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Modeling and Analysis of Bow-Tie Antenna Integrated Resonant-Tunneling-Diode Relaxation Oscillators for Wireless Radio Applications

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Abstract

We propose a self-complementary bow-tie antenna-integrated resonant-tunneling-diode relaxation oscillator and investigate its oscillation/radiation characteristics. In the investigation, we establish a physics-based equivalent circuit model of the oscillator for taking the all physical phenomena related to the diode and the antenna into consideration simultaneously. In this paper, we report the equivalent circuit modeling and the large-signal oscillation/radiation analysis of the oscillator.

Introduction

Terahertz (THz) technology has been received great deal of interest in terms of applications for spectroscopy, imaging, wireless communications, etc. In the wireless communication, the THz range is specifically one of the most attractive regions owing to the existence of the non-allocated frequency band. One of the main challenges of the THz wireless communication is the shortage of the emission RF power of the oscillator. We guess the challenge can be overcome by using the wideband-spectrum relaxation wave, whose entire emission RF power exceeds that of the narrowband sinusoidal wave, as the carrier.

From this perspective, we have proposed an oscillator consisting of a resonant tunneling diode (RTD) and a broadband bow-tie antenna (BTA) [1] for generating a wideband-spectrum relaxation wave and their theoretical models expressed by equivalent circuits respectively [2,3]. Such a semiconductor-based oscillator for THz applications is generally fabricated as a monolithically-integrated configuration of an oscillation device, a radiation antenna, etc. As the performance of such an oscillator depends upon a both of their physics, which usually differ from each other, the performance evaluation method and the design guideline for the oscillator should be established as a physical-coupled scheme considering the all effects regarding the oscillator simultaneously.

In this paper, we focus on a physics-based modeling for the entire oscillator-structure we proposed and its oscillation/radiation performance evaluated by using the physics-based model.

Modeling

Schematics of the proposed oscillator are illustrated in Figure 1. A peripheral circuit in front serves as a bias circuit and that in back is added for adjusting the total impedance of the peripheral circuits. We establish a physics-based equivalent circuit model of each component, RTD, BTA, and the circuits, since it is considerably impossible to analyze the quantum phenomena of the RTD together with the electromagnetic (EM) properties of the other components by a single solver.

Figure 1 Schematics of a proposed oscillator.

Simple block expressions of the oscillator are described in Figure 2.

The impedance variables, \( Z_f(\omega, D, w_{\text{shunt}}) \), \( Z_b(\omega, D, w_{\text{shunt}}) \), and \( Z_{\text{BTA}}(\omega, D) \), correspond to the equivalent circuit of the peripheral circuit in front, that in back, and the BTA, respectively. The RLC expression of \( Z_{\text{BTA}}(\omega, D) \) has been reported in Ref. [2]. Additionally, the circuit model of the RTD and its theoretical expression has also been reported in Refs. [3, 4], respectively.

The variables, \( Z_f(\omega, D, w_{\text{shunt}}) \) and \( Z_b(\omega, D, w_{\text{shunt}}) \), are com-
posed of several RLC elements which can be explained by the electromagnetic properties: the surface impedance due to the skin effect [5], the straight micro stripline [6], fringe capacitance [7], and parasitic components regarding the semi-insulating substrate. The circuit elements involved in $Z_{f}(\omega,D,w_{shunt})$ and $Z_{b}(\omega,D,w_{shunt})$ are evaluated by the EM field distribution in the vicinity of the circuits calculated by the finite element method-based simulator, namely, COMSOL. Their approximate numerical values of the elements are also estimated by the physical interpretation based on the structural and material properties. More precise values are numerically de-embedded by using the optimization method, namely, the particle swarm optimization [8]. More details regarding the circuit identification process have also been reported in Ref. [2].

Figure 3 summarizes the typical fitting results of the im-
pedance and radiation characteristics regarding the peripheral circuit/entire oscillator.

From Figure 3, the equivalent circuit expression is quantitatively valid for the evaluation of the radiation/oscillation characteristics by the circuit analysis around the first resonance frequency of the oscillator.

**Oscillation Analysis**

The non-linear oscillation analysis of the proposed oscillator is performed by using the equivalent circuit above mentioned. The non-linearity of the RTD is involved by considering the measured current-voltage (I-V) characteristics [9]. The supplied voltage, $V_b$, is set to a constant value to keep the negative differential conductance of the RTD maximum. More details regarding the oscillation analysis methodology have been reported in Refs. [10, 11].

Figure 4 displays the current density-voltage (J-V) characteristics of the RTD and the time-dependent orbit of the current density, $i_{rtd}(t)/S$, in the RTD, where $S$ indicates the mesa area.

We classify the oscillation modes depicted the dotted and chain lines in Figure 4 designated as the "sinusoidal" and the "relaxation" mode, respectively. The modes are quantitatively distinguished by the cycle number of the $i_{rtd}(t)/S$ trajectory in Figure 4; if the cycle number is unity, the mode is a "sinusoidal mode"; else, it is a "relaxation mode". The entire emission RF power of the relaxation mode shown in Figure 4 is ~ 6 dBm greater than that of the sinusoidal mode when the upper limit value of band is approximately set to 400 GHz. It is suggested that the relaxation wave can compensate for the shortage of the emission RF power if we employ a certain band appropriately. Moreover, it is found that the oscillation mode can be designed by adjusting the two parameters, $D$ and $w_{shunt}$, appropriately.

According to Shannon-Hartley theorem [12], the capacity of a wireless channel is directly proportional to the channel bandwidth. Therefore, the wideband-spectrum relaxation wave can contribute to the possibility of large-capacity wireless transmissions together with compensating for the RF power shortage.

**Conclusion**

We investigated radiation/oscillation characteristics of the resonant tunneling diode-based relaxation oscillator by using its physics-based equivalent circuit model. The advantage of a wideband-spectrum relaxation wave as the carrier was revealed in terms of the entire emission RF power. Our further studies will represent the link-budget analysis of the relaxation carrier wave-based THz wireless link to clarify the advantage of the relaxation carrier regarding the link performance.

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Introduction

The terahertz (THz) technology has gained much research interest worldwide. However, realization of a THz emitter with a small form factor is still a difficult task. The plasma instability in the two-dimensional (2D) electron gas has been proposed as a possible mechanism to enable an extremely fast semiconductor device operating in the THz frequency range [1].

Recently, we have reported the numerical simulation results of plasma oscillation in two-dimensional electron gas [2]. A periodic steady-state solver has been used to obtain the oscillating solution efficiently. The set of governing equations contains the 2D Poisson equation, the electron continuity equation, and the electron current density equation. The time derivative of the current density and the convective derivative are considered in the electron current density equation. It has been numerically verified that inclusion of those terms are mandatory to support sufficiently strong plasma instability in the 2D electron gas.

However, it is noted that the set of governing equations adopted in [2] is based on the moments of the Boltzmann equation. It is not sufficiently accurate to describe the behavior of plasma waves quantitatively [3]. A more advanced modelling approach is required. In order to solve this issue ultimately, the Boltzmann equation solver with the transient simulation capability is mandatory.

Although the stochastic Boltzmann equation solvers (the Monte Carlo simulators) have been used to simulate a device operating in the THz frequency range (for example, [4]), the stochastic nature of the Monte Carlo simulators introduces some numerical difficulties. Therefore, the deterministic Boltzmann equation solver is desirable. In this work, our motivation is to present some transient simulation results using a deterministic Boltzmann equation solver.

Projected Boltzmann Equation

The Boltzmann equation is a microscopic transport equation. It governs the spatiotemporal motion of the electron gas in a semiconductor device. Since it is defined on the phase space, evaluating its solution is a highly non-trivial task. As discussed above, the stochastic Monte Carlo method can be applied to solve the Boltzmann equation efficiently. However, nowadays, the deterministic Boltzmann equation solvers [5] have gained popularity. In these solvers, the angle-dependent part of the electron distribution function is expanded...
with basis functions. Depending on the dimensionality of the momentum space, there can be a few variants of deterministic Boltzmann equation solvers. In this work, the three-dimensional momentum space is understood. Therefore, the spherical harmonic expansion [6] is the appropriate technique. A numerical stabilization scheme called the H-transformation [6] has played an important role in the development of this simulation technique. In the H-transformation, the total energy (which is a sum of the kinetic energy and the potential energy) instead of the kinetic energy is used as the energy variable.

Unfortunately, when the transient simulation is concerned, the H-transformation introduces a serious difficulty [7]. The time-varying potential energy yields the time-varying transport parameters. Unavoidable interpolation procedure prevents the transient simulation capability eventually. Of course, the fundamental solution to the above problem is to invent yet another numerical stabilization scheme, which does not suffer from the time-varying transport parameters. Since it is currently a formidable task, in this work, we simply apply the kinetic-energy-based scheme [8] to get the preliminary solutions.

Numerical Results

A spherical harmonics expansion solver has been newly implemented into our in-house device simulation framework [2]. All parameters for Si are taken from [6]. The lowest expansion order is used. Various N’NN’ devices are simulated. Figure 1(a) shows the DC IV characteristics of those devices. The transient simulation results of the 1200-nm-long device are shown in Figure 1(b). The DC bias voltage is 0.8 V. The frequency of the AC voltage is 1 THz. It is clearly demonstrated that our deterministic Boltzmann equation solver can perform the transient simulation at such a high frequency without any numerical problem.

Conclusion

In this work, the transient device simulation results using a deterministic Boltzmann equation solver based on the spherical harmonics expansion has been presented. In order to avoid the numerical difficulties originated from the H-transformation, the kinetic-energy-based scheme is adopted. The numerical results demonstrate that both of DC and AC analyses can be performed without any numerical problem. It is expected that this simulator can be applied to the THz devices based on the plasma instability.

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Enhanced Catalysis by Optical Nanoantenna Reduced on Transition Metal Dichalcogenide

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Abstract

Photocatalysis of hydrogen from water is limited by incomplete absorption of solar radiation and by uncontrolled disposition of generated carriers. Nanoantenna (NA)-induced coupling of photons to excitons could enhance photocatalytic solar fuel generation by increasing broadband optical absorption and by injecting energetic electrons where NA interfaces with semiconductor catalyst. This work examined catalysis of hydrogen evolution reaction (HER) by monolayer (1L) transition metal dichalcogenide (TMD) with and without decoration by optical NA. Spectroscopic and microscopic characterization of heterostructures of 1L-TMD and NA self-assembled via exfoliation and redox chemistry was compared with discrete dipole simulation. Electrodes inked with 1L tungsten disulfide (WS2) onto which NA was electrochemically reduced exhibited higher HER relative to 1L-WS2 in neat or physically NA-decorated forms as measured by linear sweep and cyclic voltammetry. Coordinated simulation and measurement of supports improved design of 1L-TMD-NA photocatalysts and their implementation in chemical, biological, energy and water systems.

Introduction

Two-dimensional (2D) transition metal dichalcogenide (TMD) semiconductor crystals offer optoelectronic inducibility with superior electron mobility and gate tunability. This enables enhanced photocatalytic activity of interest for chemical and biological catalysis, energy, sensing, desalination, and nanoelectromechanical systems. Heterostructures of monolayer (1L) TMD crystals decorated with optical nanoantenna (NA) could improve their catalytic efficiency. However, measured catalytic efficiency of such heterostructures relative to undecorated 1L-TMD is rare and characterization of opto-electronic effects at 1L-TMD-NA heterointerfaces is largely empirical. Integration of microscopic and spectroscopic analysis with simulation of 1L-TMD-NA has been limited by computational expense and complexity, particularly for dynamic interactions at nanometer (nm) scales. Compact, multi-scale, integrated analysis of optical, electronic and catalytic effects could identify extraordinary features to guide design and implementation.

Results and Discussion

Electrochemical measurements were carried out in a three-electrode cell with a 3-mm diameter glassy carbon (GC) working electrode, a graphite rod counter electrode, and an Ag/AgCl reference electrode (Pine Research Instrumentation, Durham, NC). The cell was filled with 8 milliliters (mL) of concentrated sulfuric acid, 0.5M H2SO4. The reference electrode was kept in a salt bridge of 3M sodium chloride (NaCl). A microliter of aqueous ethanol ink containing 6 micrograms (µg) liquid phase exfoliated (LPE) tungsten disulfide (WS2) powder (Sigma-Aldrich, St. Louis, MO), 3 µg Ketjen black (EC-600JD, AkzoNobel, Chicago, IL) and 1.5 µg Nafion ionomer (Sigma-Aldrich, St. Louis, MO) was drop cast onto a glassy carbon (GC) electrode. After allowing the ink to set for 10 minutes, the three-electrode system was placed in the cell. The cell and electrodes were contained in an unlit fume hood. Electrodes were connected to a WaveNow potentiostat (Pine Research Instrumentation, Durham, NC). Linear sweep voltammetry (LSV) was performed by measuring current at a rate of 5 mV/s over a range of -0.800 mV versus the Ag/AgCl reference electrode, i.e., reversible hydrogen electrode (RHE). Bubbles forming at the electrode tip indicated hydrogen evolution. LSV was performed in triplicate. To provide background measurement, voltammetry was run with the polished working electrode prior to adding catalyst. Cyclic voltammetry (CV) performed for twenty cycles to measure persistence of catalytic activity over time.

Values of peak current density measured by LSV and CV for catalysis by LPE 1LWS2 on a GC electrode ranged from -70 to -100 mA/cm2 at -0.5 eV vs. RHE. Figure 1 shows representative CV and controls. Adjusting the position of the electrode to facilitate buoyant removal of hydrogen bubbles improved the peak current density across this range. The onset potential was approximately 250 mV without buoyant removal of hydrogen and about -300 mV after electrode adjustment. This range of measured peak current densities was greater than values previously reported for comparable measures of hydrogen evolution from WS2. From a similar experimental setup, a current density of ~25 mA/cm2 at approximately -0.45V vs SCE [1] or about -0.21V vs RHE was reported. That system consisted of three-electrodes; 25 µg of WS2 nanoflake catalyst was drop cast onto the GC electrode. Ketjen black was added to increase the conductivity of the solution. Other
similar experiments reported current densities of -20 mA/cm² at -0.5 V vs RHE and -2.2 mA/cm² at -0.2 V vs RHE, respectively [2,3]. Scarce reports of WS₂ catalysis in GC systems suggests comparison with molybdenum disulphide (MoS₂) in which disulfide edge sites similarly catalyze hydrogen evolution reaction (HER). One such study reported a current density of -13 mA/cm² at -0.2 V vs RHE [4]. Without the edge site enhancement, another study reported results for MoS₂ of ca. -5 mA/cm² at -0.5 V vs RHE [2].

Use of TMDs as photocatalysts has been constrained by difficulty tuning the intrinsic optoelectronic excitation and damping mechanisms. Optical absorption in monolayer tungsten disulfide (WS₂), for example, is limited to >2 eV (<620 nm). This keeps broadband UV-vis absorption below ca. 2%. Moreover, the photoluminescence (PL) response of WS₂ is layer-number dependent [5]. Decoration of 1L-TMD by plasmonic metal nanoparticle (NP) has enabled broadband tunability of resultant photon-exciton interactions in catalytic hydrogen generation [6,7]. The enhanced photon-exciton interactions have been attributed to hot electron transfer (HET) from plasmonic NA to TMD substrate. HET, which occurs at interfaces between non-insulating media, e.g., TMD and NA, increases plasmon damping beyond radiative and nonradiative mechanisms [8]. Such damping has been reported to transfer plasmon energy to adjacent 1L semiconducting TMDs [9,10]. Moreover, NP decoration of WS₂ was reported to enhance PL 11-fold via exciton coupling with plasmonic electric near fields [11]. However, plasmonic HET had only been evaluated across interfaces of AuNP physically deposited on TMD; and characterization of HET from NA to WS₂ has been empirical, lacking a computational framework.

