Epitaxial Growth, Chemical Doping and Electronic Applications of Two-Dimensional Transition Metal Dichalcogenides

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Epitaxial Growth, Chemical Doping and Electronic Applications of Two-Dimensional Transition Metal Dichalcogenides

A thesis presented

by

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to

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Abstract

Since the memorable discovery of graphene, enormous studies have been performed on atomically thin two-dimensional (2D) materials, such as zero-gap graphene, semiconducting transition metal dichalcogenides (TMDs), and insulating hexagonal boron nitride (h-BN). From the extensive research, many unique physical properties of these 2D materials have been revealed, and recently some of them can be synthesized in a controlled manner using chemical vapor deposition (CVD) method. Especially, the TMDs are known to have superior semiconducting transport property together with interesting electronic behavior, making them promising candidates for post-silicon electronics. However, in spite of the importance of preparation method to obtain highly crystalline and wafer-scale TMD semiconductors, still there are several issues to be solved, such as crystal defects including grain In this thesis, epitaxial growth of highly crystalline WS₂ using hydrogen gas is boundary. By strong reducing effect of hydrogen gas, crystal quality of WS_2 is obviously demonstrated. improved. In addition, by realizing epitaxial growth on c-plane sapphire, formation of grain boundary is significantly suppressed. The epitaxial growth resulted in atomically smooth stitching between merging grains and improved the chemical stability as well as electrical transport property.

C-plane sapphire is one of the most widely used growth substrate for the CVD growth of TMDs due to their similar hexagonal lattice structures. However, despite the importance of growth substrate in CVD synthesis, there is almost no report studying how the sapphire substrate changes the growth mechanism. Therefore, in this thesis, influence of c-plane sapphire on the growth behavior of WS₂ is investigated. More lattice strain was found to be induced in the epitaxially grown WS₂ grains by strong van der Waals interaction with underlying sapphire and it resulted in preferential overlayer deposition and photoluminescence (PL) quenching.

Most of semiconducting TMDs including MoS₂ and WS₂ show unipolar n-type electrical transport property. However, for practical applications, both p- and n-type transport materials are required. In this thesis, by using chemical doping technique, ambipolar CVD-grown WSe₂ is selectively converted to p- and n-type. The chemical doping significantly improved the effective carrier mobilities up to three to four orders of magnitude. Using p- and n-type doped WSe₂, a complementary metal-oxidesemiconductor (CMOS) was fabricated which shows outstanding performance. Finally, by combining chemical doping and lithography techniques, a p-n junction was fabricated within a single triangle grain of WSe₂ which shows clear rectification behavior and light response.

This thesis presents the studies on 2D TMDs from their bottom-up synthesis using CVD method to their application for advanced electronics, which proves the prospect potential of 2D materials in the coming era.

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Chapter 1

Introduction

1.1. Tow-dimensional (2D) Materials

1.1.1. Graphene

Since the memorable discovery of graphene in 2004 with its impressive electrical conductivity,¹ extensive studies have been done and accumulated so far. Graphene is a single layer of carbon sheet, in which the carbon atoms are connected in-plane by covalent bond and additional π bond while having no chemical bond in the out-of-plane direction (Figure 1.1a). When graphene layers are stacked, only weak van der Waals interaction between layers holds their layered structure. A. K. Geim and K. S. Novoselov used an adhesive tape to detach pieces of graphene layers from the bulk graphite and measured their electrical property. The monolayer graphene with subnanometer thickness showed very high carrier mobility ~200,000 cm²/Vs, and this attracted huge attention from researchers.² Since this discovery, many theoretical and experimental studies with interesting expectation and measurement were reported for several years. In spite of the superior physical properties of graphene, however, there was a lacking point in graphene that is the gapless nature. Having high electrical and thermal conductivities³ together with mechanical flexibility and transparency,³ graphene can be used for various applications in electronics. However, graphene does not have a band gap so that it is difficult to be used for semiconducting devices. Therefore, by the dedication of



Figure 1.1. Schematics of crystal structures of (a) graphene and (b) tungsten disulfide (WS₂).

many researchers, several methods have been studied to open the band gap in graphene. First, applying top and bottom dual electrical field to AB stacked bilayer (2L) graphene and ABC stacked trilayer (3L) graphene was found to open a tunable band gap (up to 250 meV for bilaver).^{4,5} Second, by confining the lateral size of graphene down to $10 \sim 100$ nm which is called a nanoribbon, small band gap can be opened (~200 meV at 20 nm width).⁶ Third, by inducing lattice strain manually or using epitaxial substrate, such as SiC, the sublattice symmetry of graphene can be broken, opening the band gap (~300 meV under 1% tensile strain⁷, \sim 260 meV on SiC⁸). Also, hydrogenation or chemical substitution doping can be used to break the sublattice symmetry of graphene and open a small band gap.⁹⁻¹² Formation of heterostructure with hexagonal boron nitride (h-BN) was also found to open a small band gap by forming superlattice structure.^{13,14} Thanks to the huge amount of research for more than a decade, now band gap can be opened in graphene. However, there are still critical limitations of them, because most of the methods need extremely delicate experimental manipulation, and the size of the band gap is too small to be used in semiconducting devices. As a result, research interest was extended from graphene to other atomically thin two-dimensional materials which have an intrinsic semiconducting nature.

1.1.2. Transition metal dichalcogenides (TMDs)

Transition metal dichalcogenides (TMDs) have a MX₂ composition with transition metal (M), such as Mo and W, and chalcogen (X), such as S, Se, and Te. Monolayer TMDs consist of a single layer of transition atom sheet which is sandwiched by single layers of chalcogen sheet (Figure 1.1b). Since the first demonstration of good semiconducting transport property of monolayer MoS₂,¹⁵ TMDs have attracted great attention. Thus, other TMDs, such as WS₂, MoSe₂, WSe₂, MoTe₂, WTe₂ have also received great attention, and a lot of their interesting physical properties have been reported so far. The biggest distinguishing property of TMDs from the graphene is that they have a sizable band gap.¹⁶ Depending on atomic composition, TMDs have different band gaps, as shown in Figure1.2.¹⁷ Due to this semiconducting property, TMDs can be used for various applications in electronics and optoelectronics, such as switching transistors, logic gates, sensors, energy harvesting, light emitting diode, and so on.¹⁸⁻²⁰

Similar to graphene, in the beginning of research, mechanical exfoliation method was widely used to prepare TMD flakes. Using this method, physical properties of various TMD



Figure 1.2. Band gap energy and corresponding wavelength of representative semiconducting monolayer transition metal dichalcogenides.

materials were well studied and diverse electronic and optoelectronic devices were fabricated and investigated. As the research advances, however, necessity for large size TMD crystals with uniform thickness became strong, because the exfoliated crystals have a limit in their crystal size and thickness uniformity. Therefore, bottom-up synthesis using chemical vapor deposition (CVD) method was introduced also for TMDs.²¹ Using CVD method, it became possible to synthesize various TMD crystals by choosing suitable precursors and substrates. However, it accompanies prolonged effort to successfully grow the desired materials. For example, there are many growth parameters which need to be optimized, such as growth temperature, time, pressure, the balance between amount of each precursor, position of growth substrate, and etc. More details about the CVD method are explained in the following chapter. From devoted works by researchers,^{22–27} CVD technology has been dramatically developed and the size of grown crystal has been scaled up to millimeter size.^{28,29} Thanks to the CVD technology, TMD research has further advanced, fabrications of wafer scale logic circuits and transistor arrays have been reported recently.^{30–33}

Compared with typical three dimensional (3D) semiconducting materials, such as Si, Ge, and InGaAs, TMDs have a notable superiority in their physical properties. In modern society, field-effect transistor (FET) is the basis of the information storage and processing. To increase the storage capacity and processing speed, the channel length of FET device needs to be shortened. However, in case of typical 3D semiconductors, as the channel length becomes shorter, the electrons finds it harder to travel through the channel as the imperfection of crystal structure scatters their flow. In contrast, in case of two dimensional (2D) semiconductors, they have a limited vertical thickness which is decided by the number of layers, and a flat lateral surface allows smooth electron flow with less scattering. Therefore, the advantage of 2D semiconductors against the short channel effect in FET application together with other superior physical properties, make TMDs a promising candidate for post-silicon electronics, although

this research is in an early stage yet.³⁴

1.1.3. Heterostructures

For special purposes, more than two different 2D materials are combined in a controlled manner, which is called heterostructure. By stacking different materials one to another, vertical heterostructure can be formed (Figure 1.3a). Thanks to the unique layered structure of 2D materials, each layer can be easily detached and stacked. Vertical stacking is useful when forming material junctions. By stacking p-type or ambipolar layer (WSe₂ or MoSe₂) on n-type layer (WS₂ or MoS₂), a p-n junction is easily formed (Figure 1.3b).^{35–38} Also, stacking of metallic (graphene) and semiconducting (WS₂) layers simply forms the metal-semiconductor junction.¹⁸ Furthermore, the relative stacking orientation between upper and



Figure 1.3. (a,c) Illustrations of (a) vertical- and (c) in-plane-heterostructures of TMDs. Blue, green spheres correspond to transition metals and yellow, orange spheres correspond to chalcogen atoms. (b) Optical micrograph of WSe₂/WS₂ stacked heterostructure on SiO₂/Si. (d) Scanning tunneling microscope image of MoS₂/WS₂ in-plane heterostructure. Figure 1.3(b) and (d) are reproduced from Ref. 38 and 48, respectively.

lower layers is known to effectively change the interlayer interaction.³⁸⁻⁴¹ For example, when MoS₂ is stacked on graphene forming MoS₂-graphene heterostructure, the optical property of MoS₂ is strongly affected by the relative orientation with the underlying graphene.⁴² More notably, some vertical heterostructures show appearance of new physical property, such as superconductivity or phase change at specific stacking configuration, which disappears when stacking configuration is altered.⁴³⁻⁴⁶ There is also an in-plane heterostructure (Figure 1.3c). In-plane heterostructures can be formed by consecutive growth of different materials. For example, by switching the precursor species from W and S to Mo and S during the growth, it is possible to make WS₂-MoS₂ in-plane heterostructure (Figure 1.3d).⁴⁷⁻⁴⁹ In a similar way, by performing consecutive growth of graphene and MoS₂, graphene-MoS₂ heterostructure can be formed.²⁴ Recent development of metal-organic CVD system made it easier to form in-plane TMD heterostructures.⁴⁸ Heterostructure itself is a very interesting research topic not only because of their usefulness in forming material junctions, but also because of the appearance of new physics with orientation-dependent tunability.

