutoCAD.

PROCESS ENGINEER EXPERIENCE
GTC (Beijing) Technology, Inc. Process Engineer Beijing, P.R. China 2008-2011
Responsible for aromatic extractive distillation process design, including process simulation, material balance, equipment sizing, piping and instrumentation design, and process flow design and analysis. Performed cost estimates, submitted project proposals, developed operating guidelines, start-up procedures and process design package.

Project Experiences
- Key person for project coordination among Sinopec Yanshan Petrochemical, SEI and GTC for Yanshan Styrene Recovery project. Both customers and PDP clients were satisfied with the immediate action upon project minute of meeting.
- Designed condensate recovery system for SECCO styrene project in Shanghai, China. Saved 10% energy of the total energy consumption; Specified instrumentation and line list for the project. Efficiently accomplished the project for customers.
- Designed GT-Styrene Recovery process for Jilin Petrochemical Co. based on specified raw pygas material with GTC's proprietary technology. Simulated the styrene extractive distillation process and optimized the process with heat integration by saving 10% energy. Prepared PFD drawing, material and utility balance.
- Designed GT-BTX process for Petrochina Daqing Co. Simulated the BTX Extractive Distillation process and Optimized the process with heat integration. Prepared PFD drawing, material and utility balance and Cost Estimation.
- Assisted start-up expert on Petrochina Dushanzi Co. BTX project start-up. Familiarized with on-site BTX unit problems and provide methods for resolving problems. Assisted foreign expert to communicate effectively with client.

TEACHING EXPERIENCE
University of WI-Milwaukee Teaching Assistant Milwaukee, WI Sep.2011- May.2012
- Fluid Mechanics Lecture and Lab. The students were satisfied about the course instruction and lab instruction. 97% of the class (32 students) got A- above score.
RESEARCH EXPERIENCE
• Research on TiO$_2$-based catalysts for application in CO$_2$ photoreduction.
• Innovated hybrid nanomaterials using as photocatalysts for CO$_2$ photoreduction with water. Improved the photocatalytic conversion activity of CO$_2$ with water to alternative fuels.
• Explored hybrid nanostructures alternative materials as anode for Lithium-ion batteries. By growing carbon on TiO$_2$, attained the good cyclability and rate capability for Li-ion batteries.
• Research, design and develop novel nanomaterial structure for use as cathodes for Li-S batteries and Li-ion batteries.
• Innovated porous microsphere carbon/sulfur nanostructure for Li-S battery cathode materials. Inhibited shuttle effect and increase the conductivity of the electrode.

HONORS & AWARDS
• "Top 3" poster award in the graduate student poster competition at the Annual Spring Symposium of the Catalysis Club of Chicago held on May 13, 2014 at the BP Research Center.
• Third place prize in the graduate student poster competition UWM-CEAS Annual Poster Competition, 2014.
• University of Wisconsin-Milwaukee Distinguished Graduate Student Fellowship Award, 2014-2015.
• UWM Chancellor’s Graduate Student Award, University of Wisconsin-Milwaukee, Milwaukee, WI, 2012-2014.
• First place prize in the graduate student poster competition at the Nano Science and Engineering Forum (NSEF) in 2012 at the AIChE Annual Meeting in Pittsburgh, PA.

PUBLICATIONS


• Zhang, Y., Zhao, C.Y., Liang, H., Liu, Y.*. Macroporous Monolithic Pt/γ-Al\textsubscript{2}O\textsubscript{3} and K–Pt/γ-Al\textsubscript{2}O\textsubscript{3} Catalysts Used for Preferential Oxidation of CO. Catalysis Letters, 2009, 127 (3), 339-347.


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