To examine effect of NA on WS₂ catalytic hydrogen evolution, Au was reduced onto edge sites of liquid phase exfoliated (LPE) 2H-WS₂ from an aqueous Au(III) solution based on a previously reported method [12,13]. Thiols dangling at WS₂ nanosheet edges underwent a redox reaction with AuCl₄⁻ to yield a covalent Au-S bond as confirmed by transmission electron microscopy (TEM; Figure 2) and X-ray photoelectron spectroscopy (XPS) [14]. In contrast, covalent bond formation was not observed when AuCl₄⁻ was reacted with chemical exfoliated 1T-WS₂. Oxidation of thiols to disulfides at edges of LPE-2H-WS₂ coincident with reduction of Au (III) to Au(0) was anticipated to increase catalytic HER in Au-decorated 2H-WS₂ [15,16]. Using LSV, AuNP-decorated 1L2H-WS₂ catalyst exhibited a 5-fold enhancement relative to undecorated monolayer and >10-fold increase relative to other reported TMD-based electrodes when exchange current density was normalized by film thickness [14]. For the comparison, equivalent mass of WS₂ per unit area electrode was used and mean flake length was maintained at < L > = 67 nm. All films exhibited 100 mV/decade slopes, consistent with reported nanolayered TMD HER electrocatalysts.

To quantitate HET from plasmonic NA to catalytic WS₂, discrete dipole approximation (DDA) of Maxwell’s equations using DDSCAT v7.3 [17] was conducted based on a previously reported method for 1L graphene [18] and MoS₂ [19]. Computed far field spectra of 1L-WS₂-NA heterostructures exhibited localized surface plasmon resonance (LSPR) and WS₂, excitonic transitions [20]. Decoration of 1LWS₂ by AuNA enhanced broadband extinction, particularly at the 620 nm A exciton red-shifted from the LSPR. Tuning the LSPR to specific excitonic transitions is anticipated to increase enhancement [21]. Edge decoration increased optical extinction efficiency relative to basal plane deposition. An 11±5% quantum efficiency of HET from ca. 20 nm AuNPs reduced on WS₂ nanosheets was estimated by comparing electron energy loss spectroscopy (EELS) measured and DDA simulation [22].

**Conclusion**

In summary, reduction of plasmonic NA onto 1L-2H-WS₂ edges formed covalent Au-sulfur bonds. Such 1L-WS₂-NA heterostructures increased catalytic HER relative to 1L-2H-WS₂ in physically decorated-NA or undecorated forms and enhanced plasmon damping attributable to direct electron transfer at the catalytic TMD-NA heterointerface. Application of this redox chemistry and integration of microscopic and spectroscopic measures and simulation enable improved photocatalysis using 1L-TMD-NA and implementation in a variety of chemical, biological, energy and water systems.

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Generation and Detection of Terahertz Radiation from Laser-Plasmas

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Abstract

Broadband terahertz (THz) radiation from laser-plasma interaction and a corresponding broadband THz time domain spectroscopy based on optically-air-biased-coherent-detection technique are presented. One of the single-shot detection techniques for THz time domain spectroscopy, the spectral-encoding technique, is also reviewed [1]. Distortions of the signals measured by this technique and the corresponding strategies to reduce them are demonstrated [2].

![Graphs showing THz waveforms and spectra](image)

Figure 1. (a) Attenuated and corrected THz waveforms using 10 Si wafers as attenuators vs. the original strong THz signal. (b) THz spectrum of the corrected THz signal with 10 Si wafers vs. that of the original one.

References


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Photonic crystal cavity optomechanics for the applications of low phase noise frequency source and high-performance sensing

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Abstract

Based on the air-slot line-defected photonic crystal nano-cavity [1], here we develop a monolithic integration of photonic crystal optomechanical oscillators and on-chip high speed Ge detectors (see Figure 1) by using the silicon CMOS platform. With the generations of both high harmonics (up to 59th order) and subharmonics (down to 1/4), due to strong mutual couplings between optomechanical self-sustained oscillation and self-pulsation oscillation, our chipset provides multiple low phase noise frequency tones [2] for applications in both frequency multipliers and dividers. The synchronization between two mechanical modes [3] and dynamical chaos [4] in the optomechanical cavity are reported as well. These characteristics enable optomechanical oscillators as a frequency reference platform for radio-frequency-photonic information processing.

Moreover, we further demonstrate a chip-scale optomechanical cavity with large mass (see Figure 2) which operates at ~77.7 kHz fundamentally and exhibits large optomechanical coupling of 44 GHz/nm. The mechanical shifting range of ~58 kHz and more than 100-order harmonics are obtained with which the free-running frequency instability is lower than 10^-6 at 0.1 sec integration time [5]. The optomechanical coupling strength can be mechanically controlled by taper fiber and the Drude self-pulsation plasma locking is also reported in this platform [6]. Such large mass optomechanical cavity can be applied for the sensing applications, such as accelerometers and magnetometers.

Figure 1. Fabricated optomechanical cavity integrated with on-chip Ge-detector

Figure 2. Fabricated large mass optomechanical cavity

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Hydration of various solutions observed by terahertz spectroscopy

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Abstract

Use of terahertz waves for sensing various type of materials has been an important subject of research. The target materials include biological tissues, various kind of liquids, industrial products, etc. In most of cases, water is the most important absorber of terahertz waves. It is crucial, therefore, to understand the terahertz absorption properties of water in various situations.

It is known that the terahertz absorption of water is strongly affected by the interaction with other molecules in solutions, gels, crystals, or other types of phases. In aqueous solutions, for example, the solute is stabilized by the interaction with surrounding water molecules, and the water molecules incorporated in this interaction are very often strongly bound to the solute, leading to reduced mobility and reduced terahertz absorption.

We have constructed a high-precision terahertz time-domain spectroscopy apparatus for hydration study. Results of hydration study of protein (hen egg white lysozyme) aqueous solution with several ammonium salts [1,2] is briefly described below. Terahertz absorption coefficient of the protein-salt aqueous solutions and salt aqueous solutions were obtained as a function of the salt concentration. The difference between the results of protein-salt aqueous solutions and those of salt aqueous solutions exhibits the nature of the hydration water of the protein and also the effect of salt on them. It was found that the hydration layer of the protein is affected by ions (mostly anions) added to the solution, and that so-called kosmotropic ions (structure makers), such as sulfate ion, reduce the amount of immobile hydration water, whereas chaotropic ions (structure breakers), such as thiocyanate ions, increase it. It is shown that the dynamical properties of hydration water molecules is correlated with the stability of the macromolecules, whereas their density with the solubility.

Measurements on other types of water-rich systems have shown that the mobility of water sometimes increases after hydration, or that the hydration number changes drastically upon a small change in the concentration. Study of hydration using terahertz waves is expected to produce fruitful results in various fields including solution chemistry, sol-gel science, biology, and medicine.

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Fast THz Modulator based on the stagger-netlike GaN HEMT active metamaterial

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Abstract

Terahertz technology promises unique applications in high speed communication, high accuracy imaging and so on [1]. However, one major bottleneck for developing Terahertz application systems is the lack of high-performance dynamic devices for effectively manipulating the Terahertz wave. In recent years, the rapid development of two-dimensional electron gas (2DEG) devices provides a promising way to develop dynamic terahertz devices [2]. Here, we combined a stagger-netlike metamaterial array with high-electron-mobility transistor (HEMT) structure to form a electronic grid-controlled THz modulator. By controlling the carrier concentration of 2DEG, the mode conversion between two kinds of dipolar resonances has been realized. Modulation depth of this device can reach up to 94%. More importantly, in the dynamic test, 600 MHz sinusoidal signals was received by a THz detector. It may provide a way to achieve effective active devices in THz wireless communication system.

Figure 1. (a) 3-D structure of a GaN HEMT metamaterial unit cell. (b) Schematic of the THz modulator. (c) Simulation transmission spectrum results with different carrier density. (d) Image of packaged THz modulator. (e) The received sinusoidal modulating signals.

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Terahertz filter and demultiplexer with photonic crystal waveguide

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Abstract

Terahertz (THz) wave is finding growing applications in various important fields such as space science, communications, and security screening [1]. Besides sources and detectors, development of THz technologies also requires devices to guide and manipulate THz waves. The demand for high performance quasi optic components such as frequency filters, demultiplexer, attenuators, splitters, and polarizers is increasing [2]. We theoretically propose and investigate a magnetically tunable narrow-band terahertz filter and a multi-channel THz wavelength division demultiplexer based on photonic crystal waveguide. The optical properties of the filter have been analyzed in detail. It is found that a single resonant peak with the central frequency of ~1 THz is existed in the transmission spectrum, which has a narrow full width at half maximum of <2 GHz. Moreover, under the control of an external magnetic field, transmission frequency and width of passband are adjustable, which reveals that the 2-D silicon photonic crystal waveguide with point and line defects can serve as a continuously tunable bandpass filter at the terahertz waveband. THz division demultiplexer consists of an input waveguide that perpendicularly coupled with a series of defects cavities, each of which captures the resonance frequency from the input waveguide. Coupled-mode theory and finite element method are used to analyze the transmission properties of the structure. It is found that the transmission wavelength centered around 1 THz can be adjusted by changing the geometrical parameters of defects cavities, which equals to THz waves generated by optical methods such as difference frequency generation and optical rectification.

References


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Abstract

Broadband terahertz absorbers have recently attracted considerable attention for their promising applications in terahertz trapping, sensing, imaging, and detecting. In this work, an efficient approach to achieve broadband terahertz absorber based on sinusoidally-patterned graphene with nearly 100% absorption is demonstrated [1]. As shown in Figure 1(a), the proposed absorber is composed of a net-shaped periodically sinusoidally-patterned graphene sheet on the top, a thin gating layer embedded dielectric spacer in the middle, and a metallic reflecting plate on the bottom. We assume the material of the dielectric spacer, the gating layer and metallic plate are the polyethylene cyclic olefin copolymer (Topas) with the permittivity of $\varepsilon_{rt} = 2.35$ [2], and the polysilicon with the permittivity of $\varepsilon_{rp} = 3$ [3], and gold, respectively. The surface conductivity of the single-layered graphene is obtained from the Kubo formula [4]. The polysilicon gating layer is placed beneath the graphene sheet to control the graphene conductivity via electrostatic doping of the graphene by applying a DC voltage $V_g$. By introducing such a unique gradient width modulation of the unit graphene sheet structure, the continuous plasmon resonances of the absorber can be excited, and over 65% normalized bandwidth of 90% terahertz absorvance can be achieved under normal incidence for both TE and TM polarizations. As one of the most exciting characteristics, the broadband absorption of this absorber are insensitive to the incident angles and the polarizations. The absorbance remains more than 70% even the incident angles reach 60° for both polarizations. Furthermore, compared to conventional multi-resonator or multi-layered structures, the continuous net-shaped single-layered graphene structure can greatly simplify the electrostatic gating structure in achieving flexible tunability. By controlling the chemical potential of the graphene, the peak absorbance can be continuously tuned from 14% to 100%, as shown in Figure 1(b). This work offers a new perspective on the design of graphene-based tunable terahertz broadband absorbers. The design scheme can be easily scaled to the infrared or visible regimes.

References


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Abstract

Terahertz quantum cascade lasers (THz-QCLs) have attracted much attention as possible carrier sources for ultra high-speed wireless communications in the future, which owes to their high output powers and the absence of relaxation oscillations in QCLs. We have recently reported a photogenerated carrier-based optical-to-THz modulation scheme for THz-QCLs [1]. In the presentation, the intensity and spectral changes in THz-QCLs induced by photo-injected carriers at low temperature will be discussed together with the relevant relaxation mechanisms for the injected carriers. Furthermore, to obtain a quantitative understanding of the rich phenomena by the optical injection, we developed a global simulation scheme applicable to a wide variety of optical excitation experiments on QCLs including strong excitation densities (Figure 1) [2]. With the most of internal parameters, e.g., electron temperatures, scattering times, gains, and waveguide loss, treated as spatiotemporal and carrier density-dependent, the rate equation was solved to obtain the number of photon in the cavity. The output power (converted from the number of photons using the time-varying reflectivity at the cavity facet) well reproduces the experimental results.

Figure 1. Model grid for the analysis of the optically excited QCL.

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THz surface Emission spectroscopy and applications

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Abstract

Terahertz (THz) wave, bridging electronics and photonics in the electromagnetic spectrum, features many exotic properties and promising applications. However, because of low THz emission efficiency, less sensitive detectors, and few manipulating devices, THz wave is still on the horizon for practical applications since 1980s. With the application of femtosecond laser, THz surface emission spectroscopy has also been developed to serve as a sensitive and contactless tool for the optoelectronic measurement of semiconductor surfaces and interfaces. When a femtosecond laser beam impinges on the semiconductor surface, photocarriers or photodipoles are excited, which then induce THz radiation with the mechanism of photoconductivity or optical rectification. As the THz surface emission is sensitive to the surfaces and interfaces, the modification of the semiconductor surface provides a significant strategy for the design and performance evaluation of many electronic and optoelectronic devices for THz applications. In this talk, we will discuss the THz radiation mechanism for traditional semiconductors by changing the crystal orientation, exciting laser intensity, surface condition, and so on [1-4]. We will also discuss THz radiation from two-dimensional layered semiconductors under linearly polarized femtosecond laser excitation.

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Tune the polarization of terahertz waves via subwavelength metallic gratings

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Abstract

Here we present our recent work on tuning the polarization of terahertz waves via subwavelength metallic gratings. Firstly, we have experimentally demonstrated a linear polarization rotator that is a three-layer metallic grating structure for manipulating the polarization of broadband terahertz waves. By mechanical rotations of the composite grating layers, this freely tunable device can rotate the polarization of a linearly polarized THz wave to any desired direction with high conversion efficiency [1]. Then we theoretically investigate the propagation of terahertz waves through a graphene-loaded metal grating under external magnetic field. It is found that resonant modes in the system can be converted between transverse-electric and transverse-magnetic polarizations due to Hall conductivity of graphene, as a consequence, asymmetric transmission of terahertz waves through this graphene-loaded metal grating is achieved, and it can be tuned by adjusting either the external magnetic field or the Fermi level of graphene [2]. These tunable terahertz devices have potential applications in various areas, such as material analysis, wireless communication, and terahertz imaging.

References


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Coherent and Continuous Terahertz Emitters from High-Tc Superconductor Mesa structures


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Abstract

After the discovery of high temperature Superconductor YBa2Cu3O7-δ with the superconducting transition temperature \(T_c=92\) K, which is well above \(77\) K, the boiling point of liquid nitrogen, this year has become a celebrative and a commemorative 30th anniversary in the history of superconductivity research. In addition to this memorable occasion, this years is another important anniversary that ten years have passed after the discovery of terahertz electromagnetic wave emission from the mesa structure of Bi2212 single crystals [1]. The important ingredients here is that the superconducting CuO2 double layers are built in a unit cell of the Bi2212 crystal as it is grown and are sandwiched by the insulating Bi2201 layers, forming a stack of intrinsic Josephson junctions (IJJs). Since the electronic structure as a result of this layered structure the superconducting as well as even normal state is highly two dimensional, the superconducting coupling becomes Josephson-junction-like, extremely weak as measured by the c-axis critical current \(J_c^c=10^6 - 10^7 \text{A/cm}^2\) compared with \(J_{cs}^c=10^6 - 10^7 \text{A/cm}^2\), resulting in the reduction of the superconducting plasma frequency to the level of \(f_\text{p} \sim 10^{11}\) c/s (0.2 ~ 1 meV), which is well below the superconducting gap \(\Delta \sim 30\) meV. This Josephson plasma mode can be excited by the dc-current through the nonlinear Josephson coupling effect by the \(dc\)-Josephson effect and the coherent THz emission is generated due to the \(ac\)-Josephson oscillation with the frequency \(f_e=(2e/h)v_e\), where \(e\) is the elementary charge of electron, \(h\) Planck constant, and \(v_e\) the voltage per intrinsic Josephson junctions, when it matches well the cavity mode frequency. Although the understanding of this phenomenon has already been well established by various experiments and theoretical works [2], the practical limitation of the device is not well understood yet. For example, the most important issue is on what determines the maximum power extracted from one intrinsic Josephson junction, and how much power can be generated from the actual mesa structure with \(N\)-intrinsic Josephson junctions, where \(N\) is the number of intrinsic Josephson junctions in a mesa. The next issue is on what limits the maximum frequency.

The essential parameter related to two issues has evidently been thought the thermal effect due to the Joule heating by the \(dc\)-current (10 - 50 mW), which produces heat of \(\sim \text{MW/cm}^2\) and naturally causes a serious temperature increase and inhomogeneity, which is often called as a hot-spot. It has been disclosed by us that the formation of the hot-spot gives only a detrimental effect on the THz radiation phenomena so that it is better to avoid it or control it to make the influence minimum on the THz emission. The heating issue has recently been studied intensively [3]. As a result, we have achieved a frequency of 2.4 THz with a power of 30 \(\mu\text{W}\) [4] quite reproducibly.