1.2. Motivation

As described in the previous section, superior physical properties of TMDs have attracted great attention from researchers. As a result, a number of studies have been reported on the characterization and CVD synthesis of TMD materials. With the advance of research, need for uniform and large size TMD crystals became stronger, because the mechanically exfoliated flakes are too small and lack of thickness uniformity. Therefore, CVD growth technique rapidly developed for last several years and now millimeter size of single crystals can be synthesized.^{28,29} To fully extract the superior property of TMDs, the crystallinity of materials

must be high enough. However, compared with exfoliated ones, CVD-grown TMD grains contain much higher density of structural defects and, consequently, CVD-grown TMDs show lower device performance compared with exfoliated counterparts.⁵⁰⁻⁵² Therefore, this crystallinity issue in CVD-grown TMDs needs to be improved. In electronic and optoelectronic applications, wafer-scale TMD films are required to fabricate devices in more systematic manner. For this, CVD method has already proved their capability of growing wafer scale continuous films of various TMDs.^{53,54} Although large-scale TMD films could be grown by CVD, still there is a grain boundary issue. During growth, as the grains are nucleated and their lateral size gradually increases, the neighboring grains start to touch each other and finally become merged. When the neighboring grains are merged, their crystal structures are deformed along the boundary, which is called grain boundary (GB). GBs are known to cause severe property variation and degradation.^{55,56} Therefore, suppressing the formation of GBs is an important issue in improving the quality of CVD-grown materials, and it can be realized by aligning the crystal orientation of grains on the substrate. In this thesis, by using H₂ gas which has a strong reducing effect on the surface of growth substrate (c-plane sapphire), epitaxial growth was achieved which effectively suppressed GB formation. The smooth stitching between merging grains was found to improve the chemical stability and electrical transport properties on the merged grains. The atomic structures of merged interfaces were studied in detail using scanning transmission electron microscopy (STEM).

To realize the CVD growth of highly crystalline large size single crystal TMD films, extensive effort has been devoted by many researchers. There are many numbers of CVD growth parameters, such as reaction temperature, time, precursor, carrier gas, and substrate. In previous literatures, many parameters have been investigated to understand how they affect and change the resulting TMDs.^{57–60} In case of growth substrate, the most widely used ones are c-plane sapphire and silicon oxide. The c-plane sapphire has a hexagonal lattice structure

same as MoS₂ and WS₂,⁶¹ therefore, it is expected that this similarity of lattice structures would influence the growth behavior of TMD crystals on it. However, still there is almost no study about this. Thus, in this thesis, I investigated how the c-plane sapphire influences and changes the growth of WS₂. Due to the similar hexagonal lattice structure of c-plane sapphire and WS₂, more lattice strain was found to be induced in epitaxially grown grains by strong van der Waals interaction. This large strain caused preferential overlayer deposition on the epitaxial WS₂ grains. Low-temperature optical study revealed that photoluminescence (PL) is strongly quenched in epitaxially grown WS₂ grains by interaction with the underlying sapphire.

To realize practical device applications of semiconducting TMDs, such as logic circuits and a p-n junction, both p- and n-type materials are required. However, most of TMDs (MoS₂, WS₂, MoSe₂, WSe₂) are known to have n-type or ambipolar transport properties and p-type materials are scarce.⁶² Therefore, various methods have been proposed to convert their electrical transport property to p-type, and the representative ones are the chemical doping,^{62,63} substitutional doping,⁶⁴⁻⁶⁶ metal nanoparticle deposition⁶⁷, and tuning Schottky barrier by changing metal electrode.^{68,69} There are several methods to achieve the substitutional doping, such as post-growth or during-growth processes, however, most of them cause severe degradation of the semiconducting property.^{64–66} Deposition of metal nanoparticles was found to be effective to convert the transport polarity to p-type.⁶⁷ But the nanoparticles degrade the optical property of underlying TMDs, which is critical for optoelectronic Metals have different Fermi levels so that the use of different metal as an applications. electrode can be employed to tune the electrical transport behavior of TMD channels.^{68,69} However, in this case, fabrication process becomes complicated requiring deposition of different metals for each TMD. In contrast, chemical doping does not degrade semiconducting and optical properties, and moreover, its experimental process is relatively simple as compared with other methods. Actually, some exfoliated TMDs were well

converted to p-type by chemical doping.⁶² However, when considering the higher density of chalcogen vacancy defects in CVD-grown TMDs compared with exfoliated flakes which causes n-type doping, it is more difficult to convert their carrier transport to p-type. Thus, in this thesis, chemical doping method is used to control the electrical transport polarity of CVD-grown WSe₂. Two different dopant molecules, 4-nitrobenzenediazonium tetrafluoroborate (4-NBD) and diethylenetriamine (DETA), were used for p- and n-type doping, respectively, and it was found that the effective carrier mobilities increase up to three to four orders of magnitude after each doping treatment. Using p- and n-type doped WSe₂, fabrication of a complementary metal-oxide-semiconductor (CMOS) circuit was demonstrated, showing good performance. In addition, by combining our chemical doping and lithography techniques, a p-n junction was fabricated within a single triangle grain of WSe₂, with clear rectification behavior as well as light response.

1.3. Outline of thesis

This thesis covers three main parts, CVD growth of high quality epitaxial WS_2 , investigation of influence of c-plane sapphire on growth behavior of WS_2 , and chemical doping of CVDgrown WSe_2 for controlling its electrical transport polarity and their device applications. Before discussing the experimental result in the main parts, basic introduction about 2D materials has been given in this Chapter (Chapter 1). For better understanding of experimental result in Chapters 3 to 5, more detailed background about physical properties and preparation methods of TMD materials are described in Chapter 2. Chapter 3 presents the epitaxial growth of high quality WS_2 using H_2 gas. In Chapter 4, influence of c-plane sapphire substrate on the growth behavior of WS_2 is investigated. Chapter 5 describes the chemical doping of CVD-grown WSe₂ for controlling its electrical transport polarity where significantly improved carrier mobilities as well as fabrication of advanced electronic devices are demonstrated. Finally, this thesis is closed with conclusions and future outlook in Chapter 6.

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Chapter 2

Property and Preparation of TMD Materials

2.1. Crystal structures and physical properties of TMDs

2.1.1. Crystal structures

Transition metal dichalcogenides, such as WS₂, MoS₂, and WSe₂, have atomically thin twodimensional structures composed of single layer atomic sheet of transition metal (W, Mo) which is sandwiched by two layers of chalcogen atom (S, Se, Te) layers, having three fold symmetry (see Figure 1.1b). They exist as polytype phases with different crystal and electronic band structures. As shown in Figure 2.1, for example, the TMDs with the 2H phase have a semiconducting property, while those with the 1T phase have a metallic property.^{1–3} In nature, semiconducting 2H phase is energetically more favorable, and artificial treatments, such as lithium intercalation shown in Figure 2.2, are necessary to obtain metallic property by conversion to 1T phase.^{4–6} Thus, all TMDs dealt with in this thesis belong to 2H phase with semiconducting property.

Depending on atomic composition of each TMD, they have slightly different lattice constants. Lattice parameters a (= b), c (indicated in Figure 2.3) of WS₂ is 3.15, 12.32 Å, MoS₂ is 3.15, 12.3 Å, and WSe₂ is 3.28, 12.95 Å, respectively.^{7,8} TMDs have strong in-plane ionic-covalent bonds among constituent atoms, while there is no chemical bond between adjacent layers. Between layers, there is weak van der Waals interaction which holds adjacent layers bound and



Figure 2.1. Top and side view of 2H and 1T MoS₂, and their band structures. Cyan and yellow atoms correspond to Mo and S, respectively. Reproduced from Ref. 3.



Figure 2.2. (a) An optical image of a monolayer WS_2 butterfly transferred on a 300 nm SiO₂/Si substrate. Red dash line indicates the boundary between the left and right wings. The surface was spin coated with PMMA (polymethyl methacrylate) and an e-beam lithographical window was opened in the lower half of the WS₂ butterfly. White dash line indicates the unmasked edge on the WS₂ butterfly. Four quadrants: A, B, C, and D are assigned to the parts of WS₂ butterfly. (b) An annular dark field STEM image of the WS₂ butterfly taken at the A/B domain boundary in (a), where the quadrant-A shows 1T phase, while the quadrant-B shows 1H phase. Reproduced from Ref. 6.



Figure 2.3. Lattice parameters *a*, *b*, and *c* of transition metal dichalcogenides. Grey and yellow spheres correspond to transition metal and chalcogen atoms, respectively.

makes a layered structure. Due to this unique 2D structure, TMDs show layer numberdependent changes in their physical properties, which is sometimes called as the layer number degree of freedom. More details are explained in the following chapters.

2.1.2. Physical properties

2.1.2.1. Phonon modes

TMDs have distinct Raman scattering not only depending on their atomic composition, but also on the number of layers. Having different atomic mass, interatomic interaction, and lattice parameters depending on atomic composition, each TMD has different phonon modes with distinctive vibrational energies (see Figure 2.4). In addition, it also changes depending on the number of layers. With unique atomically thin layered structure, phonons in monolayer TMD is strongly affected when additional layers are attached on it. Consequently, the layer number-dependent phonon scattering property becomes one of the strongest tools to distinguish the number of layers (thickness) of TMDs. Figure 2.5 shows layer number- and excitation energy-dependent Raman spectra and peak frequencies of WS₂.⁹ For example, both WS₂ and MoS₂ have E^{1}_{2g} and A_{1g} phonon modes that correspond to in-plane and out-of-plane vibration,



Figure 2.4. Raman spectra of monolayer WS₂, MoS₂, and WSe₂ measured using 473 nm excitation.



Figure 2.5. (a) Representative Raman spectra of monolayer WS_2 excited using 514.5 nm and 488 nm laser. (b) The layer number-dependent shift of A_{1g} , E^{1}_{2g} , and 2LA peaks of WS_2 . Reproduced from Ref. 9.

respectively, and the separation between those mode of around ~ 60 and ~ 20 cm⁻¹ are known to correspond to single layers.¹⁰ Additionally, use of different excitation energy is known to cause appearance of additional vibrational modes.⁹ Therefore, Raman spectroscopy using

suitable excitation energy becomes very quick and efficient way to identify the TMDs and distinguish the thickness of TMD materials.