Recently, we have done a systematic case study on the rectangular, square, circular, triangular mesas, etc. and found an interesting fact. That is concerning the missing modes, which are expected to be as the strong emission modes but systematically disappear or are missing, perhaps, due to very weak intensities. This seems to occur especially in the degenerated symmetric cases such as in square and circular mesas. We argue this effect as a mode cancellation in the degenerated cavity modes in a symmetric mesa.

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Terahertz-wave parametric wavelength conversion at room temperature

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Abstract

Over the past decade, there has been remarkable growth in the field of terahertz frequency science and engineering, which has become a vibrant, international, cross-disciplinary research activity. Wavelength conversion in nonlinear optical materials is an effective method for generating and detecting coherent terahertz waves owing to the high conversion efficiency, bandwidth, wide tunability, and room-temperature operation, and if the tuning range and the peak power can be enhanced, drastic developments in basic researches and industrial applications can be expected.

Here we demonstrate the generation of high-brightness terahertz waves using parametric wavelength conversion in a nonlinear MgO doped LiNbO3 crystal. We revealed novel parametric wavelength conversion process using stimulated Raman scattering in MgO:LiNbO3 without stimulated Brillouin scattering using recently-developed microchip Nd:YAG laser. We also demonstrated the coherent detection of generated terahertz waves using nonlinear up-conversion.

A number of applications require high brightness, that is, intense and narrowband, terahertz waves such as observing multi-photon absorption to specific excitation states. We speculate that the high-brightness terahertz wave and its visualization could be powerful tools not only for solving real world problems but also fundamental physics. We expect that these methods will open up new fields and tune up killer applications.

Acknowledgements

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Development of Small Size Terahertz Vacuum Sources in IECAS

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Abstract
Terahertz frequencies are among the most under developed electromagnetic spectra, even though their potential applications are promising in biochemical sensing, imaging for medical and security applications, astrophysics and remote atmospheric monitoring, and high-bandwidth communications. Among their wide applications, the lightweight, low voltage and broadband sources of high-power coherent THz radiation are important for military radar, electronic countermeasure systems and communications. Vacuum electronic devices such as the traveling wave tube amplifier (TWTA), backward wave oscillator and extended interaction oscillator show great potential for applications at the frequency because of wide band and high power. At THz band, the most critical part of small size THz vacuum sources is the slow wave structure, which determines the output performances, such as the output power and signal band. In the millimeter and sub-millimeter wave vacuum devices, the various of SWSs including coupling cavity, disk-loaded, helix and complex CC are used. However, they are not suitable for the THz band, resulting from the difficulty of fabricating and integrating.

Folded waveguide (FW) or serpentine circuits, for example, have ~10:1 aspect ratio in the waveguide height dimension. This is much larger than the ideal for both machining, which is limited by the length of the tool shank with acceptable wobble; and photolithography, which is limited by defocusing and absorption of light as it penetrates into deep photoresist. These requirements are the topic of ongoing research in the vacuum electronics community, motivated by the continued interest in high power mm-THz sources. Consequently, there is a great interest in the study and development of FW SWS. For the satisfied with requirement of high speed communication and ViSAR, the TWT has been developed. Based on the FW, the small size THz vacuum devices are developed in China, including UESTC, IECAS CETC 12 and CAEP, etc. In this talk, we report the development of small size THz vacuum devices in IECAS.

At IECAS, three kinds THz devices have been developed, such as the 0.3THz EIO, the peak power exceeds 4W. The 0.1THz EIO generates the output power is larger than 12W. Moreover, the THz TWT has been developed through the small size electron gun, high strength focusing magnetic field and high frequency structure welding. It produces the output power exceeds 2W and the band width is larger than 5GHz. Now we are developing the continued wave TWT for the power exceeds 10W and signal band width larger than 5GHz.

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300GHz wireless link with a CMOS transceiver

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Abstract

Since terahertz provides a wide frequency band, terahertz communication realizes a data rate exceeding 100 Gbps approaching fibre-optic speed. The frequency band from 252 to 275 GHz has already been allocated for communication. Further discussion is being made to use the frequency band exceeding 275 GHz, which has not been assigned yet, for communication use. On the other hand, since terahertz has large atmospheric attenuation and strong directivity, it is limited to short-distance fixed radio communication. However, it is long-distance and mobile application that is intrinsically expected for wireless communication. In this talk, even in terahertz, it is shown that kilometre communication is potentially possible by selecting the frequency appropriately. It is also shown that terahertz communication can be performed using CMOS process, which was said to have inferior high frequency characteristics to compound semiconductors. How will the world change when technologies beyond such conventional common sense are established? The impact of terahertz communication and the contribution of CMOS transceivers are discussed.

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Ultra-broadband Wireless Communications in the 0.1-10 Terahertz Band

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Abstract

By supporting tens or hundreds GHz bandwidths, Terahertz (THz)-band (0.1-10 THz) communication is envisioned as a key wireless technology of the next decade. The THz band will help overcome the spectrum scarcity problems and capacity limitations of current wireless networks, by providing an unprecedentedly large bandwidth. In addition, THz-band communication will enable a plethora of long-awaited applications, both at the nano-scale and at the macro-scale, ranging from wireless massive-core computing architectures and instantaneous data transfer among non-invasive nano-devices, to ultra-high-definition content streaming among mobile devices and wireless high-bandwidth secure communications.

In this invited speech, an overview of THz-band communications will be provided. First, the current progress and open research directions in terms of THz-band channel modeling will be presented. The main phenomena affecting the propagation of THz signals will be explained and their impact on the channel capacity will be assessed. Second, novel communication mechanisms such as the modulation techniques, resource allocation, timing acquisition schemes, and Ultra-Massive Multiple-Input Multiple-Output (UM-MIMO) will be presented. Finally, the state of the art and open challenges in the network layer design and other relevant research directions will be stated. This presentation is expected to provide the audience with the necessary knowledge to work in a cutting-edge research field, at the intersection of antennas and propagation, and information and communication technologies.

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Millimeter- and terahertz-wave technology for communication and radar/imaging applications by photonics technology

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Abstract

Advanced optical fiber communication technologies enable low-loss and broad-bandwidth transmission in the millimeter-wave and terahertz-wave bands via an optical fiber network. Precise optical modulation techniques can directly generate the millimeter-wave signals, and finally, an optical frequency comb signal generated by the modulation also provides the terahertz-wave signals by optical heterodyning systems. These technologies on the signal generation could be utilized for realization of distributed antenna system in the millimeter- and terahertz-wave bands based on the optical fiber network for some specific applications: foreign object debris detection systems for airport runway surveillance and high-speed railway radiocommunication systems between train and trackside. In the talk, we briefly introduce the R&D activities on the millimeter- and terahertz-wave for both wireless communication and radar/non-destructive imaging systems based on photonics.

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The fascinating split-ring-resonators: progress in design, fabrication and applications of terahertz metamaterials

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Abstract

Metamaterials with many unique optical properties are made of periodically arranged sub-wavelength metallic structures that are able to couple to external electromagnetic (EM) waves. One of such structures is the commonly used split-ring-resonators (SRR). In this talk, I will discuss and demonstrate some new progress we made on SRR-based terahertz metamaterials. By looking into the coupling between SRRs and the effect of incident polarization, we proposed a way to continuously modulate their resonances, changing the transmission intensity at resonant frequency from 20% to 80% [1]. We also designed a SRR-based polarization-insensitive broadband filter in THz range and discovered its effect in eliminating asymmetric characteristics in device structure [2]. A stop band with bandwidth of as large as 1.40 THz was achieved. To improve the fabrication process, a facile metal transfer method was employed to create SRR patterns on PDMS surface, planar and otherwise, as well as PDMS-coated surfaces, such as paper, fabric and leaf [3]. Lastly, a design of 3D THz metamaterial device was proposed to be used in biomedical field for cancer cell studies, exploiting its structural similarities with the single-cell-capturing microfluidic devices.

Acknowledgements

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Micro-fabricated L-shape metasurface terahertz biosensor

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Abstract

In recent years, more and more attention has been paid on the research and development of terahertz (THz) technologies all over the world. Biosensing is one of the most important applications of THz waves. Metasurface is an artificial two-dimensional material with flexible structures and good electromagnetic wave control capability, which is very suitable for THz wave control and sensing. In this work, a L-shape metasurface THz biosensor was designed, simulated, optimized, micro-fabricated, and finally tested by the time-domain spectroscopy (TDS) system. The simulation and experimental results agree with each other very well. The conclusions show great advantage of metasurface for the THz biosensing application, especially when it is combined with the microfluidic technology in the near future.

Figure 1. Structure of sensing array

Figure 2. Measured results of some liquid samples

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Design and electrochemical performance of nano-micro structured porous materials

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Abstract

Graphene and carbon nanotube present unique structure and excellent properties, such as high specific surface area, high conductivity, high thermal conductivity, etc., and play an increasingly important role in the electrochemical energy storage and conversion. The doping of nano-carbon with nitrogen and boron can modulate the electron and energy band structure, and further improve its physical and chemical properties. The pore structure, pore size distribution and wall thickness of the electroactive material display important effect on the electrolyte infiltration, the ion transport and adsorption, and the overall performance of the battery. The use of doped nano-carbon and porous composite structure can improve the conductivity and the electrochemical surface area and reaction sites of the electroactive materials, and significantly increase the efficiency of energy storage and conversion.

This presentation focuses on some progresses of the doped nano-carbon and porous composite structures, including the following three aspects: (1) The controllable synthesis and electrochemical lithium insertion characteristics of the porous electrode materials. Porous materials have been widely used in a variety of fields, but their large-scale controllable synthesis is still a considerable challenge. We have designed a novel templated freeze-drying method to conveniently control the porous properties of materials, which has been successfully applied to the synthesis of various porous phosphates, oxides and composites, and the electrochemical lithium storage properties have been explored. (2) The porous lithium iron phosphate and nitrogen-doped graphene composite. Three-dimensional porous microspheres composed of LiFePO$_4$ and nitrogen-doped graphene have been synthesized by a solvothermal method. The effect of graphene doping on the nucleation and growth of LiFePO$_4$ and the influence of the unique self-assembled porous microsphere structure on the electrochemical lithium insertion performance have been studied. (3) Supercapacitance properties of doped graphene. A series of dopant graphene materials have been facilely synthesized by a thermal solid state reaction. The regulation of doping configuration on the electronic structure of graphene and the influence on the supercapacitance have been systematically studied. The relevant mechanisms have been proposed and some phenomena in the literature have been reasonably explained.

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Molecular Insights into Electrical Double Layers in Graphene-Based Supercapacitors

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Abstract

Recently nano-structural carbons have become the most widely used electrode materials in supercapacitor community, because of their high specific surface area, good electrical conductivity, chemical stability in a variety of electrolytes, and relatively low cost. In particular, graphene-based carbons are emerging as an auspicious candidate due to the unique feature of graphene. Among electrolytes used for supercapacitors, ionic liquids (ILs) have been becoming a promising class of them, owing to their exceptionally wide electrochemical stability window, excellent thermal stability, non-volatility and relatively inert nature. Despite considerable work on supercapacitor with graphene-based carbon as electrodes, the details of what happens under nano-confinement, including pores, still require in-depth exploration especially for IL electrolytes.

We studied the interfacial phenomena occurring between ILs and graphene-based electrodes in supercapacitors, using the combined molecular dynamics (MD) simulation by modeling ILs-based EDLs at planar, cylindrical, spherical electrode surfaces and inside electrode pores at nano/micro-scale. This talk would include:

1) MD modeling on ILs-based EDLs at open surfaces (e.g., planar, cylindrical, spherical, with defects, etc.) [1-2] and the integration with experiments (e.g., atomic force microscopy, AFM) [3-4], which would focus on the EDL structure and its influence from ion size, ion type, applied potential, electrode curvature, etc.

2) MD modeling on ILs-based porous carbon supercapacitors [5-7], which would embody the pore size effects on capacitance, the ion dynamics under porous confinement, and pore expansion during charging.

3) The anatomy of electrosorption for water in ionic liquids at electrified interfaces [8], which would show, for the first time, the work on the adsorption of water on electrode surfaces in contact with humid ILs.

References


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Controllable Synthesis of N-Doped and Dually Doped Mesoporous Carbons for Adsorption and Catalysis Applications

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Abstract

Mesoporous carbon materials are being in vogue because of their intriguing properties and wide potentials. Doping of heteroatoms, especially N, in carbons have attracted enormous interests owing to its capability in enhancing or expanding their applicability in separation, energy conversion and catalysis. In addition, N-doping can further boost dually doped carbons, such as with S and metal as the second dopant. Such a dual doping could possibly optimize material property and maximize performance through synergistic effects. In this talk, several synthetic methods, such as post modification [1-2], one-step solvent-free nano-confining synthesis [3-4] and spray-drying-assisted assembly [5], for the synthesis of heteroatom (singly or dually) porous carbon materials with different porosities and structures will be introduced and discussed. Furthermore, the controllable synthesis of metal/heteroatom dually doped mesoporous carbons with desirable fascinating properties will be introduced. The demo-applications of these materials in typical adsorption, such as arsenic removal and CO$_2$ capture, and typical catalysis, such as oxygen reduction, biodiesel production and dehydrogenation/hydrogenation coupling reactions, will be presented and their structure-performance correlations will be discussed.

References


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Growth of TMDC Nanostructures by Chemical Vapor Deposition

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Abstract

Chemical vapor deposition (CVD) is a simple but powerful technique to synthesize thin films of various materials. It can also be used to synthesize nano-structured materials if it is used with nano-structured templates or catalysts. We have developed separate flow CVD system to make multilayered transition metal dichalcogenides (TMDCs) or doped layers [1,2]. We used the apparatus to make nanocomposite between carbon nanotube and MoS$_2$ that was applied to photovoltaic applications (Figure 1) [3]. We also describe the search for the catalysts to make TMDC nanostructures.

References


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Electrocatalytic hydrogen evolution reaction of MX$_2$ and MX$_2$ heterostructures

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Abstract

Advanced materials for electrocatalytic and photoelectrochemical water splitting are central to the area of renewable energy. Recently, two dimensional layered materials of MX$_2$ (M: Mo, W; X:S, Se, etc.) have emerged as a new kind of catalysts for such applications. Our group have reported the direct synthesis of high-quality, domain size tunable, strictly monolayer MoS$_2$ flakes on commercially available Au foils by a chemical vapor deposition (CVD) method. The nano-sized triangular MoS$_2$ flakes on Au foils are proven to be excellent electrocatalysts for hydrogen evolution reaction (HER), featured by a rather low Tafel slope (61 mV/dec) and a relative high exchange current density (38.1 μA/cm$^2$). The excellent electron coupling between MoS$_2$ and Au foils is considered to account for the extraordinary HER activity [1]. Furthermore, via a facile all-CVD approach, we have also demonstrated the direct growth of monolayer MoS$_2$ on graphene (MoS$_2$/Gr) over Au foils [2,3]. A dramatic decrease of the bandgap from ~2.20 to ~0.30 eV was detected at the domain edge of MoS$_2$ within a lateral distance of ~6 nm, as evidenced by STM/STS observations. The edges of monolayer MoS$_2$ nano-sheets were thus served as narrow-gap quantum wires, which can greatly facilitate the electrocatalytic property of MoS$_2$ in HER [4]. Meanwhile, we also synthesized either MoS$_2$/WS$_2$ or WS$_2$/MoS$_2$ vertical heterostructures on Au foils by a growth-temperature-mediated, selective two-step CVD strategy. Relative enhancement or reduction in the photocatalytic activities were observed for MoS$_2$/WS$_2$ and WS$_2$/MoS$_2$ in HER under illumination, respectively. This is explained from the type-II band alignment of the MoS$_2$/WS$_2$ stack that enables effective electron-hole separation and fast electron transfer kinetics, as well as directional electron flow from electrode to catalytically active sites [5]. The abovementioned efforts are expected to establish the internal relationship between the metallic edge states of MoS$_2$ and its HER performances, as well as the advantage of MX$_2$/MX$_2$ vertical stacks in photocatalytic HER applications.

References

Application of transition-metal dichalcogenides beyond general electronics

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Abstract

Transition-metal dichalcogenides (TMDs) are novel layered materials for various kinds of application. In particular, those formed in hexagonal prismatic structure (Figure 1a) with group-VIB transition-metal are semiconductors in nature and show good transistor performance and strong light-matter interaction. However, the potential application of group-VIB TMDs is not only limited to such general electronics and optics, but also covers next-generation electronics of spintronics and valleytronics including the coupling to the optical polarization (Figure 1b) [1]. Owing to the lack of the inversion centre in the individual layer, the six conduction band minima and valence band maxima at the edge of the hexagonal Brillouin zone split into two groups, creating a valley degree of freedom. Optical interband transition at these high symmetry points are further coupled to the helicity of light. In addition, the heavy transition-metal elements leads to a large spin-orbit interaction and a consequent spin splitting.

Here, I will report our recent research aiming at next-generation electronics, including optoelectronic device utilizing valley degree of freedom [2,3] and the fundamental investigation of the spin relaxation in TMDs [4].

References


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Prelithiated Si nanoparticles-carbon nanotubes composite anodes for Li-ion batteries

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Abstract

Freestanding flexible Si nanoparticles-multi-walled carbon nanotubes (SiNPs-MWNTs) composite paper anodes for Li-ion batteries (LIBs) have been prepared using a combination of ultra-sonication and pressure filtration. No conductive additive, binder or metal current collector is used. The SiNPs-MWNTs composite electrode material achieves first cycle specific discharge and charge capacities of 2298 and 1492 mAh/g, respectively. To address the first cycle irreversibility, stabilized Li metal powder (SLMP) has been utilized to pre-lithiate the composite anodes. As a result, the first cycle irreversible capacity loss is reduced from 806 to 28 mAh/g and the first cycle coulombic efficiency is increased from 65% to 98%. The relationship between different SLMP loadings and cell performance has been established to understand the pre-lithiation process of SLMP and to optimize the construction of Si-based cells. A cell containing the pre-lithiated anode is able to deliver charge capacity over 800 mAh/g without undergoing the initial discharge process, which enables the exploration of novel cathode materials.

It was also found out the SiNPs-MWNTs electrode with 3:2 Si/MWNT ratio exhibits the optimal balance between the high capacity of SiNPs and the high electrical conductivity and structural stabilization quality of MWNTs, leading to a high rate capability, high specific capacity, and cycle life surpassing the conventional slurry-cast SiNPs electrode using binder and Cu current collector. The reversible capacity is 1866 mAh/g (based on the total composite weight, the same below) at current density of 100 mA/g. After 100 cycles, the electrode retains capacity of 1170 mAh/g at 100 mA/g and 750 mAh/g at 500 mA/g. The superior performance is believed to be due to the cooperative or even synergistic effect achieved by the optimal combination. Furthermore, the freestanding feature of our electrode eliminates the non-active mass, which is promising for enhanced capacity and energy density of Li-ion cells.

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Investigating structure and stress evolution in Si anode using in situ high pressure technique and Raman microscopy

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Abstract

Silicon is widely regarded as one of the most promising anode materials for next-generation lithium-ion batteries, making Li-Si an important energy storage system. During Li insertion into Si, stress of gigapascal level is introduced accompanied by a volume expansion up to 280%, alters properties of materials, and leads to mechanical failure of Si anodes.

\( \text{Li}_{15}\text{Si}_4 \) (alpha- \( \text{Li}_{15}\text{Si}_4 \), space group: I-43d), the only crystalline phase that forms during lithiation of the Si anode in lithium-ion batteries, was found to undergo a structural transition to a new phase (beta- \( \text{Li}_{15}\text{Si}_4 \)) at approximately 7 GPa (see Figure 1). Ab initio evolutionary metadynamics calculations suggest beta-\( \text{Li}_{15}\text{Si}_4 \) has an orthorhombic structure with an Fdd2 space group. This new beta- \( \text{Li}_{15}\text{Si}_4 \) has substantially larger elastic moduli compared with alpha- \( \text{Li}_{15}\text{Si}_4 \), and has good electrical conductivity. As a result, beta- \( \text{Li}_{15}\text{Si}_4 \) has superior resistance to deformation and fracture under stress. The theoretical volume expansion of Si would decrease 25% if it transformed to beta- \( \text{Li}_{15}\text{Si}_4 \), instead of alpha- \( \text{Li}_{15}\text{Si}_4 \), during lithiation. In addition, the fact that beta- \( \text{Li}_{15}\text{Si}_4 \) can be recovered back to ambient pressure, provides opportunities to further investigate its properties and potential applications [1].

Nanostructured Si are important materials to address mechanical stress issues in batteries although their stress were only calculated and no experimental data are available. Using in situ Raman microscopy to monitor the shift of the first-order Raman peak of Si, we were able to measure for the first time the lithiation-induced stress in Si nanoparticles. The shift of Raman peak of Si under hydrostatic stress was calibrated via an in situ high pressure Raman experiment. We observed a transition in the stress in Si core of nanoparticles during lithiation, from tensile to compressive (see Figure 2). At the beginning of lithiation, the reduction of the native oxide surface layer of the Si particle results in a tensile stress of approximately 0.2 GPa in Si. During the formation of amorphous Li\( \text{Si}_x \) in the outer layer of the nanoparticles, an increasing compressive stress up to 0.3 GPa is introduced to the Si core. This evolving stress explains the cracks that developed in the amorphous Li\( \text{Si}_x \) layer during lithiation of the Si nanoparticles, and is also consistent with modeling results. These results improve our understanding of lithiation-induced stress in nanostructured Si anodes, and provide valuable information for their theoretical study and future design [2].

Figure 1. The atomic structure of alpha-\( \text{Li}_{15}\text{Si}_4 \) (lower left) and beta-\( \text{Li}_{15}\text{Si}_4 \) (upper right). Each silicon atom is surrounded by lithium atoms (green spheres) with the coordination number of 12 (shown in blue polyhedra) or 13 (shown in pink polyhedra).

Figure 2. The stress in Si nanoparticles (left y-axis) and the potential of the half cell versus Li\(^+\)/Li (right y-axis) as a function of lithiation duration.
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Abstract

Developing an anode material with high-energy capacity is crucial to improve the performance of rechargeable batteries. Si is an important anode candidate because of its large lithium storage capacity (4200 mAhg\(^{-1}\), about 10 times higher than graphite, 372 mAhg\(^{-1}\)), low lithium alloying/dealloying potential, long discharge plateau, and abundance. However, the huge volume expansion/shrinkage of Si particles during charge/discharge cycles results in the pulverization of electrode structure and causes considerable capacity decay. The lack of electronic contacts between Si particles or between active coating materials and current collectors reduces its capacity. Many investigations have been carried out to accommodate this severe volume expansion including novel nanostructured Si, such as Si wires, Si tubes, porous thin films, and nest-like Si nanospheres, and multiphase composites with active Si and other active/inactive phases. However, the nano-sized Si particles would agglomerate during cycling which will decrease the cycling performance, due to high electrochemical activity. In this presentation, micro-sized polycrystalline Si particles have reduced agglomeration effect compared to nano-sized ones. We find that the electrochemical performances can be improved greatly by introducing proper binder to protect the electrode from cracking and using suitable conductive agent to provide an efficient conductive channel. By introducing nano-porous structures through chemical etching, the huge volume expansion can be effectively relieved by maintaining the complete structure of Si particles during charge/discharge cycles. Moreover, combined with carbon coating which is attractive owing to the conductive and ductile features of carbon, the conductivity can be improved and the volume expansion/shrinkage of Si can be further decreased. Developing nano-porous structure combining with pyrolyzed polyacrylonitrile could form a novel composite electrode. The electrochemical performance can be considerably improved with a conductive carbon network and nano-porous structure together with carbon layer accommodate the volume expansion/shrinkage during charge/discharge cycles.

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Efficient Solar-Rechargeable Lithium Ion Battery Energy Storage

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Abstract

Use of energy storage devices such as lithium ion batteries (LIBs) can help to mitigate the problem of intermittent photovoltaic (PV) power to achieve higher PV penetration into the electric grid. Also, the benefit of deployment of electric vehicles for energy sustainable future seems rather irrelevant unless they are charged using electricity generated from renewables. In addition, large scale practical applications of battery based electric vehicles is still challenging because of the inflexibility it has with the charging stations. All these issues can be addressed by use of solar cells as a viable energy source to charge lithium ion batteries. Here we demonstrate simple, efficient and cost effective photo-charging design approach where the use of promising low cost solar cells such as perovskite solar cell or dye sensitized solar cell with the help of DC-DC power conversion can efficiently charge a Li$_4$Ti$_5$O$_{12}$-LiCoO$_2$ LIB.

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High performances flexible energy storage devices

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Abstract

In this work, the flexible fiber-shaped supercapacitors have been fabricated by using the conductive polymers and graphene as the electrodes. The prepared fiber-shaped electrodes show good mechanical properties and excellent electrochemical performances. They can be easily knotted, twisted and woven into different shapes without sacrificing their electrochemical properties [1-2].

References


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Figure 1. Schematic illustration and the SEM images of HCFs and formation of hollow structures.
High Temperature Flexible Supercapacitors Using Graphene Electrodes

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Abstract

With increasing demand for high performance energy storage systems, the feasibility of reliable and functional energy storage devices that well operates under extreme conditions is of prime importance for expanding applicative fields as well as for understanding materials’ intrinsic and extrinsic properties and device physics. Our group has been investigating the control in the physical structure and chemical composition of 2D graphenes and beyond for ultracapacitive energy storage devices under limited circumstances, where conditions are classified into thermodynamic (e.g. pressure, volume and temperature) and kinetic (e.g. high rate and frequency) variables. We also studied a fundamental foundation via in-situ spectroscopic techniques to understand charge storage phenomenon of new materials and devices occurring on a nanoscale under various circumstances. In this talk, I will introduce high temperature operating, flexible supercapacitors based on graphene electrodes that can efficiently deliver electrical energy under electrochemical, mechanical and thermal stresses [1-4]. In order to achieve high performance supercapacitor devices under thermal, mechanical and electrochemical stresses, the micro- and macroscopic structures and chemical compositions of graphenes are delicately controlled by chemical modification. A new generation of flexible supercapacitors, with the long-term durability and outstanding electrochemical properties, were realized, showing a high position of the Ragone plot, even under severe conditions.

References


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Laser direct writing of high-performance micro-supercapacitors on graphene oxide and polymer films

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Abstract

The rapid development of portable miniaturized electronics has improved the research demand for compact energy storage components with high energy and power densities. In recent several years, a new type on-chip energy storage unit, which is called micro-supercapacitor (MSC) with two interdigitated electrodes in the same plane, has attracted much research attention, because MSCs possess not only the advantages of the supercapacitors, such as high power densities, robust cycle performance, pollution-free operation, and maintenance-free features, but also the small size, light weight, and flexibility, as well as the simplified packaging processes and compatibility to the integrated circuits. However, the materials and fabrication methods should be cost-effective, scalable, and compatible with current electronic industries. Carbon materials, especially graphene, which possess high specific surface areas, electrochemical stability, high conductivity, and high mechanical tolerance, can meet the requirement for energy storage unit in flexible wearable devices. Laser induced carbonization from polymers and laser induced reduction of graphene oxide have been reported for the preparation of MSCs due to the high power at the focused area. Compared to the commonly used printing and lithographic techniques, laser direct writing is a non-contact fast single-step fabrication technique with no need for masks, post-processing, and complex clean environments. Moreover, the laser direct writing method has the potential to be integrated to current product lines for commercial use. Therefore, it is necessary and important to study the preparation of high-performance micro-supercapacitors on graphene oxide or polymer films by laser direct writing technique.

In this paper, we will introduce our recent studies on the laser direct writing of high-performance micro-supercapacitors on polyimide and graphene oxide films, which consists of the following several parts: the preparation of carbon MSCs by laser direct writing on polyimide films in air using a continuous-wave blue-violet semiconductor laser (Figure 1) [1]; the improvement of the carbon MSCs by laser direct writing in Ar [2]; the carbon/Au MSC with high-rate charge-discharge capacitive performance [3]; and electrolyte-free high-performance reduced graphene oxide (RGO)-GO-RGO MSCs prepared by laser direct writing on graphene oxide films (Figure 2) [4].

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Rational Design of Novel Carbon Catalysts for Clean Energy Conversion and Storage

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Abstract

In fuel cells and metal-air batteries, there are critical chemical reactions: oxygen reduction reaction (ORR), and oxygen evolution reaction (OER), respectively. These reactions, however, are sluggish and require noble metals (e.g., platinum) or their oxides as catalysts. The scarcity and high cost of noble metals have hampered the commercial applications of these technologies [1]. Therefore, it is necessary to search for alternative materials to replace Pt. Carbon nanomaterials, such as carbon nanotubes (CNTs) and graphene, are appealing as an alternative for metal-free catalytic applications because of their structures and excellent properties. Although the superior catalytic capabilities of heteroatom-doped carbon nanomaterials for ORR have been demonstrated, trial-and-error approaches are still used to date for the development of highly-efficient catalysts. To rationally design a catalyst, it is critical to correlate intrinsic material characteristics with catalytic activities. Through first-principles calculations, we have identified a material property that serves as the activity descriptor for both ORR and OER, and established a volcano relationship between the descriptor and the catalytic activities of the carbon-based nanomaterials [2]. The design principles can be used as a guidance to develop various new carbon-based materials for clean energy conversion and storage.

Figure 1. Volcano relationships between the descriptor and the catalytic activities of the carbon-based nanomaterials.

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Abstract

The exploitation of high-performance lithium ion batteries is an effective way to promote the practicality of electric vehicles and the large scale development of renewable energy. We have designed and used a low temperature solution approach which is simple, low cost, scalable to synthesize Zn₄GeO₅@g-C₃N₄ hybrid structure. Furthermore the synergistic effect on their lithium storage has been discussed. The Zn₄GeO₅@g-C₃N₄ hybrids exhibited highly reversible capacity of 1370 mA h g⁻¹ at 200 mA g⁻¹ after 140 cycles and excellent rate capability of 950 mA h g⁻¹ at 2000 mA g⁻¹. On the other hand, molybdenum sulfide, one of the transition metal sulfides, has been considered as a hopeful anode material, because of the small volume change (~103%) and the high theoretical capacity (669–1675 mA h g⁻¹). In order to improve the electrochemical properties of molybdenum sulfide, we have designed and prepared TiO₂@MoS₂ core-shell structure. This composite structure provides a plenty of surface active sites for rapid transportation of lithium ions and orderly path for electrons. As a result, the electrochemical performance of TiO₂@MoS₂ is much higher than that of molybdenum sulfide as well as titanium dioxide. In addition, transition metal oxides also have been attracted widespread attention, because of it versatile nanostructures, high theoretical capacity and small volume change. We have synthesized 3D NiO microsphere architecture assembled from porous nanosheets via easy hydrothermal method. The advantage of large specific surface area endows the as-prepared 3D NiO microspheres with a good performance of stable and high reversible discharge capacity up to 820 mA h g⁻¹ even after 100 cycles at a current density of 100 mA g⁻¹, and good rate capability of 634 mA h g⁻¹ at a high current density of 1 A g⁻¹.

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Functionalized graphene with both physical and chemical adsorptions of charges for high-performance supercapacitors

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Abstract

Carbon-based supercapacitor is also called electric double-layer capacitor that store energy via physical adsorption and desorption of ions from the electrolyte. Pseudocapacitors based on metal oxides or conductive polymers store energy via a redox process and generally have higher specific capacitance compared to the EDL type. Graphene has been regarded as an ideal candidate for supercapacitor applications, while the specific capacitance achieved so far in the lab, normally 200-300 F/g, is much lower than its theoretical value of 550 F/g. In order to enhance the charge storage capacity of graphene, we functionalized reduced graphene oxide by N-doping and adsorption of small molecules of hydrolysized polyimide (PI). In N-doped graphene, the N-O bonds are responsible for the enhanced capacitance owing to their pseudo capacitive property [1]. Further, we found that the hydrolysis of PI can release small molecules into water solution, and these aromatic molecules adsorbed onto graphene via π-π interaction have a significant effect in increasing the capacitance. With merely 3% weight increase after adsorption, the specific capacitance is about 40% increased. High capacitance over 420 F/g can be easily achieved from the functionalized graphene electrode in H₂SO₄ aqueous electrolyte, even the electrode has high mass loading around 5 mg/cm². In Li₂SO₄ aqueous electrolyte that can extend the operation voltage window to 1.6 V, the specific capacitance also remains high around 400 F.