2.1.2.2. Electronic band structure

As aforementioned, TMDs exist as polytype phases. The distinctive electronic band structures of 2H MoS₂ with sizable band gap at *K* valley and 1T MoS₂ without band gap are shown in Figure 2.1.³ Also, the calculated layer number-dependent electronic band structures and thickness-dependent band gap change of representative TMDs are shown in Figure 2.6.¹¹ Monolayer TMDs have the largest band gap at *K* valley with direct transition from conduction to valence band. As they become thicker than monolayer, however, the band gap is gradually reduced and the transition changes to be indirect followed by the change of the global band extrema depending on the number of layer.¹² Thus, the monolayer TMDs with a broad choice of band gap size and direct transition become a good base to study optics and optoelectronics of 2D materials.

2.1.2.3. Spin degree of freedom

Different from graphene which consists only of carbon, TMDs have diatomic structures composed of transition metal and chalcogen atoms, thus, the in-plane inversion symmetry is broken in monolayer TMDs (Figure 1.1). The outer layers of electronic state of TMDs consist of d-orbital of transition metal atoms so that this together with the broken inversion symmetry makes electrons in monolayer TMDs experience the asymmetrical potential gradient. Owing to the asymmetrical in-plane electric field together with the resultant out-of-plane Zeeman-like effective magnetic field, spin-orbit coupling-induced spin-orbit splitting occurs in conduction



Figure 2.6. (a) Calculated electronic band structures of representative TMDs with different thicknesses. 1L and 2L mean mono- and bi-layer. (b) Thickness-dependent band gap change of representative TMDs. Reproduced from Ref. 11.



Figure 2.7. (a) Calculated electronic band structures of monolayer MoS_2 with and without spin orbit coupling (SOC). (b) Band diagram at *K* valley with spin-orbit splitting in conduction and valence bands. Electron with spin-up state has slightly smaller transition energy (E_g) compared with electron with spin-down state. Figure 2.7a is reproduced from Ref. 13.

and valence bands.^{13–15} Figure 2.7a shows the calculated electronic band structure of monolayer MoS₂ with and without spin-orbit coupling and the band splitting at *K* point can be seen when spin-orbit coupling is induced.¹⁴ The split energy states at *K* and *K'* valleys have sizable separation, especially in valence band (~ 400 meV in WS₂).¹³ The electron residing in each split state has an opposite spin state, consequently, the spin of electrons can be controlled by choosing the excitation energy (see Figure 2.7b). This spin controllability of electrons (additional spin degree of freedom) in TMDs allows additional data storage in electrons.

2.1.2.4. Valley degree of freedom

Due to the diatomic composition and broken inversion symmetry in monolayer TMDs, spinorbit splitting occurs in conduction and valence bands. Also, due to the broken inversion



Figure 2.8. Electronic band dispersions of monolayer TMDs near *K* and *K*' valleys of Brillouin zone. The interband transition in each valley has opposite helicity and optical selection rule.

symmetry in monolayer TMDs, degenerate inequivalent *K* valleys (*K* and *K'*) exist in 2D hexagonal Brillouin zone, as shown in Figure 2.8.¹⁶ Electrons at respective *K* and *K'* valleys experience the same size but oppositely directed out-of-plane Zeeman-like efficient magnetic field. Therefore, electrons at *K* and *K'* valleys are oppositely spin polarized and have different optical selectivity, electrons at *K* (*K'*) valley interacting with right- (left-) circular polarized light, respectively.^{13,14,17} This valley-dependent spin polarization together with the opposite optical selectivity of electrons allow additional degree of freedom, which can be used for additional information storage in carriers.
2.1.2.5. Optical property

2.1.2.5.1. Direct and indirect transition

The band gap of TMDs shows interesting change in their transition mechanism from direct (monolayer) to indirect as get thicker (see Figure 2.6). Also, the size of band gap gradually decreases as the number of layers increases. When thicker than monolayer, the indirect transition of electrons from conduction to valence band accompanies phonon-assisted momentum change which dissipates electrons' energy as heat. As a result, the optical efficiency is highest in monolayer (*K-K* direct transition) and exponentially deteriorates with increasing the thickness. Regarding the PL efficiency of TMDs, up to 60 % of PL quantum yield was reported for CVD-grown WS₂.¹⁸ The broad range of band gap energy (0.95 ~ 2.02 eV) of various TMD materials offers flexible choice of active material in optoelectronic engineering depending on the purpose, and it makes TMDs promising materials for optoelectronics.¹⁹

2.1.2.5.2. Spin-valley polarization

As described in previous sections, monolayers of TMDs possess inequivalent degenerate K and K' valleys in momentum space with global extrema located on it. Having oppositely spinpolarized electrons, each K valley has different optical selectivity. Therefore, by using rightor left-circular polarized excitation laser, it is possible to selectively shine the desired valley and obtain the spin-valley polarized excitons.^{20–25} Originating in the large exciton binding energy (~ hundreds of meV) and long exciton lifetime by large spin-orbit splitting, high efficiency of spin-valley polarization has been reported even at room temperature.²⁶

2.1.2.5.3. Interlayer exciton

Strong quantum confinement effect in monolayer TMDs results in various interesting optical properties. Stacking two layers of TMDs offers a new opportunity to obtain exotic physical properties. Once two different TMD monolayers are stacked with type II band alignment as shown in Figure 2.9, excitons can be formed between the layers which is called interlayer exciton.^{27,27–29} Formation of interlayer exciton relies not only on the band alignment, but also on the lattice mismatch and electronic band structures.³⁰ Once interlayer excitons are formed, their energy is determined by the electronic band structures of substituent monolayers. Therefore, by choosing suitable combinations of substituent monolayers, the energy of interlayer excitons can be engineered.³¹



Figure 2.9. Type II band alignment between MoS_2 and WSe_2 . Blue and green arrows represent intralayer transitions within MoS_2 and WSe_2 , respectively. Red arrow indicates interlayer transition which accompanies tunneling of carriers between layers.

2.1.2.5.4. Exciton control

Due to the large exciton binding energy and spin-valley degree of freedoms in electron, TMD materials become an interesting platform to study the spin-valley optics. In TMD heterostructures, interlayer excitons can be formed by spatially separated electron and hole pair staying at respective layers. Different from the intralayer excitons, the interlayer excitons have vertical dipole moment which causes Stark effect. Therefore, by inducing vertical electric field to the heterostructure with interlayer excitons, it is possible to further control the energy of interlayer excitons.^{32,33} Figure 2.10 represents the energy change of interlayer exciton In MoSe₂-WSe₂ heterostructure depending on the applied vertical electric field. In addition, spatial separation of electron and hole pair of interlayer exciton results in much longer



Figure 2.10. The energy of interlayer (up) and intralayer (down) excitons versus electric field applied to $MoSe_2$ -WSe_2 heterostructure. The top inset is an optical image of a representative device. The bottom inset is schematic of the heterostructure cross section. The white arrow represents positive direction of electric field. Reproduced from Ref. 33.

exciton lifetime compared with intralayer one. This long lifetime allows microscale lateral diffusion of excitons even at room temperature, in which the diffusion direction can be further controlled by electrical manipulation.^{33,34} Moreover, the flow of valley-polarized excitons excited by circularly polarized laser could be controlled by inducing electrical potential.³⁵ The operation of exciton transistor devices made of bulk semiconductors-based coupled quantum wells were already reported.^{36,37} However, they require low temperature operation, which is not suitable for practical applications. In contrast, long lifetime of interlayer excitons in TMD heterostructures makes it feasible to realize room-temperature operating TMD-based exciton transistors. Therefore, TMDs are promising optoelectronic materials although more studies and investigations are necessary.

2.1.2.6. Electrical transport

2.1.2.6.1. Field-effect transistor

Having sizable band gaps, TMDs can be used for switching transistors. Since the first demonstration of single layer MoS₂ transistor that showed impressive performance (carrier mobility (μ) ~ 217 cm²V⁻¹s⁻¹),³⁸ the electrical transport of TMD materials has been actively investigated. For evaluating the electrical transport properties, usually bottom- or top-gated FETs are fabricated followed by 2- or 4-probe measurement. Figure 2.11 depicts three typical device structures used for TMD FETs. The carrier mobility is extracted using the following equation.

$$\mu = \frac{g_{\rm m}}{C_{\rm ox}V_{\rm d}} \frac{L}{W} \tag{2.1}$$

where $g_m = dI_d/dV_g$ is the transconductance obtained from the transfer curve; C_{ox} is the capacitance per unit area of insulator; V_d is the applied drain voltage; L and W represent the



Figure 2.11. Schematics of (a,b) bottom-gated FETs on SiO₂/Si substrate with 2 and 4 electrodes. (c) Top-gated structure with additional insulating layer and top electrode deposited on the channel.

effective length and width of the channel.

Generally, mechanically exfoliated flakes show better transport and switching performance compared with CVD-grown samples. This is because the native crystal defects exist in CVD-grown materials, which consist of sulfur defect, broken structure, undesired extrinsic substitutional doping, or grain boundary, trap and scatter the carriers hindering their flow.^{39–46} Many theoretical calculations expected promising electrical transport properties for WS₂, MoS₂, and WSe₂.^{47,48} However, still the experimental demonstration, especially when using CVD-grown materials, cannot keep up with the theoretical values and there are several reasons need to be noted. First, as already mentioned, the crystal quality needs to be high enough to suppress the defect-induced phonon scattering. Second, for efficient carrier injection from electrode to channel, the electrode contact needs to form a good Ohmic contact without Schottky barrier.^{49,50} Third, due to the atomically thin structure of 2D materials, they are vulnerable to surrounding dielectric environment. Any undesired charge impurities around the channel can strongly affect the device performance. Therefore, removing the charge

impurities in the oxide (insulating) layer of transistor device or protecting TMD channels with highly dielectric, covalent bonding-free h-BN will help extract the intrinsic properties of TMDs.^{51–53}

Depending on atomic composition, each monolayer TMD has different electrical transport polarity. For example, MoS₂ and WS₂ typically show n-type transport, while WSe₂ and MoSe₂ show ambipolar transport behavior although the polarity of majority carriers changes depending on measurement environment. Figure 2.12 indicates the transfer characteristics of various exfoliated TMDs and their polarity change by chemical doping.⁵⁴ The transport polarity is important when fabricating electronic devices, such as p-n junction and logic circuits. For these applications, both of n- and p-type channels are necessary. However, there is almost no p-type TMDs and few of them are toxic or air-unstable. As a result, several methods have



Figure 2.12. Representative transfer characteristics of a pristine MoS_2 , WS_2 , $MoSe_2$, and WSe_2 FETs before and after doping using n-type dopant ($C_{30}H_{40}Rh_2$ (RhCp*Cp)₂) or p-type dopant ($[N(C_6H_4-p-Br)_3]SbCl_6$) of various concentration and exposure time. Reproduced from Ref. 54.

been reported to control the electrical transport polarity of TMD sheets, such as substitutional doping, ^{55–57} chemical doping (see Figure 2.12)^{54,58} and tuning the Schottky barrier by changing the electrode metals.^{59,60}

2.1.2.6.2. Logic circuits

Based on good transport and switching performance, logic circuits become one of the most promising applications of TMD materials. The importance of fabricating high-performance logic gates, such as CMOS, NOT, OR, AND is needless to be emphasized. As shown in Figure 2.13, various logic gates have been fabricated using exfoliated MoS₂, WSe₂, and MoTe₂, which showed good performance.^{61–65} Also, CVD-grown WSe₂ and MoS₂ were used to fabricate a flexible CMOS inverter on a polyimide film as shown in Figure 2.14a, which showed high voltage gain of ~30 (Figure 2.14b).⁶⁶ When fabricating logic circuits, use of pure p-type channel together with typical n-channel is beneficial for improving the device performance, as compared with when ambipolar channel is used as p-channel or both channels are n-type.⁶⁶ Therefore, asymmetric chemical doping was introduced to achieve CMOS operation by forming p- and n-doped MoS₂ channels. In similar tactics, chemical doping of WSe₂^{67,68} and defect-engineered p-doping of MoS₂ were reported.⁶⁹ Thus, TMD materials have bright promise not only in applications of typical logic gates but also in flexible and transparent electronics.



Figure 2.13. Three kinds of fundamental logic units based on different configuration of MoS_2 and WSe_2 . Optical images, logic circuit diagrams, and schematic diagrams of (a, b, and c) complementary logic transmission gate, (d, e, and f) inverter, and (g, h, and i) Schmidt-like flip-flop. Reproduced from Ref. 62.



Figure 2.14. (a) Schematic illustration and optical image of high-gain quasi-CMOS inverter made of WSe₂ monolayer film on a flexible polyimide substrate. (b) $V_{in}-V_{out}$ transfer characteristics of WSe₂ inverter at different values of curvature radius *R*. Reproduced from Ref. 66.

2.1.2.6.3. Valley Hall effect

As mentioned in the previous section, one of the greatest interesting physical properties of monolayer TMDs is the additional spin-valley degree of freedoms of electrons. The existence of spin and valley degree of freedoms together with electric charge in electrons results in valley Hall effect (VHE).⁷⁰ When circularly polarized laser is illuminated on the monolayer TMD which has a broken inversion symmetry, valley-dependent (*K* and *K*^{*}) oppositely spin-polarized electrons are excited. Once the channel is biased, electrons at each *K* valley experience oppositely directed Lorentz-like force, thus, move to the opposite directions perpendicular to the drift current in the channel.⁷⁰ Figure 2.15 illustrates the circularly polarized laser-induced valley Hall effect. In 2014, observation of the VHE from monolayer MoS₂ was reported.⁷¹ In case of bilayer MoS₂, however, VHE was not observed reflecting the preserved inversion symmetry. More recently, VHE of interlayer exciton was observed from MoS₂-WSe₂ heterostructure, even at room temperature.⁷² The VHE in TMD materials implies their promise to be used in valleytronic applications for information storage and processing.



Figure 2.15. Illustration of circularly polarized excitation-induced valley Hall effect. Oppositely circular polarized lasers induce selective excitation in K or K' valley. By opposite spin polarization of electrons in each valley, opposite Hall voltage is induced and electrons move to opposite direction on the channel perpendicular to the drift current.

2.1.2.7. Twistronics

Since the observation of magic angle-induced superconductivity in bilayer graphene,⁷³ a new field of research (called Twistronics) has emerged, which investigates the stacking of layered materials with controlled configuration and investigating their physical properties. Figure 2.16 indicates an example of this kind of study, where the rotation angle-dependent change of Raman 2D peak of graphene-hBN heterostructure was investigated.⁷⁴ As other 2D materials, vertical stacking of different TMD layers is known to form unique superlattices together with other interesting physical properties that are sensitively affected by tiny change of twist angle. For example, MoSe₂-WSe₂ and WSe₂-WS₂ heterostructures showed stacking angle-dependent valley exciton trapping in Moiré superlattice potential.^{75,76} Also, Mott insulating phases were observed in WSe₂-WS₂ heterostructure at a specific stacking angle.⁷⁷ In case of MoS₂-WS₂



Figure 2.16. (a) Schematic of device structure and experimental technique using rotatable hexagonal boron nitride (BN) on graphene. (b-d) AFM image of a fabricated device showing three different orientations of top BN. Scale bar is 1 μ m. (e) Schematic illustration of Moiré superlattice arising between graphene (red) and BN (blue) at 0° and 30°. (f) Raman spectrum of orientation-variable heterostructure shown in (b-d). (g) FWHM of Raman 2D peak as a function of stacking orientation angle. Reproduced from Ref. 74.

and MoSe₂-WSe₂, atomic reconstruction and corresponding change in electronic band structures were observed at small stacking angles ($< 2^{\circ}$).^{78,79} Therefore, combination of materials together with controlling their stacking configuration open a new world of physics. Here, the materials are not limited to semiconducting TMDs. The heterostructure of ABCtrilayer graphene and h-BN was found to form a special Moiré superlattice and showed tunable phase change between Mott-insulator and superconductor depending on temperature and electromagnetic environment.⁸⁰ Therefore, combining TMDs with graphene, h-BN, or other 2D materials is also expected to be an interesting research topic.^{81–85}

2.2. Preparation of TMD materials

2.2.1. Mechanical exfoliation

In the beginning of graphene research, an adhesive tape was used to detach thin pieces of layered flakes from the bulk crystal. A similar method has been applied to the TMD research as well.⁸⁶ By detaching layered crystals repeatedly using the adhesive tape, thin flakes with a thickness of ~ nm can be obtained and easily transferred on target substrates such as SiO₂.⁸⁶ Figure 2.17 shows a schematic of mechanical exfoliation and successive transfer. This simplicity is one of the advantages of the mechanical exfoliation method. Another advantage is the high crystal quality of obtained flakes which is determined by the crystal quality of the bulk. Using this method, fundamental properties of various 2D materials including TMDs could have been investigated in details and various types of devices have been fabricated and studied. However, there exist several critical limitations in this mechanical exfoliation method. First, the size of the exfoliated flakes is very limited up to tens of μ m, which does



Figure 2.17. Mechanical exfoliation of 2D crystals. (a) Adhesive tape is pressed against a 2D crystal so that the top few layers are attached to the tape (b). (c) The tape with detached layered crystals is pressed against a target substrate. (d)Upon peeling off, the bottom layer is left on the substrate. Reproduced from Ref. 86.



Figure 2.18. Optical image of exfoliated WS_2 flakes. Points 1, 2 and point 3 correspond to monolayer and multilayer areas, respectively. Reproduced from Ref. 87.

not fit to wafer scale device fabrication or commercialization of TMDs. Second, the thickness uniformity of obtained flakes is very poor in spite that uniform thickness film is required for systematic characterization or device fabrication. Figure 2.18 displays an optical image of WS_2 flakes prepared by mechanical exfoliation.⁸⁷ The flake thickness is ununiform and the size in only 20 μ m. Therefore, as the research becomes more and more advanced, need for large size TMD films with uniform thickness became stronger. Thus, researchers began to study the bottom-to-up synthesis of TMD films using CVD method.

2.2.2. Chemical vapor deposition (CVD)

2.2.2.1. Conventional CVD

CVD is a bottom-up material synthesis method. By reacting precursors in the optimized condition, desired materials can be synthesized in a controlled manner. For conventional CVD growth of TMDs, typically a tubular furnace is used to heat precursors and growth substrate under the flow of carrier gas. Figure 2.19 shows typical CVD set up used for TMD synthesis, in which metal oxide and chalcogen powers are used as feedstocks. As a transport carrier, inert gases, such as Ar and N₂, are widely used, and mixture with H₂ or O₂ also can be used for special purposes.^{88–90} Depending on target materials, low-pressure or plasma-



Figure 2.19. Illustration representing a typical thermal CVD set up where a tubular furnace is used for heating the precursors and growth substrate. During growth reaction, an inert carrier gas is flowed.



Figure 2.20. Optical images of CVD-grown MoS₂ and WS₂.

enhanced low temperature reactions are employed as well.^{91,92} As precursor materials for typical TMD CVD, metal oxides (WO₃, MoO₃) and chalcogen (S, Se, Te) powders are widely used. Optimization of the amount of each precursor, reaction temperature, reaction time, gas flow rate, positions of precursors, and growth substrate, is performed to obtain TMDs with desired thickness and grain size. One big advantage of the CVD method compared with mechanical exfoliation is the capability of synthesizing large size TMD flakes (films) with much better thickness controllability. For specific purposes, wafer-scale TMD flakes and films with precisely controlled thickness are needed, thus, extensive effort has been devoted by the CVD method.^{93–95} However, several problems and limitations were found in conventional CVD method where the powder type precursors are used. First, high reaction temperature which is required for lateral growth of grains causes excessive density of grain nucleation which hinders the large size growth, making a number of GBs. Figure 2.20 shows the optical images of CVD-grown MoS₂ and WS₂ on c-plane sapphire. Second, as the reaction time is increased for large size growth, the surface of metal oxide precursor is thermally reduced and stiffened, and it causes gradual reduction of supply of metal species to the growth reaction. Consequently, there is a limitation in growth reaction time which results in the size limit of the grains.

2.2.2.2. Alkali halide-assisted CVD

To reduce the high reaction temperature-induced high nucleation density and also to increase the lateral size of grown crystal, use of growth promoters was introduced.⁹⁶ As promoters, alkali-metal halides (MX where M = Na, or K and X = Cl, Br, or I) are used, and they are placed inside of the reaction chamber together with a metal oxide precursor. Figure 2.21 shows the schematic of CVD set up where the alkali halide powders are mixed with metal



Figure 2.21. Illustration representing the alkali-halide-assisted CVD set up where alkali-halide powder is placed together with metal oxide precursor.