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How to improve the hydrogen storage of graphene based systems?

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Abstract

In the last two decades, the significant efforts have been made to develop alternative energy sources instead of fossil fuels because of increasing CO₂ emissions and the environmental impacts. Besides; hydrogen has been concerned to be an ideal clean energy carrier among the other renewable energy sources because of its environmental friendliness. However, some challenges have to be addressed before hydrogen will become a conventional and commonly available energy carrier. Carbon-based materials such as graphene and carbon nanotubes have been designed for hydrogen storage due to their large surface area, light weight, and tunable properties. Recently, we proposed a new strategy in which we considered three pure transition metal (TM) atoms or/and a combination of two TM atoms and one alkali earth metal atom (AEM) with high, medium and low hydrogen adsorption energies. These different metal atoms are used to decorate the Boron doped graphene sheet (BDG) and investigated their performance towards hydrogen storage capacity through spillover mechanism using first-principles calculations. Our results indicate that that the activation energies for H atom diffusion are much smaller, indicating that a fast H diffusion on this proposed surface can be achieved. These TM and AEM atoms decorated BDG surface can have the maximum hydrogen gravimetric capacity of 6.4% for double-sided adsorptions. To achieve higher gravimetric density, we also considered the Boron and Nitrogen co-doped graphene surface (BNDG) because B–N pair is isoelectronic to the C–C pair. However, controlling the binding strength of considered metal atoms with that of the BNDG surface is an important issue in the application of hydrogen storage. The recent studies have shown that the binding strength between the metal atom and the substrate can be controlled by means of applying an external electric field. Thus, the effects of the external electric field, as well as the effects of applying point charges on the designed medium towards its hydrogen storage capacity, will be discussed.

References


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Nano-Micro Conference

Abstract

Ternary Nanostructured Photocatalysts for Photoelectrochemical Water Splitting

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Abstract

The world’s population is growing continuously and in close proportionality to the demand on fossil fuels, which may lead to its depletion in the coming decades. The ability to harness energy from renewable resources is highly desirable for sustainable energy economy. In recent years, photoelectrochemical (PEC) water splitting process based on Earth-abundant semiconductor photocatalysts has gained considerable research interests for resolving energy and environmental issues. The PEC watersplitting that mimics the natural photosynthesis process can convert solar energy into storable form of hydrogen (H₂) energy, which is a good energy vector to meet the escalating energy demand. To date, however, almost all singular semiconductor photocatalysts used demonstrated poor PEC performance for solar-to-H₂ energy conversion. In this study, two different novel ternary nanostructured hematite-based photocatalysts of eRGO/C₆₀/α-Fe₂O₃ and eRGO/NiO/α-Fe₂O₃ were synthesized, characterised and tested as photoanodes toward PEC water splitting application. The ternary nanostructured hematite-based photoanodes were characterised using FE-SEM, EDX, XRD, XPS, as well as Raman, UV-Vis and EIS spectroscopic methods. It was found that the ternary nanostructured photoanodes of eRGO/C₆₀/α-Fe₂O₃ and eRGO/NiO/α-Fe₂O₃ showed 5-fold and 9-fold enhancement in current density and significant reduction in charge transfer resistance when compared to the pristine hematite photoanode. In this instance, the enhancement in PEC performance of ternary nanostructured eRGO/C₆₀/α-Fe₂O₃ photoanode was attributed to the electrons cavenging property of C₆₀ as well as the highly conducting eRGO property that have mitigated the high interfacial recombination rate of photogenerated electron-hole pairs. Whilst for the ternary nanostructured eRGO/NiO/α-Fe₂O₃ photoanode, eRGO transferred the electrons efficiently in the p-n heterojunction without causing substantial bulk recombination. Additionally, the internal electrostatic field in eRGO/NiO/α-Fe₂O₃ could facilitate the effective separation of photogenerated electron-hole pairs so that more holes could participate in the water oxidation reaction instead of recombination process. It is anticipated that the fundamental understanding gained through this study is helpful to design and construct high-performance photoelectrodes for the application in PEC water splitting in the near future.

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Van der Waals Oxide Heteroepitaxy for Transparent and Flexible Electronics

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Abstract

In the diligent pursuit of low-power consumption, multifunctional, and environmentally friendly electronics, more sophisticated requirements on functional materials are on demand. For example, flexible electronics represents a fast-developing field and has a great potential to impact our daily life. In building up flexible electronics, the materials with controllable conduction, transparency, and good flexibility are required. Recently, the discovery of free-standing 2D materials has created a revolution to this field. Pioneered by graphene, these new 2D materials exhibit abundant unusual physical phenomena that is undiscovered in bulk forms. In the meantime, it also possesses very high transparency to the visible light. However, the extensively studied pristine graphene naturally has no bandgap and become restricted in many field-effect based applications. Hence, looking for various types of new 2D materials has been a focal research direction nowadays. In this talk, we intend to take the same concept, but to integrate a family of functional materials in order to open new avenue to flexible electronics. Due to the interplay of lattice, charge, orbital, and spin degrees of freedom, correlated electrons in oxides generate a rich spectrum of competing phases and physical properties. However, a generic approach to build up flexible electronics based on functional oxides is yet to be developed. In this study, we use a 2D material as the substrate. And we take several functional oxides as model systems, including transparent conducting oxides, VO$_2$, NiO, Fe$_3$O$_4$, Pb(Zr,Ti)O$_3$, and oxide nanocomposites, to demonstrate a pathway to build up functional oxides for transparent and flexible electronics.

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Nanowire-plasmonic photocatalysts and thermal emitters

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Abstract

Optical absorption enhancement using plasmonic structures enables a wide range of applications such as solar energy harvesting devices, light emitting devices and photothermal management. For example, in plasmonic photocatalysis, it has recently attracted great interest in enhancing photocatalytic efficiency not only by the plasmon-enhanced near field but also by the plasmon-enhanced hot-carrier injection, which could boost the visible response of wide bandgap photocatalysts [1]. Here we report measurements and simulations of the efficient sunlight-driven and visible-active photocatalysts composed of plasmonic metals and ZnO nanowire (NW) arrays fabricated via an all-wetchemical route (Figure 1a) [2]. Another application of plasmon-enhanced light absorption is the perfect absorber and thermal emitter [3]. It is found that with proper designs supported by the electromagnetic simulation, the plasmonic structures could exhibit near perfect absorption at desired resonant wavelengths, making them promising for a number of potential application such as thermal emitters (Figure 1b) [4], molecular sensors [5] and IR sensors [6].

References


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Abstract

Water splitting and CO₂ fixation on heterogeneous photocatalysts are important reactions from the viewpoint of solar-to-fuel energy conversion. To achieve these reactions, it is important to improve both bulk and surface properties of a photocatalyst so as to suppress electron–hole recombination and promote surface redox catalysis. In this presentation, recent progress on the development of new photocatalysts that are active for such artificial photosynthetic reactions will be given. In particular, surface modification techniques developed by our group to construct active sites and light-absorbing centers will be presented. For example, we developed a new powdered photocatalyst consisting of Co(OH)₂ and TiO₂ [1]. It is well known that TiO₂ is an active photocatalyst, but only works under UV irradiation. By contrast, the Co(OH)₂/TiO₂ hybrid photocatalyst is capable of absorbing visible light with wavelengths of up to 850 nm and oxidizing water into oxygen gas, even though it consists of only earth-abundant elements only. To our knowledge, this system provides the first demonstration of a photocatalytic material capable of water oxidation upon excitation by visible light up to such a long wavelength.

References

Photoelectrocatalytic Production of Solar Fuels from Water and CO₂

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Abstract

There is renewed interest in the photocatalytic and photoelectrochemical conversion of CO₂ into value-added chemicals using various semiconductor particles and electrodes. Common CO₂ reduction products are C1 chemicals (CO, HCOOH, CH₂OH, and CH₄) in aqueous media, while the production of C2-C4 hydrocarbons (e.g., C₂H₆ and C₃H₈) has also been reported. A number of solar-active materials have been reported, but they still suffer from low selectivity, poor energy efficiency, and instability, while failing to drive simultaneous water oxidation. In this regard, p-type CuMOₓ (where M = Al³⁺ and Fe³⁺) is highly promising because of its unique structure; suitable bandgap energies; high conduction band level, which is sufficient for H₂ production and CO₂ reduction; and relative stability in aqueous solution. Unfortunately, the typical synthetic route for CuMOₓ is annealing a Cu(I) and M(III) salt mixture at high temperature, which inevitably results in irregular, coarse particles of several micrometers. Furthermore, the as-synthesized particles are difficult to fabricate into durable films on transparent conducting oxide (TCO) substrates because of the absence of particle-to-particle interaction. Even if they are fabricated, the films have less intimate and looser interparticle connections undergoing a significant charge recombination at the solid/solid interface. This difficulty in synthesizing CuMOₓ films has caused this material to be less studied despite its potential as a promising photocathode. With this in mind, we have attempted to synthesize high-efficiency CuMOₓ films on TCO substrates via electrochemical deposition. This talk presents our recent studies on the solar CO₂ conversion to value-added chemicals while using water as an electron donor in various photo-systems [1-11].

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Photocatalytic elimination of organic chemicals and aerosol-associated influenza virus infectivity in the air

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Abstract

The efficiency of photocatalysis depends on the surface area and materials, and we have prepared a nanosized-titanium dioxide (TiO₂)-coated ceramic irradiated by UV-LED lamps as a photocatalytic air cleaner. Ceramic filter system decomposed 80% of acetaldehyde and Particulate dioxins (40 pg/m³) and gaseous dioxins (16 pg/m³) were removed by 7.5 and 2.8 pg/m³ by passing through four TiO₂-coated ceramic (30 × 30 × 2 cm) under black-light, indicating about 80% of dioxin was decomposed by the photocatalysis. Ceramic was changed to aluminum plate and the efficiency was improved. The 90% of 5 ppm acetaldehyde (12.4 μmol/h) was decomposed and generated carbon dioxide (25.43 μmol/h; RC: 92.5% carbon dioxide conversion rate) efficiently and continuously for 200 min with the ratio of one acetaldehyde (12.40 μmol/h) to two carbon dioxide (25.43 μmol/h) at their molar ratios by being passed through the TiO₂-coated aluminum plate (5 × 10 × 1 cm) under black light, indicating complete decomposition of acetaldehyde with high efficiency. This photocatalysis system was applied for elimination of acetaldehyde and inactivation of influenza aerosol in a closed cubic space using aluminum plates. Acetaldehyde at 20 ppm in a cubic 1 m³ space was eliminated by 60 min at a half-life of 8 min. The aerosol-associated infectivity and the RNA genome of influenza virus produced by a nebulizer in a 779 liter cubic space were eliminated within 7 minutes but were detectable up to 28 minutes without the function of a photocatalytic air cleaner. Influenza virus was broken down by photocatalysis rather than being trapped by Hepafilter as intermediate breakdown products of influenza virus were observed. Thus, a photocatalytic air cleaner efficiently decomposed and eliminated organic chemicals, acetaldehyde, and aerosol-associated influenza virus infectivity and viral RNA, indicating a photocatalytic air cleaner functioned in the cleaning and detoxification of the air in the closed space for maintaining a safer environment.

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Photocatalytic Degradation of Polybrominated Diphenyl Ethers on TiO$_2$-based composites

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Abstract

Polybrominated diphenyl ethers (PBDEs) are widely used as flame retardants, and become a new class of global contaminants. Because PBDEs possess typical characteristics of persistent organic pollutants (POPs) like persistent, bioaccumulation, and biotoxicity, their elimination therefore attracts much attention of researchers.

The reductive debromination is a common strategy to treat PBDEs. Among various reductive methods, the photogenerated electron of TiO$_2$ is considered to be highly efficient to reduce decabromodiphenyl ether (BDE209) [1]. However, this leads to accumulation of brominated intermediates, but the debromination products with less bromine atoms are much more difficult to get further reductive debromination.

To promote the photocatalytic reductive debromination, we developed several TiO$_2$-based composites including reduced graphene oxide (RGO) loaded TiO$_2$ (RGO/TiO$_2$), Ag/TiO$_2$, and CuO/TiO$_2$ to reduce BDE209 and/or 2,2',4,4'-tetrabromodiphenyl ether (BDE47) [2-4]. These heterostuctured photocatalysts have two beneficial roles: charge separation in space, and enhanced adsorption of PBDEs. Thus, the photocatalytic reduction of PBDEs was greatly improved. However, the complete debromination of PBDEs to diphenyl ether through the photocatalytic reduction process is rather difficult. For example, although all the added BDE47 was rapidly reduced, the debromination efficiency was still less than 50% over Ag/TiO$_2$ and CuO/TiO$_2$ [3,4]. This is because that the reduction of low-brominated PBDEs is rather difficult, due to their weak electron affinity.

Although most studies indicate that BDE209 is strongly resistant to oxidation, we firstly demonstrated that the h$^+$/•OH-involved oxidative degradation of BDE209 took place slowly in the UV-irradiated TiO$_2$ aqueous dispersions [5]. We also noted that the oxidation of BDE209 is much slower than the following oxidation of the less-brominated organic intermediates. Since the highly brominated PBDEs are more easily reduced, and the lower brominated PBDEs become more susceptible to the oxidation, we then developed an effective “one-pot” photocatalytic system for driving concurrently the pre-reduction and consecutive oxidation of BDE47.

Acknowledgements

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Nanostructured TiO$_2$ with oxygen vacancies for the decomposition of organics

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Abstract

Titanium dioxide (TiO$_2$) has been widely not only studied but also applied in industrial fields as photocatalyst because of its environmentally and economically advantages with high chemical stability, earth abundant and bio compatible properties. However, its large band-gap for the activity to only UV light region, and the high recombination rate of photogenerated electron and hole pairs have to be overcome to utilize effectively sunlight and to enhance the photocatalytic performance. Recent enormous efforts to overcome the above-mentioned drawbacks have resulted in the one-dimensional TiO$_2$ nanotubes, nanofibers and nanorods to suppress the carrier recombination, and/or the heterojunction structure of TiO$_2$ with another semiconductor to achieve larger separation of the photogenerated electron and hole, as well as the modification of TiO$_2$ nanoparticles with gold clusters to expand the light conversion from UV to visible and near-infrared region. Here, we report simple and effective modification of nano-sized TiO$_2$ materials by in-liquid plasma processing, which is a non-thermal plasma discharged in liquid. The present study focused to treat pristine TiO$_2$ nanoparticles by the discharge in water-based solution and to investigate the material properties as well as the photocatalytic activities for decomposing organics. As a result of plasma treatment, we found the incorporation of oxygen vacancies on the sub-surface of TiO$_2$ nanoparticles, and concluded that the origin of photocatalytic enhancement for acetaldehyde decomposition under fluorescent lamp attributed in the recombination suppression by the surface trap of oxygen vacancies near the nanoparticles surface.

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One-dimensional nanoarrays for solar cells

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Abstract

Various kinds of modifying methods towards ZnO and TiO$_2$ one-dimensional nanostructures such as nanorods and nanotubes have been carried out for their applications in Dye-sensitized solar cell (DSSC), perovskite solar cell (PSC) and photo detectors (PD). Core-sheath ZnO/CdTe and double-sheath ZnO/CdSe/CdTe nanocable arrays as effective photoanode have been developed for solar cells, which resulted in a saturated current as high as 14.3 mA/cm$^2$. Large area free-standing highly ordered TiO$_2$ nanotube arrays on the fluorine-doped tin oxide (FTO) conductive glass substrates have been successfully obtained to serve as photo-anodes of DSSCs. The certified photovoltaic conversioncy of TiO$_2$ nanoarrays based DSSCs is up to 10.3% by using N719 as Dye and I$_3^-$ as electrolyte. Perovskite solar cell with efficiency of up to 18.6% based on TiO$_2$ nanorod arrays is also presented. Self-powered broadband photodetectors based on CH$_3$NH$_3$PbI$_3$/ZnO nanorod arrays heterostructure have been achieved with high detectivity of $3.56 \times 10^{14}$ cm Hz$^{1/2}$ W$^{-1}$ and high responsivity of 24.3 A W$^{-1}$.