Figure 2.22. (a,b) Schematic of alkali-halide-assisted growth reaction. Metal oxychlorides are formed, and these promote the reactions. Chalcogens are not shown here. (1)-(3) The proposed process of the added salt decreasing the melting point of the precursors. (4) Scanning electron microscopic images of Nb nucleus with (left) and without (right) added to salt. (b) (5)-(7) The growth process of 2D atomic layer, with intermediate products. (8) Different single-crystalline monolayers with large size of monolayer and growth time of less than 3 min. Reproduced from Ref. 97.

oxide precursor. The alkali halide modifies the reaction between metal oxide and chalcogen atoms by forming intermediate volatile transition metal-based halide species.^{96,97} Due to the lower activation energy of intermediate species formation, it is possible to nucleate and supply the enough amount of precursor flux to the growth substrate at lower temperature, which can reduce the nucleation density. Therefore, the reduced nucleation density together with enough supply of precursors further assist the lateral growth, which is beneficial for achieving growth of large single crystals and reducing the GBs. Figure 2.22 shows the example of using alkali halides to grow large single crystals of TMDs with decreased nucleation density.⁹⁷ Regarding the use growth promoters, there is a concern about possible substitutional doping and deterioration of crystal quality. However, microscopic, optical, and device studies revealed the high crystal quality free from substitutional doping together with high electrical mobility.^{18,96,98}

2.2.2.3. Metal-organic CVD (MOCVD)

To solve the issue of thermal reduction-induced stiffening of metal precursor which results in limitation of growth reaction time, use of metal-organic type precursors was introduced.⁹⁹ Different from the conventional CVD, in which powder type precursors are used, in this system, metal-organic and organic chalcogen vapors are used as precursors. Figure 2.23 shows a typical CVD set up using metal-organic precursors. By bubbling the metal-organic and organic-chalcogen liquids, we can achieve constant supplies of metal and chalcogen feedstocks. Thus, uniform thickness of continuous TMD films could be obtained.⁹⁹ Due to this sizeunlimited growth capability of metal organic CVD (MOCVD) system, recently, it is regarded as the most promising synthesis method for large-scale production of TMDs. Furthermore, there is an additional advantage of MOCVD system which is the tunability of precursor species during the growth. By connecting multiple precursor source lines to the furnace and switching the precursors during the reaction, it is possible to fabricate heterostructures.^{100,101} By using MOCVD method, various kinds of single-layer TMDs and their heterostructures were fabricated and their physical properties were studied.^{98–101}

Thanks to the significant development of CVD technology, the controllability of grown materials has improved a lot. However, there is still an argument in CVD research regarding the low crystal quality of CVD grown TMDs. For this, several solutions have been reported to increase the crystallinity by finding better composition of carrier gas or using new types of metal-organic precursors.^{89,90}



Figure 2.23. Schematic of MOCVD set up where metal-organic and chalcogen-organic gases are used as precursors. Each precursor is connected to growth chamber with separate line

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Chapter 3

Hydrogen-Assisted Epitaxial Growth of Monolayer Tungsten Disulfide and Seamless Grain Stitching

Abstract

Recently, research on TMDs has been accelerated by the development of large-scale synthesis based on CVD. However, in most cases, CVD-grown TMD sheets are composed of randomly oriented grains, and thus contain many distorted GBs which deteriorate the physical properties of the TMD. In this chapter, the epitaxial growth of monolayer tungsten disulfide (WS_2) on sapphire is demonstrate by introducing a high concentration of hydrogen during the CVD process. As opposed to the randomly oriented grains obtained in conventional growth, the presence of H₂ resulted in the formation of triangular WS₂ grains with the well-defined orientation determined by the underlying sapphire substrate. Photoluminescence of the aligned WS₂ grains was significantly suppressed compared to that of the randomly oriented grains, indicating a hydrogen-induced strong coupling between WS₂ and the sapphire surface that has been confirmed by density functional theory (DFT) calculations. Scanning transmission electron microscope observations revealed that the epitaxially grown WS₂ has less structural defects and impurities. Furthermore, sparsely distributed unique dislocations were observed between merging aligned grains, indicating an effective stitching of the merged grains. This contrasts with the GBs that are observed between randomly oriented grains, which include series of 8-, 7-, and alternating 7/5-membered rings along the GB. The GB structures were also found to have a strong impact on the chemical stability and carrier transport of merged WS₂ grains. This chapter demonstrates a novel method to grow high-quality TMD sheets with much less structural defects, contributing to the future development of TMD-based electronic and photonic applications.

3.1. Introduction

Monolayers of TMDs have been attracting an intensive interest as one of the representative families of 2D materials. In contrast to the gapless nature of monolayer graphene, monolayers of TMDs, such as molybdenum disulfide (MoS₂) and tungsten disulfide (WS₂), have direct band gaps of $1 \sim 2$ eV. Thus, these materials show strong PL in the visible range and also allow excellent semiconductor device operation with high on/off ratios.^{1,2} Large-scale synthesis of TMD monolayers is essential to develop electronic and optoelectronic applications. While recent development of the CVD methods has enabled the wafer-scale production of TMD monolayers,³⁻¹⁰ there still exist several issues that degrade the device performance of the CVDgrown TMD sheets. First, the TMD growth is not based on surface self-limiting mechanism so that the growth occurs not only on the substrate surface but also on the surface of the TMD itself. This makes difficult to synthesize highly uniform monolayer films, as multilayer spots are frequently observed at the center of monolayer grains.^{3–6} Accordingly, many multilayer grains are formed when the TMD growth is continued to fully cover the substrate.^{6,7} Second, the lattice orientation of each TMD grain is not controlled, which results in the formation of polycrystalline TMD sheets having many GBs with a wide range of misorientation angles.¹¹⁻¹⁵ Although the pre-annealing process of sapphire substrates in air was reported to result in aligned growth of MoS₂,¹⁶ this method has not been successfully applied to other TMD

monolayers, such as WS₂. Graphene and h-BN act as good templates to obtain aligned TMD grains due to interlayer van der Waals interaction, but the TMD grain size is limited, typically less than 1 μ m,^{17–20} and the use of other 2D materials complicates the process and increases the costs.

Among the various existing TMDs, WS₂ possesses promising properties, such as strong photo- and electroluminescence, the spin-valley coupling phenomenon, non-toxicity of the source materials, and a lower effective mass of electrons compared with MoS₂ and MoSe₂.²¹⁻ 24 These properties make monolayer WS₂ a promising candidate for electronics, photonics, and valleytronics. In most of the CVD studies, WS2 is grown from tungsten oxide (WO3) and sulfur (S) precursors in the flow of an inert gas, mainly Ar.^{8,14,25} Recently, the addition of hydrogen has been found to improve the quality of TMDs.^{26–29} The added H₂ gas, usually at concentrations below 10%, can produce defined grain edges,²⁶ increase the photoluminescence intensity,²⁷ promote reduction of a metal oxide source,²⁸ and suppress the multilayer growth.²⁹ In this chapter, high concentrations of H_2 gas (> 40%) during the CVD growth in shown to assist the alignment of triangular WS₂ grains along the c-plane sapphire substrate. Compared with the case of the randomly oriented grains obtained for pure Ar, the PL and Raman signals from the aligned WS₂ grains were strongly suppressed. This suggests enhancement of the WS₂-sapphire interaction that ultimately assists the WS₂ alignment. DFT calculations were also performed to explain the enhanced WS_2 -sapphire interaction induced by the H_2 supply. STEM observations revealed the presence of unique and defined dislocations between connected aligned grains. On the contrary, randomly oriented WS₂ grains have various distorted GBs which are consisting of arrays of 8-, 7-, and alternating 7/5-membered rings. This study also indicates that the merged aligned WS₂ grains show higher chemical stability and improved transport properties compared with the randomly oriented WS₂ grains. The results of this

work offers a new tactic for the large-scale growth of highly crystalline TMD sheets by minimizing the effects of GBs, enhancing the potential applications of 2D materials.

3.2. Experimental method

3.2.1. CVD growth of WS₂

WS₂ was synthesized using WO₃ (99.998%) and sulfur powder (99.999%) on c-plane sapphire substrate using a tubular furnace with a heating belt. The CVD setup is illustrated in Figure 3.1. The inner diameter of the quartz tube is 26 mm. Pure Ar gas (99.999%) and mixture with H₂ gas (99.99999%) were flown during the CVD process, while keeping a total flow rate of 100 ccm. The temperatures of the WO₃ precursor (with the sapphire substrate on top) and sulfur were set at 825 °C and 210 °C, respectively. The growth time was 30 minutes. When H₂ gas is used in addition to Ar, caution is required not to inhale H₂S gas by venting the out-line gas after neutralization.



Figure 3.1. CVD setup used for the growth of monolayer WS_2 . A sapphire substrate was placed on top of the WO_3 boat.

3.2.2. Characterization methods

Scanning electron microscope (SEM) images were measured with a HITACHI S-4800 with 1 kV acceleration voltage. Atomic force microscope (AFM) images were measured with a Bruker Nanoscope V. PL and Raman spectra were measured with a Tokyo Instruments, Nanofinder 30 using a 473 nm laser beam with low power (100 μ W and 600 μ W for PL and Raman, respectively). Annular dark-field STEM (ADF-STEM) images were measured by using JEOL 2100F equipped with double delta correctors operating at the acceleration voltage of 60 kV. The probe current was about 25 pA. The convergence angle and the inner acquisition angle were 35 mrad and 79 mrad. The electron energy-loss spectroscopy (EELS) spectra were taken by using a Gatan low-voltage quantum spectrometer.

3.2.3. DFT calculations

Theoretical calculations were performed using DFT with the generalized gradient approximation (GGA-PBE)³⁰ and van der Waals (vdW) corrections (vdW-DF2-C09)^{31,32} implemented in the STATE package.³³ Vanderbilt ultrasoft pseudopotentials were used to describe electron-ion interactions.³⁴ Valence wave function and deficit charge density were expanded in terms of plane-wave basis with cutoff energy of 25 and 225 Ry, respectively. Surfaces of Al₂O₃ were simulated by a repeated slab model of α -Al₂O₃ thin films with 8 atomic layers. Al- and H-terminated Al₂O₃ surfaces were used to model H₂-WS₂ and Ar-WS₂ growth, respectively. To calculate the adsorption energy of a WS₂ grain on the Al₂O₃ slab, the 4 × 4 lateral cell was adopted to exclude the interaction with the periodic images of WS₂ grain. The triangular WS₂ grain with lateral length of 1.2 nm was constructed using 15 W and 36 S atoms. To avoid an unphysical dipole interaction with periodic images normal to the α -Al₂O₃ slab arising from the polar (111)/($\overline{111}$) surfaces, the effective screening medium (ESM) method

was adopted.³⁵ The geometric structures were optimized until the force acting on atoms was less than 0.01 Ry/Å under the fixed lattice parameter.

3.2.4. Device fabrication and measurement

SiO₂(90 nm)/n⁺-Si substrate was cleaned with acetone and isopropyl alcohol (IPA) by sonication for 10 minutes each. WS₂ flakes were transferred onto prepared SiO₂/Si substrates using the polystyrene-assisted transfer technique.³⁶ Transferred WS₂ was spin coated by polymethyl methacrylate (PMMA) and patterned using electron beam (EB) lithography, followed by thermal evaporation of Ti/Au electrodes in vacuum and lift-off. Device measurement was conducted at room temperature in vacuum at 10⁻³ Pa with a bias voltage of 1 V.

3.3. Results and discussion

3.3.1. Hydrogen-assisted epitaxial growth of WS₂

Figure 3.2a,b shows SEM images of WS₂ grains synthesized by ambient-pressure CVD using H_2 -Ar mixed gas (H_2 : 50%) and pure Ar gas, which are denoted hereafter as H_2 -WS₂ and Ar-WS₂, respectively. For the WS₂ synthesis, WO₃ and sulfur powders were used as precursor with the c-plane sapphire substrate placed on top of the WO₃ boat without any nucleation seeds. Most of the CVD parameters, such as total gas flow rate, growth time, and substrate temperature were set to the same values for both H_2 -WS₂ and Ar-WS₂, except for the gas composition and the amount of sulfur powder (a larger amount of sulfur was necessary for the H_2 -WS₂ growth, as the formation of H_2 S leads to higher consumption of sulfur).


Figure 3.2. SEM images and angle distributions of the as-grown H_2 -WS₂ (a,c) and Ar-WS₂ (b,d) grains. Arrows in (a) and (b) show the crystal orientations of the c-sapphire substrate. (e) Atomic structure of WS₂ grains oriented on c-plane sapphire observed for H_2 -WS₂ samples.

In the case of Ar-WS₂, triangular grains are randomly oriented on the sapphire substrate (Figure 3.2b), as reported in the previous literatures.^{5,9,26,37} In contrast, most of the H₂-WS₂ grains are oriented along only two directions (Figure 3.2a). One side of these oriented WS₂ grains is aligned parallel to the [1 $\overline{1}00$] direction of the underlying sapphire substrate, as seen in Figure 3.2a. This result suggests an epitaxial growth of WS₂ grains assisted by the H₂. Figure 3.2c,d shows the angle distribution of WS₂ grains. In the presence of H₂ gas, 94% of the WS₂ grains are aligned to the [1 $\overline{1}00$] direction within ±5° dispersion, signifying the orientation of WS₂ is determined by the sapphire surface. The CVD results obtained with other H₂ concentrations are shown in Figure 3.3. Relatively high H₂ concentrations over 40% were found to improve the WS₂ alignment (see Figure 3.4). Figure 3.2e depicts the



Figure 3.3. SEM images of WS₂ grains grown under different H_2 concentrations. Total flow rate was fixed to be 100 sccm. All the samples are set in the same direction designated in H_2 0% image.

orientation of WS₂ grains with respect to the underlying sapphire lattice. Although the lattice mismatch between WS₂ (0.315 nm) and c-plane sapphire (0.476 nm) is large (34.4%),^{38,39} van der Waals epitaxy allows the alignment of the H₂-WS₂ grains on sapphire. The aligned growth can lead to perfect connection of the neighboring WS₂ grains having the same orientation without any GBs, as illustrated in the left part of Figure 3.2e, which will be discussed later.

AFM images of H₂-WS₂ and Ar-WS₂ grains are shown in Figures 3.5a,b, respectively. The height of the H₂-WS₂ grains was ~0.8 nm, suggesting the formation of monolayer WS₂.²⁶ It can be seen that the H₂-WS₂ grains are almost free from multilayer areas at their centers (Figure 3.5a). In contrast, the Ar-WS₂ grains contain thick overgrown multilayers at the center of monolayer grains (bright triangles in Figure 3.5b). The height of the multilayer grains is



Figure 3.4. Histograms of grain orientation distribution corresponding to the samples shown in Figure 3.3. The $[1\overline{1}00]$ direction of the sapphire substrate is set as 0°. Grains of WS₂ show the preferential alignment under the hydrogen concentration higher than 40%.

generally higher than 100 nm. These multilayer grains are widely observed in samples of different TMDs grown by CVD, and it becomes more severe when the growth time is extended to fully cover the sapphire surface, because the coverage of the overgrown multilayers increases with the growth time.^{3–7} It is speculated that H_2 gas introduced during the WS₂ growth process etches multilayer nuclei when they appear on the monolayer WS₂, thus suppressing the



Figure 3.5. AFM and ADF-STEM images the H₂-WS₂ (a,c) and Ar-WS₂ (b,d) grains.

formation of multilayer WS₂ growth. This is consistent with the recent literature reporting that the addition of H_2 in low concentrations (5%) suppresses the formation of multilayer WS₂.²⁹ The H₂-assisted etching, at the same time, reduces the growth rate of H₂-WS₂, resulting in smaller grains compared with the Ar-WS₂ grains (see Figure 3.2a,b).

To evaluate the crystallinity of the WS₂ grains, ADF-STEM was measured for both H₂-WS₂ and Ar-WS₂ samples, as shown in Figures 3.5c,d. Bright and dark spots in the images correspond to W and S atoms, respectively. The Ar-WS₂ grain contains several point defects (Figure 3.5d), which include vacancies and impurities. Metal impurities are assigned to Mo from the EELS analysis (see Figure 3.6). Although the origin of these metal impurities is not clear, contamination due to impurities in the feedstock and/or from sources used for MoS₂ growth in our laboratory (even though different setup is used for each TMD) may occur, because TMD monolayers are highly susceptible to the unintentional substitutional doping.⁴⁰ However, as seen in Figure 3.6a, the number of defects and impurities was much smaller for



Figure 3.6. ADF-STEM images of H_2 -WS₂ (a) and Ar-WS₂ (b) grains. The Ar-WS₂ grain has a higher amount of impurities and vacancies as compared with H_2 -WS₂. (c-f) ADF-STEM images and EELS data of vacancy (c,d) and impurity (e,f). The right panel of (f) shows the ADF intensity, suggesting the presence of a Mo atom in the lattice.

the H₂-WS₂ grains. There are two possible reasons for this lack of defects when using H₂. First, H₂ can reduce precursors more effectively, converting WO₃ and S to WO_{3-x} and H₂S, respectively, which helps to improve the crystallinity by reducing structural defects caused by oxygen included in WO₃ source.²⁷ This was confirmed by the X-ray photoelectron spectroscopy (XPS) measurements, as shown in Figure 3.7. The H₂-WS₂ sample shows the preferential formation of WS₂, while the Ar-WS₂ shows the presence of WO₃ and/or WO_xS_y species on the sapphire surface in addition to WS₂.



Figure 3.7. XPS spectra of sapphire surfaces after WS_2 growth with and without H_2 gas. The surface coverage of WS_2 is around 10%.

defects in the H_2 -WS₂ grains is that H_2 may be able to heal the defects formed within the WS₂ lattices. Since defects are more energetically unstable, they can be easily eliminated by H_2 during the growth, which allows to reconstruct the WS₂ crystalline lattice. This phenomenon is related to the observed suppression of overgrown multilayer WS₂ by removing the multilayer nuclei from monolayer grains. As shown earlier, the use of H_2 during the CVD assists the epitaxial growth of aligned WS_2 grains with almost no structural defects. Therefore, physical properties of TMD monolayers grown with the aid of H_2 are expected to be improved, as will be discussed later.

It has been reported that MoS_2 grains can be aligned on sapphire in pure Ar flow (i.e. without H_2 gas).^{16,41} We have also observed the grain alignment of MoS_2 on sapphire in the absence of H_2 .⁴² In contrast, for WS₂ grains, I have not observed the grain alignment when no H_2 is supplied regardless of the substrate position (either above the WO₃ precursor or downstream of it) and the growth condition (different temperatures and Ar flow rates). Also, there has been no previous reports of the WS₂ alignment as far as I know. Thus, the aligned growth of WS₂ is much more difficult to achieve than that of MoS_2 , which may be related to the surface condition of the sapphire substrate. As the H_2 gas can clean the sapphire surface and gives the WS₂ alignment, it is expected that H_2 -assisted CVD method can be applied to other TMDs, such as WSe₂.

3.3.2. Optical properties of WS₂ grains

To understand the optical properties of the WS₂ grains, PL spectra from WS₂ grains grown with and without H₂ gas during the CVD process were measured. Figure 3.8a shows the PL intensity map of an as-grown Ar-WS₂ grain on sapphire substrate. While the PL intensity is weak at the center due to the presence of multilayer grain,^{10,21,43} most of the area shows strong emission. On the other hand, the PL from the as-grown H₂-WS₂ was strongly suppressed at the central parts of the grains, although the edges show clear PL emission (Figures 3.8b and Figure 3.9). There are two possible reasons for the observed reduction of the PL intensity of the H₂-WS₂ grains; structural defects in the grains^{44,45} or a strong coupling (or charge transfer)) with the underlying sapphire.^{17,46,47} As shown in Figures 3.8c and Figure 3.9, the PL became



Figure 3.8. (a) PL intensity maps of as-grown Ar-WS₂ (a) and H_2 -WS₂ (b) on sapphire. (c) PL spectra collected from the H_2 -WS₂ grain shown in (b) before and after the transfer to SiO₂/Si substrate. The averaged PL spectra collected from the square indicated in (b) of the identical WS₂ grain are compared.



Figure 3.9. PL analysis of (a-c) as-grown and (d-f) transferred H_2 -WS₂. (a,d) PL intensity. (b,e) PL peak position. Normalized PL spectra (c,f) were measured from the marked points in (a). (c) and (f) show slight red-shift of the PL peak with increasing intensity from the center to the edge of the H_2 -WS₂ grains, suggesting doping at the grain edges by gaseous molecules in ambient air.

much stronger after transferring the H_2 -WS₂ grains onto a SiO₂/Si substrate. If the PL quenching in the as-grown sample (Figure 3.8b) would be related to structural defects, then the PL should remain weak after the transfer. However, given that the transferred H_2 -WS₂ grains

show PL from the whole grains, the PL quenching in the as-grown H₂-WS₂ grains seems to originate from the interaction between the WS₂ and the sapphire surface. It is noted that the light reflection by the underlying SiO₂ substrate also contributes to the enhanced PL from the transferred H₂-WS₂ grains. The PL intensity from the Ar-WS₂ grains also increased after the transfer. However, the suppression of the PL mainly at the central part of the H₂-WS₂ grains is quite different from the Ar-WS₂, signifying the substantial influence of sapphire to the H₂-WS₂ grains.