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Facile Development of Nanostructured Photocatalysts for CO₂ Capture and Conversion

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Abstract

The continuous reliance on fossil fuel-based energy is inevitable. Rational strategies to reduce carbon dioxide (CO₂) emissions are thus highly demanded. Developing efficient photocatalysts that can harness solar energy appears to be a promising methodology to capture and recycle CO₂ as a fuel feedstock. The conversion efficiency of the current photocatalysts, however, is generally very low due to various limiting factors, such as fast electron-hole recombination rates, narrow light absorption range, and backward reactions. Thus, developing strategies to overcome the above limitations is an important task in this field.

Here we present several strategies through controlled synthesis to address the aforementioned limitations toward enhancing the overall CO₂ conversion efficiency. Examples of novel photocatalysts being explored include mesoporous nanocomposite particles (i.e., Cu-TiO₂-SiO₂) [1,2], 1D structured Pt-TiO₂ thin films [3], CuO-ZnO heterojunction nanowires [4], and crumpled graphene-based nanoballs [5,6]. Systematic materials characterization and photocatalysis analysis of the materials, by electron microscopy, X-ray diffraction, gas chromatography, X-ray photoelectron spectroscopy, in-situ diffuse reflectance infrared Fourier transform spectroscopy, and femtosecond time-resolved transient absorption spectroscopy, aid in understanding the quantitative CO₂ photoreduction pathways and the correlations between materials properties and CO₂ photoreduction performance.

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Highly Concentrated CO Evolution for Photocatalytic Conversion of CO₂ by H₂O as an Electron Donor

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Abstract

The reduction in human-induced emissions of CO₂ from automobiles, factories, power stations etc., over the next 15 years is currently one of the most important issues facing the planet. We should therefore attempt to develop industrial processes using CO₂ as a feedstock in order to build a sustainable society in the near future. Linear CO₂ molecules adsorbed on the solid bases are converted into unique structures, such as bicarbonate and carbonate species possessing lattice oxygen atoms. We believe that the process involves the capture and distortion of CO₂ upon adsorption on a solid base through activation by photoirradiation. Unstable CO₂ species adsorbed onto the surface can then be reduced by electrons with protons derived from H₂O. These days, we succeeded in designing highly selective photocatalytic conversion of CO₂ by H₂O as the electron donor, by the simultaneous use of an inhibitor of the production of H₂ and a material for CO₂ capture and storage, such as ZnGa₂O₄/Ga₂O₃ [1,2], La₂Ti₅O₁₄ [3], SrO/Ta₂O₅ [4], ZnGa₂O₄ [5] and ZnTa₂O₆ [6], and Sr₂KTaO₁₅ [7] with the modification of Ag cocatalyst. An isotope experiment using CO and mass spectrometry clarified that the carbon source of the evolved CO is not the residual carbon species on the photocatalyst surface, but the CO₂ introduced in the gas phase. In addition, stoichiometric amounts of O₂ evolved were generated together with CO.

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Highly efficient visible-light-driven BiVO$_4$-based Photoelectrode Materials

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Abstract

BiVO$_4$ is an excellent visible-light-driven photocatalyst that can split water into O$_2$ for its more positive valence band than that of O$_2$/H$_2$O, whereas its positive conduction band makes it difficult to directly generate H$_2$ from H$_2$O. But using the PEC, BiVO$_4$ can become a promising candidate as photoanode for hydrogen evolution performance. In our group, the leaf-like structure BiVO$_4$ photoelectrodes were prepared by electrochemical deposition, in which Zn$^{2+}$ ions were introduced as a direct agent to control the morphology and size of Bi nanoparticles [1]. NiFe$_2$O$_4$ and CoFe$_2$O$_4$ nanoparticles were loading on the surface of BiVO$_4$ to construct heterojunction. The heterojunctions can effectively prevent carriers from recombining and accelerate the separation of electrons and holes [2]. Besides, Bi/BiVO$_4$[3] and FeF$_2$/BiVO$_4$ can enhanced PEC hydrogen evolution performance.

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**Nano Photocatalytic Materials Design Toward Small Molecular Hydrocarbons Oxidation**

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**Abstract**

The search for active catalysts that efficiently oxidize methane under ambient conditions remains a challenging task for both C1 utilization and atmospheric cleansing [1-6]. Here, we show that when the particle size of zinc oxide is reduced down to the nanoscale, it exhibits high activity for methane oxidation under simulated sunlight illumination, and nano silver decoration further enhances the photo-activity via the surface plasmon resonance. The high quantum yield of 8% at wavelengths <400 nm and over 0.1% at wavelengths ~470 nm achieved on the silver decorated zinc oxide nanostructures shows great promise for atmospheric methane oxidation. Moreover, the nanoparticulate composites can efficiently photo-oxidize other small molecular hydrocarbons such as ethane, propane and ethylene, and in particular, can dehydrogenize methane to generate ethane, ethylene and so on. On the basis of the experimental results, a two-step photocatalytic reaction process is suggested to account for the methane photo-oxidation.

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TiO$_2$-C hybrid aerogel photocatalysts for methylene blue degradation

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Abstract

TiO$_2$-C hybrid aerogels were prepared by one-pot sol-gel process in ethanol, followed by supercritical drying and carbonization, using TiCl$_4$ as TiO$_2$ precursor, resorcinol-furfural as carbon precursors, ethyl acetoacetate (EA) as a chelating agent and propylene epoxide (PO) as a gel initiator. Ce doping was performed by adding cerous nitrate into the solutions that form gels to modify the photocatalytic properties. Microstructures of samples were characterized by XRD, SEM, TEM, UV-Vis, Raman spectroscopy, nitrogen adsorption, mercury porosimetry, XPS and IR spectroscopy and the photocatalytic properties for methylene blue degradation were tested under UV and Vis light irradiation. Results showed that the porous textures of the hybrid aerogels were related to TiCl$_4$/resorcinol-furfural mass ratio, the molar ratios of EA/Ti and PO/Ti. The samples had large specific surface areas, high adsorption capacities for methylene blue, uniform distribution of TiO$_2$ nanoparticles as anatase in the amorphous carbon structure. The presence of the amorphous carbon inhibited both the growth of TiO$_2$ nanoparticles and their conversion from anatase to rutile phase. The adsorption of methylene blue followed the pseudo-second-order kinetics model and its degradation followed the first-order kinetics model. The maximum photocatalytic activity for methylene degradation was up to 4.23 times that of P25 for the TiO$_2$-C hybrid aerogels carbonized at 800 °C. A partial reduction of Ti$^{4+}$ to Ti$^{3+}$ was found for the samples carbonized at 900 °C, which improved significantly the catalytic activity under visible light. Methylene blue can be degraded under visible light within 60 min with the Ce-doped TiO$_2$-C hybrid aerogels or the undoped one carbonized at 900 °C. Adsorption coupled with photo excitation, reduced recombination rate of e/h pair, band gap narrowing by interaction of carbon with TiO$_2$, partial reduction of Ti$^{4+}$ to Ti$^{3+}$ during carbonization or Ce-doping, and the increased light utilization via scattering by macropores are responsible for the improved catalytic performance as compared with P25 photocatalyst from Degussa.

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Noble metal-free Metal Sulphides as Highly Efficient Visible Light Driven Photocatalysts for H₂ Production from H₂S

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Abstract

Hydrogen sulfide is an extremely toxic gas which is generated from both nature and human factors. Recently, photocatalytic splitting of H₂S into H₂ and S has attracted great attention because hydrogen production and H₂S removal are simultaneously achieved. However, the deactivation of the photocatalysts and lack of suitable setup for photocleavage of H₂S to H₂ limit its wide application. Herein, we constructed a complete setup for H₂ production from H₂S. This setup has some functions including H₂S absorption, decomposition and product recovery. Simultaneously, a series of MnS/In₂S₃ composites were successfully fabricated by a solvothermal method. A maximum H₂ production rate of 8360 μmol g⁻¹ h⁻¹ can be achieved over MnS/In₂S₃₀.₇ catalyst, and the corresponding QE of this sample is as high as 34.2% at 450 nm even in the absence of any noble-metal co-catalysts. Importantly, MnS/In₂S₃ composite displays a good stability and also anti-photocorrosion. Additionally, in order to further enhance visible-light photocatalytic H₂ production activity of MnS/In₂S₃, MnS/In₂S₃/CuS composites were prepared through solvothermal treatment. And a maximum H₂ production rate of 29252 μmol g⁻¹ h⁻¹ can be achieved over a MnS/In₂S₃/CuS with optimized composition, which is 3.5 times higher than that of MnS/In₂S₃₀.₇, and this reveals that the addition of CuS can effectively increase the photocatalytic activity for splitting H₂S into H₂. All in all, suitable setup for photocatalytic splitting of H₂S and noble-metal free metal sulphides photocatalysts have great significance for photocleavage of H₂S to H₂.

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Figure 1. Photocatalytic process of splitting H₂S in 0.6 M Na₂SO₃/0.1 M Na₂S/3 M H₂S solution.
CO$_2$ Photocatalytic Reduction over TiO$_2$ Nanocrystals with Coexposed \{001\} and \{101\} Facets

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Abstract

CO$_2$ photocatalytic reduction with water is one of the most popular and challenging technologies to produce renewable energy. Engineering TiO$_2$ with coexposed \{001\} and \{101\} facets could enhance the conversion efficiency of CO$_2$ due to the effective separation of photogenerated charges caused by the formation of \{001\}/\{101\} surface heterojunction. However, faceted TiO$_2$ nanocrystals still suffers from low conversion efficiency and low selectivity for CO$_2$ reduction. Faceted TiO$_2$ nanocrystals were combined with graphene and metal nanoparticles respectively to improve the activity and selectivity of CO$_2$ photocatalytic reduction. The results show that the faceted TiO$_2$/graphene composites exhibited higher CO yield than that of pristine TiO$_2$ due to the formation of \{001\}/\{101\} surface heterojunction and supporting of graphene, which can effectively promote the spatial separation of photogenerated electrons and holes. Differing from graphene, Pt loading tended to promote the production of CH$_4$ and H$_2$ while Cu$_2$O suppressed H$_2$ evolution and exhibited lower CH$_4$ selectivity comparing with Pt. Furthermore, when Pt and Cu$_2$O were co-deposited on TiO$_2$ crystals, H$_2$ and CO production were both inhibited and CO$_2$ was selectively reduced to CH$_4$. Pt could not only capture photogenerated electrons but also increase the electrons density on the surface of TiO$_2$. Meanwhile, Cu$_2$O loading enhanced the CO$_2$ chemisorption on TiO$_2$ while inhibited that of water. As a result, Pt and Cu$_2$O co-deposited TiO$_2$ crystals exhibited high selectivity for CH$_4$ production.

Figure 1. Schematic diagram of graphene and metal modified faceted TiO$_2$ nanocrystals.
Abstract

Contrast agents play a vital role in the enhanced examination of biomedical imaging. However, traditional clinical small-molecule agents face a variety of drawbacks, such as low blood circulating time, difficult modification and potential toxic and side effects. Herein, a simple albumin directed fabrication of gold (Au) or platinum (Pt) nanoparticles was achieved for exploring the utilization in CT and MR imaging. Firstly, ultra-small nanoagents (Pt@BSA) with a mean core size of 2.1 nm were obtained through a facile one-pot synthesis by the reduction of chloroplatinic acid hexahydrate using bovine serum albumin (BSA) as the biotemplate under room temperature. It was demonstrated that the nanocrystals could serve as potential new and potent CT contrast agents, especially vital for in vivo imaging with prominent enhancement and metabolizable behaviours due to the combination of the higher X-ray attenuation property and prolonged imaging time [1]. Then gold nanoparticles (Au@BSA) were also prepared with BSA as a biotemplate following with conjugation of diatrizoic acid (DTA) for a potential CT imaging contrast agent (Au@BSA-DTA). The biomimetic material Au@BSA-DTA with double radiodense elements of Au and iodine displayed much stronger CT imaging effect compared with the traditional small molecule contrast agents [2]. Finally, a novel CT/MR contrast agent Au@BSA-Gd-DTPA was fabricated by modifying the as-prepared Au@BSA with diethylene triamine pentoacetic acid (DTPA), followed by the chelation of Gd (III) ions. For the CT phantoms, the formed nanocomplex showed an improved contrast in CT scanning than that of Au@BSA as well as small molecule iodine-based CT contrast agents, and for the $T_1$-weighted MRI images, the nanoagents displayed a relatively higher $r_1$ relaxivity than that of the commercial MR contrast agents. Importantly, the above mentioned nanoagents exhibited not only good colloid stability and water dispersibility, but also satisfying low-cytotoxicity and hemocompatibility. In summary, we have constructed a series of novel biomaterials that can be used as contrast agents for both X-ray CT and MR phantoms, which paves the potential clinical applications in cancer early diagnosis.

Figure 1. A schematic illustration of the preparation of the Au@BSA-Gd-DTPA.

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Nanoparticles in Magnetic Resonance Imaging

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Abstract

Magnetic resonance imaging (MRI) provides crucial roles in diagnosis and treatment of human diseases. More and more new MRI techniques have been developed recently. Among them, chemical exchange saturation transfer (CEST) imaging (Figure 1) has shown its promising for noninvasive pH imaging and metabolic imaging [1]. Nanoparticles have also been studied widely in the field of magnetic resonance imaging, including disease detection and stem cell migration. For example, superparamagnetic iron oxide nanoparticles (SPIONs) have been intensively studied for their biomedical applications as T2 contrast agents in MRI. We collaborated with Zhang BL group [2] and found that compared with other nanoparticles, SPIONs exhibit high magnetic responsivity which can reduce the amount of the contrast agents needed for calcium-responsive MRI, low cytotoxicity, higher biocompatibility and chemical stability. The assessment of changes in the extracellular calcium concentration by magnetic resonance imaging would be a valuable biomedical research tool to monitor brain neuronal activity. The nanoparticles, EGTA-SPIONs, have potential as smart contrast agents for Ca\(^{2+}\)-sensitive MRI. We also collaborated with Bu WB group [3] and found that both T1-weighted imaging and in vivo pH mapping can be successfully acquired on the kidney and glioblastoma (GBM) of the mouse after intravenous injection of the T1/CEST NaGdF\(_4@PLL\) nanodots (NDs), demonstrating the feasibility of such an anatomical and functional dual-mode imaging technique on one magnetic resonance machine by the rational design of MRI contrast agents. Meanwhile, the PLL shell exhibits a sensitive CEST effect that depends on the pH value of the lesions. Attractively, these ultrasmall nanoagents could be excreted through urine with negligible toxicity to body tissues, which has been demonstrated by the blood biochemistry, hematology, and tissue H&E staining analysis.

Figure 1. CEST images of creatine tubes of different concentrations.

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Proteins promote cancer nanotheranostics

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Abstract

Utilizing molecular imaging probes can increase the detection rate and diagnosis accuracy of tumors. However, the non-specific adsorption of nanoprobes is seriously affecting the detection accuracy and sensitivity. Leveraging on the unique domains and amino acid sequences of bio-endogenous proteins, some specific tactics, including surface engineering of nanoparticles with proteins, and protein-mediated biomimetic synthesis of nanoprobes have been developed in a green chemistry fashion in our group [1-4]. This protein-mediated cancer nanotheranostic strategy is of great significance in reducing nonspecific adsorption of nanoprobes for in vitro diagnosis and in vivo applications. Capitalizing on this strategy, protein-coated nanoparticles with single-/dual- imaging modality were fabricated. Furthermore, therapeutical moieties can be imparted to the above structures in a one-pot manner for enhanced tumor theranostics. Particularly, the operating mechanisms of these reactions are preliminarily studied [5].

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Tracking of Stem Cells with Magnetic Resonance Imaging

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Abstract

Recently various cell therapies have improved outcome of many diseases. A critical component of cell therapies is the ability to track the cells after transplantation. Monitoring a cell’s survival, migration, differentiation, and integration within host tissue is crucial for assessing the safety and efficacy of cellular treatments. In this talk magnetic resonance imaging (MRI) based tracking of stem cells is presented, where cells labeled with superparamagnetic iron oxide (SPIO) nanoparticles can be tracked in a chicken embryo model up to 10 days post transplantation. Nevertheless, SPIO nanoparticles as a $T_2^*$ contrast agent are usually associated with signal loss in MR images, leading to difficulties for cell tracking. To overcome this problem, a new imaging sequence, SWIFT with water and fat suppression, is introduced. Compared with other conventional pulse sequences, such as spin echo and gradient-recalled echo, the SWIFT approach enhances in vivo mapping of SPIO distribution in tissues, and improves detection sensitivity of SPIO nanoparticles.

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Tuning solar absorbance and reflection of high-temperature solar spectrally selective surfaces

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Abstract

Spectrally-selective solar absorbers are widely used in solar hot water and concentrating solar power (CSP) systems [1,2]. However, the performance at high temperatures (>450 °C) is still not satisfactory due to their high infrared (IR) emittance and long term thermal stability. Recent progress on cermet-based solar absorbers has shown promising temperature thermal stability and wavelength selectivity. Thus, here we explore W-Ni-Al2O3, W-Ni-YSZ (yttria-stabilized-zirconia) and W-Ni-SiO2 cermet based spectrally selective surfaces for high-temperature solar absorber applications [3,4]. The developed multilayer selective surfaces are deposited on a polished stainless steel substrate comprising two sunlight absorbing cermet layers with different W-Ni volume fraction inside the dielectric matrix, one or two anti-reflection coatings (ARCs) and one tungsten IR reflection layer for reduced IR emittance and improved thermal stability.

Regarding the W-Ni-Al2O3 cermet based solar absorbers; we observed a detrimental change in the morphology, phase, and optical properties if the cermet layers are deposited on a stainless steel substrate with a thin nickel adhesion layer, which is due to the diffusion of iron atoms from the stainless steel into the cermet layer forming a FeWO4 phase. A 100 nm thick tungsten layer can suppress the degradation of the optical properties at high temperatures and lowers the emittance relative to the stainless steel substrate, which improves the spectral selectivity of the solar absorber. We experimentally demonstrated a solar absorber with a solar absorptance of ~0.9 and total hemispherical emittance of ~0.15 at an operating temperature of 500 °C.

The fabricated W-Ni-YSZ cermet based solar absorbers are tested for their long term thermal stability at 600 °C. A distinct change in surface morphology of the solar absorbers with high oxygen deficiency in their YSZ-ARC layers, suggests to causing the degradation of the optical properties at high temperature. The oxygen deficiency can be effectively overcome through increasing the oxygen partial pressure during sputtering, which leads to a stable solar absorber with an experimentally demonstrated solar absorptance of ~0.91 and a total hemispherical emittance of ~0.13 at 500 °C.

We also developed a new kind of absorber that reflects a certain range of wavelength and absorbs the rest of the whole solar spectra [5]. The absorbed part is used for electrical power generation by steam engine and the reflected part is used for solar photovoltaic conversion. The thermal energy can be easily stored for later conversion to provide electrical power around the clock without worrying the Sun’s night time.

References

Titanium Dioxide Films with Pure Anatase Phase Synthesized by Mist Chemical Vapour Deposition

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Abstract

Pure anatase structured TiO\textsubscript{2} films were successfully synthesized by a novel fine channel mist chemical vapour deposition which is a vacuum free, low temperature method [1]. The effects of TTIP concentration on the morphological, structural and optical properties of TiO\textsubscript{2} films were investigated. It was confirmed that anatase crystallinity of TiO\textsubscript{2} film increased with the increase of TTIP concentration. Figure 1 showed that XRD patterns of TiO\textsubscript{2} thin films which were synthesized with the different concentration of TTIP in ethanol varying from 0.025 to 0.4 mol/L. The high catalytic activity will be expected by using obtained stable anatase TiO\textsubscript{2} films.

Figure 1. XRD patterns of TiO\textsubscript{2} films synthesized at TTIP concentration of 0.025, 0.05, 0.1, 0.2, and 0.4 mol/L.

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Abstract

Plasmonic metamaterial absorbers have garnered significant interest due to their unique ability to trap light beyond diffraction limit and potential applications in energy harvesting and information processing. Especially, the broadband absorbers show fascinating applications in photovoltaics and thermophotovoltaics, bolometers, thermal emitters, and photodetectors. In this talk, I will review our recent work on broadband optical absorption based on plasmonic based on ultra-thin metal-insulator-metal (MIM) plasmonic absorbers [1,2].

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One-Dimensional Nanomaterials for Energy Storage

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Abstract

One-dimensional nanomaterials can offer large surface area, facile strain relaxation upon cycling and efficient electron transport pathway to achieve high electrochemical performance. Hence, nanowires have attracted increasing interest in energy related fields. We designed the single nanowire electrochemical device for in situ probing the direct relationship between electrical transport, structure, and electrochemical properties of the single nanowire electrode to understand intrinsic reason of capacity fading. The results show that during the electrochemical reaction, conductivity of the nanowire electrode decreased, which limits the cycle life of the devices [1]. We have fabricated hierarchical MnMoO$_4$/CoMoO$_4$ heterostructured nanowires by combining “oriented attachment” and “self-assembly” [2]. The asymmetric supercapacitors based on the hierarchical heterostructured nanowires show a high specific capacitance and good reversibility with a cycling efficiency of 98% after 1,000 cycles. Then, we designed the general synthesis of complex nanotubes by gradient electrospinning, including Li$_3$V$_2$(PO$_4$)$_3$, Na$_{0.7}$Fe$_{0.7}$Mn$_{0.3}$O$_2$ and Co$_3$O$_4$ mesoporous nanotubes, which exhibit ultrastable electrochemical performance when used in lithium-ion batteries, sodium-ion batteries and supercapacitors, respectively [3]. In addition, we have successfully fabricated a field-tuned hydrogen evolution reaction (HER) device with an individual MoS$_2$ nanosheet to explore the impact of field effect on catalysis [4]. We also constructed a new-type carbon coated K$_{0.7}$Fe$_{0.5}$Mn$_{0.5}$O$_2$ interconnected nanowires through a simply electrospinning method. The interconnected nanowires exhibit a discharge capacity of 101 mAh g$^{-1}$ after 60 cycles, when measured as a cathode for K-ion batteries [5]. Our work presented here can inspire new thought in constructing novel one-dimensional structures and accelerate the development of energy storage applications.

Reference


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Green Semiconductor Vertical-Cavity Surface-Emitting Lasers based on Quantum Dots

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Abstract

Green VCSELs emitting in the spectral range from 491.8 nm to 565.7 nm, covering most of the ‘green gap’, are demonstrated. These devices are featured with low threshold current and CW lasing at room temperature. A few technologies such as laser liftoff of sapphire substrate, the removal of high-defect GaN buffer and Cu plating to increase the heat dissipation. Two dielectric DBRs were adopted as the cavity mirrors. The results presented here open up opportunities to design and fabricate semiconductor green lasers with excellent performance that may lead to wide-gamut, low consumption power and compact displays and projectors. The VCSELs could also be bonded on to Si for integration with other optoelectronic devices/circuits.

Figure 1. Lasing spectra (left) in this work and comparison (right) of threshold currents with literatures.

References


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Spin Texture And Spin Injection In A 3D Topological Insulator

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Abstract

One of the most critical steps towards spin functionalized electronics and optoelectronics is to generate and manipulate spin current in a desirable way. In 3D topological insulators (TIs), a strong spin-orbit interaction and the time-reversal symmetry result in spin-momentum locking of the surface electrons, which leads to a unique surface spin texture and the prospect of generating directional and dissipationless spin current running across the surface that is promising for spintronic applications. However, the metallic nature that is often found to be inherent to many 3D TIs due to residual defects has unfortunately imposed a severe obstacle to controlling surface spin current. As a result, very little experimental work has been done so far on this issue. Moreover, since most of the early studies have been limited to Bi₂Se₃ - a prototypical TI with a rather weak hexagonal warping effect, the contribution of the out-of-plane spin texture to the photocurrent remains elusive so far. In this work, we show that, with circular polarized light, helicity driven photocurrent is obtained in another 3D TI Bi₂Te₃ that exhibits a stronger hexagonal warping effect. We find the helicity-dependent photocurrent to be sensitive to the incident angle of the light, which could be explained within the framework of the circular photo-galvanic effect (CPGE) by taking into account the spin texture of the topological surface state. By correlating the light incident angle and probing surface current directions, we are able to identify photocurrent components associated with the in-plane and out-of-plane spin texture of the TI and thereby directly uncover the impact of the out-of-plane spin texture on surface spin current promoted by the strong hexagonal warping effect. By exploring the out-of-plane spin texture, we demonstrate spin injection from GaAs to TI and its significant contribution to the surface current [1]. We further show that the spin current of TI can be manipulated by the precession of injected electron spins in an external magnetic field. These discoveries pave the way to not only intriguing new physics but also enriched spin functionalities by integrating TI with conventional semiconductors, such that spin-enabled optoelectronic devices may be fabricated in such hybrid structures.

References


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Quantum phase transitions and order parameters of a topological insulator

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Abstract

Using quantum entanglement, quantum coherence, and the reduced density matrix, we study the quantum phase transitions and propose order parameters for the phases of a topological insulator, specifically a spinless Su-Schrieffer-Heeger (SSH) model, and consider the effect of short-range interactions. All the derived order parameters and their possible corresponding quantum phases are verified by the entanglement entropy and electronic configuration analysis results. The order parameter appropriate to the topological regions is further proved by calculating the Berry phase under twisted boundary conditions. It is found that the topological nontrivial phase is robust to the introduction of repulsive intersite interactions and can appear in the topological trivial parameter region when appropriate interactions are added.

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Novel Topological Phase with a Zero Berry Curvature

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Abstract
Recalling the scenario of Aharonov-Bohm effect that electrons experience a phase shift induced by a magnetic vector potential in spite of a zero-magnetic field, we have a similar story in topological version to tell here, where the magnetic field and the magnetic vector potential are replaced by their geometric counterparts, Berry curvature and Berry connection, respectively.

Starting from a simple two-dimensional tight-binding model with two types of hopping, i.e. intercellular $\gamma$ and intracellular hopping $\gamma'$ based on square lattice, we show that nontrivial topological phase emerges when $|\gamma'| < |\gamma|$ under a zero Berry curvature. Our work offers a new way to design topological materials without Berry curvatures [1].

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Spin texture of flatten Dirac-cone surface state on W(110)

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Abstract

Topological insulators and Rashba systems with spin-split energy band structure induced by strong spin-orbit interaction have attracted a great attention for the dissipationless spin current transport. The spin orientation of such spin-split state is locked with their crystal momentum and is strongly influenced by the symmetry of surface crystal. However, so-far, most of topological material and Rashba systems are sp-electrons system with C₃ᵥ point group symmetry [1].

Recently, we have reported spin polarized Dirac-cone surface state on W(110) with C₂ᵥ symmetry [2]. This surface state is formed by d-electrons and strongly influence by two-fold symmetry: the massless and massive band dispersion along ΓH and ΓN. Moreover, by model Hamiltonian based on k•p theorem, it have been predicted that the spin-polarized flatten Dirac-cone surface state shows quasi-one dimensional spin texture as shown in Figure 1 [3]. However, there is no evidence for the spin texture on W(110).

In this presentation, we have clarified the spin texture of flatten Dirac-cone surface state on W(110) studied by spin- and angle-resolved photoemission spectroscopy. The observed spin texture is good agreement with our predicted one. This research is the model case of d-electron –based surface state with C₂ᵥ symmetry. The finding opens a new avenue in the study of d-electrons-based spin texture with C₂ᵥ symmetry. If I have remaining time enough to talk another symmetry surface of tungsten, I will introduce you.

Figure 1. (a) Energy Contours of the surface state as function of kₓ, kᵧ for energies around crossing point based on model calculation. (b) Spin texture for the constant-energy surface (solid line) at 0.02eV above crossing point. The in-plain spin components are shown as arrows.

In this presentation, we have clarified the spin texture of flatten Dirac-cone surface state on W(110) studied by spin- and angle-resolved photoemission spectroscopy. The observed spin texture is good agreement with our predicted one. This research is the model case of d-electron –based surface state with C₂ᵥ symmetry. The finding opens a new avenue in the study of d-electrons-based spin texture with C₂ᵥ symmetry. If I have remaining time enough to talk another symmetry surface of tungsten, I will introduce you.
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One-dimensional edge states with spin splitting in bismuth

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Abstract

A two-dimensional (2D) system with strong spin-orbit coupling like a topological-insulator surface and semiconductor-heterostructure interface has provided a useful platform for realizing novel quantum phenomena applicable to advanced spintronic devices. 1 bilyer (BL) bismuth is theoretically predicted to be 2D topological insulator and have a spin-polarized state at edge [1,2]. However, it is not certain experimentally because to preparer a free-standing 1 BL bismuth is very difficult. Here we challenged to observe a spin-polarized electric state at edge by different approach. As observed by the atomic force microscopy (AFM) of our Bi thin film (Figure 1a), triangular-shaped bismuth BL islands with typically ~0.1 μm edge length are formed on the top surface of the Bi thin film, and the edge of each island runs along the ΓK direction in the k space. And we have also observed the 1D band dispersion from the edge state of bismuth islands measured by ARPES. In this presentation, we show the result of ARPES and spin-resolved ARPES for bismuth thin film, and discuss the origin of the 1D spin-splitting band compared with our first-principles band-calculations.

References


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Figure 1. (a) AFM image measured at 300 K in Air and (b) Band dispersion near EF along ΓK line measured at 30 K in UHV for a Bi thin film, respectively.
Magnetic Topological Insulators and Their Heterostructures

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Abstract

When magnetic order is introduced into topological insulators (TIs), the time-reversal-symmetry is broken, and the non-trivial topological surface is driven into a new massive Dirac-fermions state. By adjusting the Fermi level position, quantum anomalous Hall effect (QAHE) emerges in the Cr-doped (BiSb)2Te3 samples where dissipationless chiral edge conduction is realized in the macroscopic millimeter-size devices without the presence of an external magnetic field, and the stability of the dissipationless chiral edge conductance is well-maintained as the film thickness varies across the 2D hybridization limit. By further manipulating the topological surface gap, we realize the metal-to-insulator quantum phase transition in the system.

In addition to the uniform magnetic TIs, our recent work on several magnetic TI based heterostructures will be presented. First, in the TI/Cr-doped TI system, we demonstrate that the spin-orbit torque is highly efficient that the critical charge current density required for the magnetization switching is three orders of magnitude smaller than that of heavy metals. In addition, by constructing novel AFM/TI heterostructures, we realize emergent interfacial magnetic effects, which can be tailored through artificial structural engineering. Finally, by introducing additional superconductivity (SC), we observe the presence of the chiral Majorana edge mode in the QAHE-SC hybrid system. All these exotic magnetic TI-based phenomena will serve as fundamental steps to further explore the TRS-breaking TI systems.

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Charge-current induced spin polarization in BiSbTeSe$_2$ topological insulators

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Abstract

The surface states of 3D topological insulators (TIs) possess a helical spin texture in which the spin and momentum are perpendicularly locked to each other. Due to this spin-momentum locking, a net spin polarization can be induced by a charge current and vice versa. However, topological surface states are expected to give rise to only one type of spin polarization for a given current direction, which has been a limiting factor for spin manipulations. In this talk we report that in devices based on the bulk-insulating topological insulator BiSbTeSe$_2$, two different kinds of spin polarizations were observed in different devices: The spin polarization expected from the topological surface states was detected in a heavily electron-doped device, whereas the opposite polarization was reproducibly observed in devices with low carrier densities [1]. We propose that the latter type of spin polarization stems from topologically-trivial two-dimensional states with a large Rashba spin splitting, which are caused by a strong band bending at the surface of BiSbTeSe$_2$ beneath the ferromagnetic electrode used as a spin detector. This finding paves the way for realizing the “spin transistor” operation in future topological spintronic devices.