It is speculated that the emission from the edges of H_2 -WS₂ grains reflects the reduction of the WS₂-sapphire interaction induced by the intercalation of gaseous molecules in ambient air and/or the successive p-type doping from the molecules (see Figure 3.9), which may occur gradually in air.⁴⁸ After the transfer, the PL peak shifted from 2.00 eV to 2.03 eV (see Figure 3.8c). It is likely that tensile strain, arising during the cooling down process due to the different thermal expansion coefficients of WS₂ and sapphire, is relaxed during the transfer process.⁴⁹

I also investigated the influence of gas environment by switching from pure Ar to a H₂-Ar gas mixture during the WS₂ growth period. The result is shown in Figure 3.10. Two different kinds of WS₂ grains were observed on the sapphire surface after the CVD; large misaligned grains nucleated during the Ar period at the beginning, and small aligned grains grown during the H₂-Ar period. The latter showed a reduced PL, while the former gave the intense PL emission. These results confirm the strong influence of H₂ gas on the characteristics of WS₂ grains.

Figure 3.11a shows the Raman spectra of the as-grown Ar-WS₂ (black) and H₂-WS₂ (red) grains measured using a 473 nm laser excitation. Both samples showed clear E^{1}_{2g} and A_{1g} peaks with a peak separation of 62 cm⁻¹, evidencing the formation of monolayer WS₂.⁴⁹ The Raman intensity of the H₂-grown sample was found to be weaker, less than half of that of the



Figure 3.10. CVD growth of WS₂ by switching pure Ar to H₂-Ar mixed gas during the CVD growth. (a) CVD profile and the corresponding carrier gas composition. (b) Optical micrograph of the WS₂ grains on sapphire. (c) The same image with (b) highlight of the growth periods. (d) PL intensity map taken at the dashed square shown in (b). It is clearly seen the WS₂ grains that grow during flowing H₂-Ar are aligned and show weaker PL, while the grain formed during Ar flow is misoriented and exhibits stronger PL.



Figure 3.11. (a) Raman spectra of the as-grown samples. (b) Raman spectra of the identical H_2 -WS₂ grain taken before and after the transfer.

Ar-grown sample measured under the same condition. This lowering of the Raman intensity seen for the H_2 -WS₂ also can be attributed to the effective WS₂-sapphire coupling, considering the fewer structural defects and higher purity compared to Ar-WS₂ grains (see Figure 3.5c,d). The enhancement of the Raman intensity after the transfer process, shown in Figure 3.11b, also supports this conclusion. While the suppression of the PL from the H₂-WS₂ is ascribed to the strong orbital interaction between WS₂ and sapphire, the reduced Raman intensity can be explained by modulation of phonon modes of the H₂-WS₂ due to strong interaction between WS₂ and Al-terminated sapphire surface. This is supported by the DFT calculations, as mentioned later. In addition, the release of strain can be seen from the Raman shift after the transfer.⁴⁹

3.3.3. Analyses of sapphire surfaces

Considering the previous literature that reports the desorption of surface oxygen from sapphire surface by heating in hydrogen at 823 °C,⁵⁰ it is highly likely that hydrogen supplied during the CVD removes the surface oxygen at elevated temperature. This is expected to result in oxygen-deficient, Al-rich sapphire surface. Thus, the chemical states of sapphire surfaces were measured by XPS (see Figure 3.7c,d). In the H₂-assisted CVD, the main Al_{2p} peak associated with α -Al₂O₃ (74.2 eV) and a small peak are observed at lower binding energy (peak A (72.7 eV) in Figure 3.7). The low binding energy indicates that some of Al is less positively charged than that of α -Al₂O₃, being consistent with the expected oxygen-deficient surface state. On the other hand, the sapphire after the CVD with pure Ar showed an additional Al_{2p} peak at a higher binding energy (peak B in Figure 3.7c). This can be assigned to oxygen-rich sapphire, presumably suggesting the presence of OH-terminated sapphire



Figure 3.12. Water contact angle measurements for the pristine sapphire and the sapphire after the CVD with H₂-Ar and pure Ar flows.

surface. The oxygen concentration ([O]/([A1]+[O])) of the sapphire surface submitted to the H₂-assisted CVD (47.1%) was lower than that of sapphire submitted to standard Ar CVD (48.1%). As the sapphire substrates were exposed to air during the transport to the XPS chamber after the CVD process, which inevitably causes surface adsorption of oxygen and water molecules, the difference of the measured oxygen concentrations may have become smaller. The deviation from the stoichiometry of α -Al₂O₃ (60%) can be also due to the heat-induced desorption of surface oxygen.⁵¹

For macroscopic inspection of the sapphire surface, the water contact angle was measured. As shown in Figure 3.12, the sapphire surface after the H_2 -WS₂ growth showed a much higher contact angle than that after the Ar-WS₂ growth. This is consistent with the XPS results; the sapphire reacted with H_2 has less surface oxygen, making the sapphire surface more hydrophobic. It is considered that the Al-rich, active sapphire surface, formed by H_2 , induces the strong interaction with the growing WS₂ grains, guiding their lattice orientation to align along the sapphire's surface potential.

I also investigated the CVD growth in pure Ar flow using a sapphire substrate which was annealed in H_2 flow (100%) prior to the CVD. Figure 3.13 shows the SEM image and PL spectrum. Although the WS₂ grain shape is not highly defined, a partial alignment was observed, with the WS₂ grains elongated along a crystal direction of the sapphire. Also, the PL was weaker than that of grains grown on non-annealed sapphire substrate. Thus, the high temperature treatment in H_2 gas influences the chemical structure of the sapphire surface, highly likely assisting the WS₂ alignment. It was reported that the step edges of c-plane sapphire substrates can guide the growth of aligned TMDs.^{52,53} However, although step/terrace structures on the sapphire surfaces after the CVD with H₂-Ar gas (Figure 3.14), the WS₂ orientation did not follow the step edge direction. This proves that the WS₂ alignment in our H₂-assisted CVD is based on the atomic lattice of c-plane sapphire surface.



Figure 3.13. Effect of the H_2 pre-annealing of sapphire prior to the CVD growth with pure Ar gas. (a) Experimental procedure. SEM images of the sapphire surface pre-annealed in H_2 (b,c) and that without pre-annealing (d). (d) corresponds to the standard Ar-WS₂ grains. (e) PL spectra of WS₂ grains.

Energy (eV)



Figure 3.14. AFM images of the sapphire surface after the H_2 -assisted CVD growth. The WS_2 grains are oriented in two directions, but not aligned along the step/terrace direction.

For further understanding of the influence of H₂-induced WS₂ alignment, DFT calculations were performed using the slab models shown in Figure 3.15. Here, triangular WS₂ grains with 1.2 nm lateral size are adsorbed on α -Al₂O₃ slabs whose surfaces are terminated with Al



* Adsorption energy per one triangular WS₂ grain consisting of 51 atoms

Figure 3.15. DFT calculation results of triangular WS_2 grains adsorbed on Al-terminated (a,b) and O-terminated (c,d) sapphire surfaces. (a,c) Atomic structures after geometry optimization. Other structural images can be found in Figure 3.16. (b,d) Spatial distribution of valence electrons. Yellow: S, gray: W, blue: Al, red: O atoms. The arrow in (b) indicates the distorted S $2p_z$ orbital due to strong interaction with the atomic orbitals of Al atoms of sapphire surface. (e) Comparison of WS₂-sapphire distances and adsorption energies.



Figure 3.16. Atomic structures of WS_2 grains adsorbed on Al- (a) and O-terminated (b) sapphire. Left and right images show the side and top view images, respectively. These structures are identical to those shown in Figure 3.15. It is clearly seen that the WS_2 -sapphire distance after geometry optimization is shorter for the Al-terminated surface (a) than the O-terminated one, indicating the stronger interaction between WS_2 and the Al-terminated sapphire.

atoms (Figure 3.15a) and O atoms (Figure 3.15c). As discussed above, the Al- and Oterminated surfaces correspond to the H₂-WS₂ and Ar-WS₂ cases, respectively. It was found that after geometry optimization, the WS₂-sapphire distance is shorter for the Al-terminated surface (see Figure 3.15a, 0.15 nm) than for the O-terminated one (Figure 3.15c, 0.29 nm). In addition, in the valence electron distribution shown in Figure 3.15b, the S 2p orbital is extended to the sapphire surface, suggesting the orbital hybridization of WS₂ and the Al-terminated sapphire surface. Reflecting this strong coupling, the measured adsorption energy was much higher for the Al-terminated than the O-terminated surfaces (Figure 3.15e). Thus, the PL quenching and the reduced Raman intensity observed in the H₂-WS₂ can be ascribed to this strong WS₂-sapphire coupling.

3.3.4. Observation of WS₂ grain boundaries

The hydrogen-assisted epitaxy found in this work motivated us to measure the atomic structures of the interfaces between merged WS₂ grains, as misorientation angles greatly influence the GB structures and transport properties.¹⁵ In this study, we analyzed the interface between aligned WS₂ grains merged with a low misorientation angle realized by our H₂-assisted CVD growth technique, in addition to the GBs of rotated grains with large misorientation angles (Ar-WS₂). Figure 3.17a shows the low magnification STEM image of a GB with a low misorientation angle of ~4°. In the SEM and AFM images shown in Figures 3.2a and 3.5a, the triangular WS₂ grains look highly aligned, but there may be slight misalignment between neighboring grains due to the tapered or round triangular structures as well as thermal fluctuation during the CVD growth.

Interestingly, as shown in Figure 3.17a, there was no linear defect (or GB) along the interface between the two merged H_2 -WS₂ grains, which is widely observed in the merged TMD



Figure 3.17. (a-d) ADF-STEM images of GBs of H₂-WS₂. (a-c) GB structures with misorientation angle of ~4°. (d) Three-fold merging GBs with 2° and -4° of mismatch angles. The marked lines indicate dislocations. (e-i) ADF-STEM images of GBs of large angle misoriented Ar-WS₂. Different merging angles of (e-g) 19°, (h) 24° and (i) 10° resulted in different GB structures: (e-g) 7-5-membered rings, (h) 7-membered rings and (i) 8-membered rings.

grains.^{12–15} Instead, one-dimensional (1D) dislocation with small point defects (dislocation cores) were sparsely observed, as can be seen from the magnified images (Figure 3.17b,c). It is surprising that the crystalline structure is almost preserved even at the merged interface. Here, only few dislocations are expected to be enough to compensate the lattice discontinuity due to the very small misorientation angle and offset of the neighboring lattices. This allows the merging grains to form structurally continuous interface without forming severe structural defects along the boundary. Therefore, the electronic properties across the merged interface are not likely to be affected by such dislocations, resulting in nice atomic stitching. It is noted that such stitching has not been reported so far for merged TMD grains.