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Abstract

Ferrofluids have demonstrated great potential for a variety of manipulations of diamagnetic (or non-magnetic) micro-particles/cells in microfluidics, including sorting, focusing, and enriching. By utilizing size-dependent magnetophoresis velocity, most of the existing techniques employ single phase ferrofluids to push particles towards channel walls. In this work, we demonstrate a novel strategy for focusing and separating diamagnetic micro-particles by using the laminar fluid interface of two co-flowing fluids -- a ferrofluid and a non-magnetic fluid [1]. As shown in Figure 1, next to the microfluidic channel, microscale magnets are fabricated to generate strong localized magnetic field gradients and forces. Due to the negative magnetophoresis force, diamagnetic particles suspended in the ferrofluid phase migrate across the ferrofluid stream at size-dependent velocities. Because of the low Reynolds number and high Peclet number associated with the flow, the fluid interface is sharp and stable. When the micro-particles migrate to the interface, they are accumulated near the interface, resulting in effective focusing and separation of particles. We investigated several factors that affect the focusing and separation efficiency, including susceptibility of the ferrofluid, distance between the microfluidic channel and microscale magnet, and width of the microfluidic channel. This concept can be extended to multiple fluid interfaces. As an example, complete separation of micro-particles was demonstrated by using a three-stream multiphase flow configuration.

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Porous carbon fibers for electromagnetic wave absorption

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Abstract

Modern warfare urgently needs radar stealth technology to meet the shelter requirements of weapons and equipment. In this paper, porous carbon fibers were developed as an efficient microwave absorbent filler to prepare microwave absorbing composites. Firstly, polyacrylonitrile (PAN) was used as carbon precursor polymer (CPP), while polymethyl methacrylate (PMMA) as thermally decomposed polymer (TDP). PAN/PMMA (70/30) blend fibers were prepared by a wet spinning of PAN/PMMA solutions. Secondly, Porous carbon fibers (PCF) were obtained through carbonization of the blend fibers. The molecular weight of PAN could be used to control the size of pores in PCF, that is, when the average molecular weight of PAN was 51,000 g/mol and 83,000 g/mol respectively, PCF with pores ranging 1-10μm in diameter (designated PCF-L); and the other one with pores with diameter in the range of 0.1-1 μm (designated PCF-S) were obtained. The resultant porous fibers, PCF-L and PCF-S, were used as microwave absorbing fillers to mix with epoxy respectively, the composites containing 2-6 wt% of the filler were fabricated. It was found that PCF-S filled composites showed much better microwave absorption performance than that PCF-L filled one. The composite containing 6 wt% of PCF-S reached the lowest reflection loss of -32dB; and the reflection loss below -10 dB covered the whole X band. The change or destruction of the porous structure in PCF showed great effect on the microwave absorption properties, which demonstrate that the pores in PCF could make great contribution to the microwave absorption.

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Chiral modes of topological semimetals under magnetic field

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Abstract

Topological Dirac/Weyl semimetals, two new quantum phases of matter, attract broad interests from both condensed matter and particle physicists [1-3]. A Dirac (Weyl) semimetal with degenerate (nondegenerate) linear touchings, dubbed as Dirac (Weyl) points, in the electronic band structure is protected by various crystal symmetries (topology) [4-12]. While a Weyl point bears a topological charge in terms of the momentum-space Berry gauge flux, a Dirac point is neutral since it consists of two Weyl points of the opposite topological charge [13]. In addition, the Dirac (Weyl) semimetal exhibits cusps (Fermi arcs) instead of the conventional Fermi ring at the boundary of the Brillouin zone. On the other hand, because of the Landau level formation under an external magnetic field, these band touchings gain to hold massless chiral one-dimensional channels rarely seen outside the discussion of fundamental particles. As a result of the famous chiral anomaly [14-21], the chiral magnetic effect [22-24] is realized in such systems and is observed as the negative magnetoresistance [25-28].

Here, we try to provide a natural but yet missing analysis of the chiral matter, Weyl semimetal, in terms of the powerful framework of Tomonaga-Luttinger liquid, which enables us to examine the correlation and localization effects largely enhanced in this system under a strong magnetic field. We found new features unique to the 1D channels such as the independent critical exponents for the Greens function and the resistivity, which can be directly compared with experiments of realistic materials. The ubiquitous presence of a large number of Weyl points is also taken into account.

Besides, we consider the Dirac semimetal in the form of a nanowire, i.e., new ingredient of confinement geometry is added to this conventional gapless topological semimetal. Once a magnetic field along the nanowire direction is further applied, there will occur a competition between the effects of the confinement and the magnetic field, which strongly affects how the band gap is opened in the system. Expectedly, the system at finite temperature will show distinct transport features as one turns on and gradually increases the external magnetic field.

There has been an increase of interest recently by the micro- or nano-technology community in considering topological materials emerged in the last decade, whose novel topological properties may bring about new possibilities in various applications. Based on these findings, we hope to clarify a few aspects from the viewpoint of either fundamental science or nano-micro engineering.

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Novel Self-powered UV-Visible Photodetector with Fast Response and High Photosensitivity Employing Fe:TiO$_2$/n-Si Heterojunction

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Abstract

A UV-Visible photodetector employing heterojunction between the Fe:TiO$_2$ and Si was fabricated via a facile solution process. The existence of built-in electric field between TiO$_2$ and Si help facilitate the separation of photogenerated electron-hole pairs and regulate the electron transport. Under zero bias, the device exhibited high responsivity of 46 mA/W (350 nm) and 60 mA/W (600 nm) with a 0.5 mw∙cm$^{-2}$ light irradiation. At a small reverse bias of -0.5V, the quantum efficiency of the heterojunction rise up beyond 100% with a broad wavelength range. The exploring of Fe:TiO$_2$/n-Si heterojunction photodetector demonstrates an ultrasensitive (on/off ratio up to 10$^3$), fast (rise/decay time of <10/15 ms), and broad-band (UV-visible) photodetection with no or low external energy supply. Such novel photodetector with Fe:TiO$_2$/n-Si Heterojunction might be potentially useful for relative applications with weak-signal fast detection in UV-visible band.

Figure 1. Responsivity and EQE of Fe:TiO$_2$/n-Si heterojunction under each bias in UV-visible band.

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Gram-scale production of nanoporous graphene by Mg-thermoreduction of CS$_2$ for electrochemical energy storage

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Abstract

Mass production of porous carbon at low cost for supercapacitor applications is highly desired. Herein, for the first time we report a novel method to grow porous 3D graphene by using Mg to reduce CS$_2$ vapor. In a tube furnace the mixture of Mg and NaCl powders are heated at 570 $^\circ$C and reacted with CS$_2$ vapor carried by an Ar flow. The Mg powder can be about 95% consumed and the weight of resultant carbon nanomaterial is in gram scale after one experiment. This 3D graphene product with specific surface area of 980 cm$^2$/g is processed into electrodes for electrochemical test, showing a specific capacitance of 212 F/g at 1 A/g in H$_2$SO$_4$ aqueous electrolyte. Besides, the electrode is continuously charged/discharged at 5 A/g for 10,000 cycles and its capacitance remains 98.5% of the original value.

![SEM images of the 3D graphene product at low and high magnifications.](image)

Figure 1. (a) and (b) SEM images of the 3D graphene product at low and high magnifications.

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Enhanced electrochemical energy storage of N-doped graphene by adsorbing molecules of hydrolysized polyimide

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Abstract
A novel approach for increasing the specific capacitance of graphene-based supercapacitors is reported. In this work, Kapton, which is polyimide (PI) film widely used in industry, is used as the starting material to functionalize graphene. Hydrolysis of PI in alkalic solution released small aromatic molecules containing pyrrolic nitrogen that can be dissolved in water and easily adsorbed onto graphene sheets via π–π interaction (Figure 1). These small molecules store charge via a redox process, endowing graphene with pseudocapacitance and increasing the specific capacitance. N-doped graphene films are obtained by hydrothermally reducing solid composite films consisting of graphene oxide, ammonium acetate, and salt. These films are firstly used as electrodes to make symmetric supercapacitors with H₂SO₄ aqueous electrolyte as electrolyte, and show specific capacitances around 310 F/g. After the adsorption of hydrolysized PI molecules, the weight of graphene film is ~3% increased, and the specific capacitance was remarkable increased up to 467 F/g. Notably, the areal specific capacitance is in the scale of 1.2 F/cm². When using the Li₂SO₄ aqueous electrolyte that can extend the potential window to 1.5 V, the device also remains high specific capacitance around 445 F/g, and exhibits high energy densities up to 35 Wh/Kg. Our devices not only deliver excellent capacitive performances in aqueous electrolyte (89% capacitance retention at 20 A/g and 85% capacitance retention over 5 000 cycles), but also exhibit extraordinary mechanical flexibility. This novel strategy by adsorbing small molecules from hydrolysized PI provides a promising route towards high-performance supercapacitors.

Figure 1. Illustration of the adsorption of hydrolysized PI molecules onto graphene sheet.
Recent research in the continuous terahertz wave on nondestructive detection of carbon fibre composite

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Abstract

Terahertz nondestructive detection system is to test the image of the carbon fibre composite material. We use the continuous back wave oscillator (BWO) to pump the terahertz radiation, which the frequency is range from 210-250 GHz. By the reflection type detection system, the terahertz probe focus moves from the surface of the concealed carbon fibre composite material. In order to get real-time scanning the material image, the material is located by the 2D translation machine, which is connected to the computer. The data is collected by the phase-locked amplifier and calculated by the Matlab using the least square method (LSM). The interior structure, such as cracks, inclusions, empty and bubbles, can be detected by the images nearly 1mm resolution, which is a more convenient and simple method of nondestructive detection of the carbon fibre composite material.

Introduction

The composite material is made from different materials, which is produced by physical or chemical mechanism in the macroscopic. It is largely depended on the performance of various materials, which can make it better than the original composition. For this reason, it has played an important role in many fields, such as aerospace, automobile, shipbuilding, construction and other products. As the biggest aircraft, the carbon fibre composite is used to cut down the weight of 50% less than before. So it has been in a new fashion to lightweight, energy saving, safety development and green manufacturing on changing the traditional materials. But the carbon fibre composite material is difficult to the cutting processing, the blade wear in the interaction process of fiber is still very serious. In addition, cause it has very different material properties, low bond strength between the layers, such as fiber pull out, torn between the layers, voids and the other serious problems influence the quality of the product. So how to detect the quality and nondestructive, have been the key to restrict its further application.

Continuous terahertz wave have found the unique capacity to penetrate composite material and identify defects such as cracks, inclusions, voids and other [1–7]. Many years, there have been much research work done on the terahertz detecting cause of its importance character as high transmittance, safety for human beings and for other organisms. The ability of imaging the defects in aerospace composite materials, acquired during operation, by THz time-domain spectrometer (TDS) was researched in [1]. As the TDS system always needs a stable and complex optical roots, our work shows the ability of non-destructive detection in composite materials, reinforced by carbon fiber, by using continuous terahertz wave imaging system. The experimental is easy to build up and results show that the resolution is better than 1mm, and the information to collect the image of the measured sample can be obtained by two-dimensional (2D) scanning system.

Principe of the methods

With the complex structure of fibre carbon composite materials, there is a bunch of characteristic parameters that have to be focused on: fibre volume content, porosity, inclusions, voids, regions of insufficient resin infiltration and others. Continuous terahertz wave imaging system is a non-destructive and contactless inspection technique that is based on the propagation and reflection of optical waves. All the imaging system is based on source, our work is used the continuous back wave oscillator (BWO) as the source of terahertz radiation. It can easily operate the frequency and range from 160 to 2100 GHz. The output power has the relationship with the frequency, which is given by:

$$U\left(f\right) = \left(U_0 + U_1 f + U_2 f^2 + U_3 f^3\right)^2$$  (1)

from the front surface of the material and the other one reflected from the material attached to the high reflective surface. From the difference for the reflective of the thickness and the surface of the sample, the intensity of the continuous terahertz wave is not the same. For the BWO is a continuous terahertz source of fixed frequency output, obtained the intensity by the detector, we use point to point scanning imaging method to record the intensity and put the relevant information into computer, which can image the composition of the sample, such as the shape and the size.

Experimental setup

Figure 1 presented the reflected imaging system from the sample of the composite material. The BWO with an output power in 0.4mW is build up as the terahertz wave source. We choose the output frequency is from 206.2 to 260 GHz,
which is extremely coherent, polarized, and monochromatic (\(\Delta v/v \approx 10^{-5}\)). The continuous terahertz wave generated by the BWO is modulated by a chopper. The wave exiting from BWO is divided into two parts by a high-resistivity (R \(\geq 10000\Omega\)) silicon wafer, with 0.5mm thick, acting as a wave splitter. One of the split beams directly incidents into the focusing lens with collecting the wave light focusing onto the sample of composite material. The focusing lens always consist with two lens, which are collimator lens. With the attenuated reflection, the polyethylene lens can get the wave focus well to the sample. The length of optical root should be short and high transmittance. In this lens group, we choose the polyethylene lens, which are well performance in terahertz wave focusing.

The other split beam incidents into the detector, which converts the intensity of the terahertz into voltage signal as the input signal for the lock-in amplifier, and reference frequency for the lock-in amplifier is the same as the chopper. By this way the noise of the signal is filtered, we can obtained the intensity of the surface of the sample. After, with the acquisition unit, the data is transferred into the computer. Finally we can calculate each point by matlab and imaging the sample.

As depicted in Figure 2 is the imaging system we set up in the lab. We put one piece of paper before the sample in order to achieve the non-destructive of the material. Due to the simple system, it can easy to connect all the instrument without the external interference.

![Figure 1](image1.jpg)  
**Figure 1 Schematic of the terahertz reflected imaging system.**

Compared with the traditional transmission optical non-destructive system, the terahertz reflection non-destructive imaging system is a little different to set up, firstly, the terahertz is invisible, and the reflected direction is always changed, so the collection is easy to be affected by the surface of the composite material orientation, component space distribution and other factors. Secondly, the reflected wave consists of lens reflect wave diffuse wave and scattered wave. And the roughness, the size of the focusing and the signal to noise ratio of the sample can also influence the composition of the reflected wave and the data processing method.

![Figure 2](image2.jpg)  
**Figure 2 Schematic view of the experimental non-destructive system.**

![Figure 3](image3.jpg)  
**Figure 3 2D energy spot of the continuous terahertz wave.**

**Experimental results**

For the continuous terahertz wave generated from the BWO, the optical focusing should adjust to the plate of the sample, which is highest energy point of the sample. We use the point-to-point scanning system to detect the sample, so the wave spot from the polyethylene lens is changed by the focal. The Figure 3 is the measured result of energy density distribution in the plane view, which is the sample position located, acquired by using a 2D terahertz camera. The terahertz camera is Ophir-Spiricon, made from America. In this case, the optical roots can be ready to detect the sample of the composite material.

![Figure 4](image4.jpg)  
**Figure 4 The sample of fibre cabin composite material.**

![Figure 5](image5.jpg)  
**Figure 5 The image result of scanning the sample.**

In order to verify the experimentally non-destructive system of modeling, Before the sample of carbon fibre composite material (as shown in Figure 4), we put a piece of paper to set up the detection system. The sample is like square with 5\(\times\)4\(\times\)0.5cm of the LWH size. With place the sample at the focal point of the focusing lens, the reflect wave can reach into the detector. After the data processing in the computer, the detection image of the sample is obtained with the resolution within 1 mm. The resolution depends on the output of the BWO frequency and the distance of the X-Y plane. But if the resolution
is higher, the time of the scanning image would be longer. And the theoretical focal spot diameter at 206.2 GHz is 0.93 mm by the relationship of NA. Figure 5 have shown the image result of the composite material with the Matlab. It can easily present the profiles of the sample’s surface with non-destructive the material. Therefore, the system can be used as a non-contact scanning image system for detecting various composite materials.

Conclusion

In this paper, the method of a reflection terahertz wave non-destructive system is demonstrated to image the carbon fibre composite material. By the pyroelectric detector to get the signal to phase-lock amplifier, it converts the optical intensity to electric signal, which we can test the sample. Compared with other image systems, the reflection terahertz imaging system has more practical application value than the transmission imaging system, it can apply on nondestructive, non-contact and extraction of material surface profile. The resolution of the measurement is higher than 1 mm. But in some practical situations, the target object needs higher resolution with fast way, such as 3D imaging, recognition and tracking. In these cases, the detection imaging system will be replace as new detector, but means of reflection detection would never be replace. Meanwhile, the terahertz wave can be a powerful and easy way for imaging and detecting the composite materials.

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