Figure 3.17d shows the GBs formed by three merging grains with small misorientation angles; the grains B and C are rotated by 2° and -4° with respect to the grain A, respectively. In this case, two parallel dislocation lines existing in one grain are connected to one dislocation in another grain at a dislocation core. Accordingly, the dislocation core is slightly larger than that seen in Figure 3.17b,c. This result also demonstrate that the small misorientation angles can give very smooth atomic-scale connection by simply forming a few excess atomic lines to compensate lattice discontinuity and release the possible strain.

Figure 3.17e displays the GB of two merged grains with misorientation angle of 19°, which was grown in pure Ar flow. This large misorientation angle resulted in a continuous defect line along the boundary. In this case, a series of connected 7-5 membered rings was observed along the GB, as shown in the magnified images of Figure 3.17f,g. Moreover, different types of defect structures were widely observed along other large-angle GBs. Figures 3.17h,i show the GBs formed for grains merged with angles of 24° and 10°, respectively. Series of 7-membered ring defects (Figure 3.17h) and 8-membered rings (Figure 3.17i) were observed along the respective GBs. The observation of these linearly arranged defects is consistent with the previous literatures on TMDs that report different types of GBs consisting of a number of irregular atomic rings along the GB.^{13–15} Therefore, the low misorientation angle realized by the CVD with a high H₂ concentration produces unique GB structures, which would give much less influence on the physical and chemical properties.

It is difficult to differentiate the role of hydrogen from the effect of low angle misorientationinduced connection observed here, because the WS₂ grains cannot be aligned in the absence of H_2 during the growth. Given that large misorientation angles lead to defects forming along the GBs, the misorientation angle is considered to be a key factor in forming consistent stitching between neighboring WS₂ grains. However, as discussed in Figure 3.2e, the H_2 supplied during CVD growth may also contribute to effective stitching by minimizing (or eliminating) formation of lattice defects.

3.3.5. Influence of GBs on chemical and physical properties of WS₂

There are several methods to visualize GBs in TMD monolayers, such as oxidation^{26,54,55} and Joule heating.⁵⁶ Most of them utilize the relatively high chemical reactivity of GBs. In order to investigate the chemical stability of GBs, I oxidized the merged WS₂ grains in air at 300 °C for two hours. Figures 3.18a,b show the SEM images of H₂- and Ar-WS₂ grains, respectively, taken after the oxidation. As previously reported,²⁶ large-angle misoriented Ar-WS₂ grains showed clear visible defects along the GBs after this mild oxidation (Figure 3.18b). On the other hand, the H₂-WS₂ sample (Figure 3.18a) showed difference in oxidation behavior. The boundary between WS₂ grains oriented in the opposite directions (marked with red and blue triangles) exhibited a clear oxidation line similar to those observed for the misaligned grains (Figure 3.18b). This is because a mirror twin boundary is present between these 180°-rotated grains.¹¹ On the contrary, no visible line was observed for the grains connected with the same orientation (two red triangles). This demonstrates that the low angle misorientation gives consistent stitching that is chemically more stable than other rotated GBs, being consistent with our ADF-STEM observations (see Figure 3.17a-c).

The presence of GBs is also known to deteriorate the carrier transport properties of 2D materials, because they act as a scattering site for carriers while also introduce defect states in



Figure 3.18. SEM images of (a) H_2 -WS₂ and (b) Ar-WS₂ grains after oxidation. Red and blue triangles in (a) indicate upward- and downward-oriented aligned grains, respectively. Inset in (b) shows the magnified image of dotted square.



Figure 3.19. Transfer curves of merged H_2 -WS₂ (a) and Ar-WS₂ grains (b). Insets show optical micrographs of the measured devices.

the middle of the energy band that trap carriers.^{11,13,15} Four-terminal devices were fabricated on merged WS₂ grains with Ti/Au electrodes defined by EB lithography. Insets in Figure 3.19 show optical micrographs of the devices made with H₂-WS₂ (a) and Ar-WS₂ (b) grains with merged angles of 1-2° and ~11°, respectively. In the case of the Ar-WS₂ (Figure 3.19b), intergrain conductance was clearly lower than the intra-grain conductance. These merged grains may have a GB similar to that shown in Figure 3.17i, as the misorientation angle (~11°) is very close. Thus, a linear chain of the irregular and distorted atomic rings formed along the GB should be responsible for the observed low conductance. In contrast, the H₂-WS₂ grains show almost no conductance degradation across the interface, as shown in Figure 3.19a. This result suggests that the sparsely distributed dislocations (see Figure 3.17a-c) do not significantly affect the carrier transport, which is promising for realization of high performance electronic devices.

3.4. Conclusions

The epitaxial growth of aligned WS₂ on c-plane sapphire substrates is demonstrated using high concentration of hydrogen during the CVD growth. The presence of hydrogen can enhance WS₂-sapphire interaction, likely by exposing an Al-rich surface on the sapphire substrate. This change of the chemical state of the sapphire surface induced by hydrogen was supported by XPS and water contact angle measurements. The DFT calculations suggest the Al-rich surface enhances the coupling between a WS₂ grain and sapphire, which can potentially affect the grain alignment and PL intensity. The H₂-induced alignment is critical for obtaining low misorientation angles between merged WS₂ grains. Inspection of the merging areas by ADF-STEM revealed the unique and smooth reconstructed interface structure with sparsely located 1D dislocations, which can release the small strain between adjacent grains. In stark contrast, the highly misaligned WS₂ grains grown in pure Ar showed linear arrays of distorted atomic rings along the boundaries between merged grains. These boundary structures were found to influence the chemical stability of the WS₂ and the carrier transport, implying that realizing low angle connection by the epitaxial growth of WS2 is essential to obtain high quality This study is expected to shed light on the formation of well-stitched grains and at the sheets. same time offers an important guiding principle to obtain high-quality TMD films that can be applied to future electronic and optoelectronic devices.

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Chapter 4

Influence of van der Waals Interaction on Growth and Physical Properties of Epitaxially Aligned WS₂

Abstract

In the previous chapter, the epitaxial growth of high quality WS₂ on c-plane sapphire substrate was demonstrated. Having similar hexagonal crystal structure with many TMDs, c-plane sapphire is commonly utilized as a growth substrate in CVD synthesis. Due to the similarity of crystal structure, c-sapphire substrate is expected to influence the growth of TMDs while few studies have been reported about this. Thus, in this chapter, I demonstrate that higher strain is induced in the epitaxially grown WS₂ grains via van der Waals interaction with sapphire as compared with misaligned WS₂ grains. In addition, this strain was found to enhance overlayer deposition on monolayer WS₂, while multilayer growth was not observed in non-epitaxial WS₂. PL of the epitaxially grown WS₂ grains was reduced reflecting effective van der Waals interaction with sapphire. Moreover, low-temperature PL measurement revealed strong influence of the c-plane sapphire surface on the optical property of WS₂. DFT calculation supports that the aligned WS₂ grains are more strongly bound to the sapphire surface, as compared with misaligned WS₂.

4.1. Introduction

The CVD growth of monolayer TMDs has been widely studied using various kinds of precursors, such as metal oxide and metalorganic gas, and different carrier gases, such as pure Ar and the mixed gas with H₂ or O₂, in order to increase the grain size and improve the film uniformity.^{1–8} Also, different types of substrates have been studied, including insulators (SiO₂, sapphire) and metal substrates (Au and Ni foils).^{2,9–12} Among them, c-plane (0001) sapphire which has a hexagonal surface crystal structure similar to those of WS₂ and MoS₂ is one of the most suitable template. Epitaxial growth of WS₂ was achieved on the sapphire substrate by flowing hydrogen during the CVD growth.⁸ This epitaxial growth on sapphire is expected to give highly crystallized TMD films as compared with polycrystalline films consisting of randomly oriented grains that are widely formed on the SiO₂ substrate, because stitching of the merged grains can occur for the aligned TMD grains.^{13–19} However, it is not clearly understood how the sapphire surface affects physical properties of the aligned TMDs.²⁰

Therefore, in this work, I have investigated physical properties of the epitaxially aligned WS_2 grains by comparing with randomly oriented grains which are formed on the same c-plane sapphire substrate. It was found that the PL from aligned WS_2 grains is weaker than that of misaligned grains. This result is interpreted by substrate-induced higher strain in the aligned WS_2 due to effective van der Waals interaction between WS_2 and the sapphire surface. Selective overlayer deposition of multilayer WS_2 was also observed on the aligned grains, reflecting the induced strain. DFT calculation was performed to investigate the interaction between WS_2 grains and sapphire. Low-temperature PL revealed significant difference in the optical properties after detaching from the sapphire substrate. Carrier transport measurements were also performed to further understand the crystallinity of aligned and misaligned WS_2 grains.

4.2. Experimental method

4.2.1. CVD growth of WS₂

WS₂ was synthesized from WO₃ (99.998%, Alfa Aesar) and sulfur (99.999%, Mitsuwa Chemicals) precursors by ambient pressure CVD method. C-plane sapphire (Kyocera) with a low miscut angle (< 0.2°) was used as substrate which was cleaned by sonicating in acetone and isopropyl alcohol prior to the CVD. For the CVD growth, a three-zone tubular furnace with a quartz tube (outer diameter is 26 mm) and a heating belt were used, as illustrated in Figure 4.1. Different from Figure 3.1, an additional inner quartz tube was used for more efficient reaction between precursors on growth substrate. During the CVD reaction, 500 ccm of pure Ar gas (99.999%) was flowed as a carrier gas. The temperatures of sulfur, WO₃, and sapphire substrate were set at 160 °C, 1080 °C, and 970 °C, respectively. After 30 min reaction, the sample was cooled down naturally inside the furnace while flowing Ar gas.



Figure 4.1. CVD setup used for growth of WS_2 . A three-zone tubular furnace was used for heating WO_3 and sapphire heating with a ribbon heater for sulfur heating. Different from the set up in Figure 3.1, additional inner quartz tube is used for more efficient reaction between precursors on growth substrate.

4.2.2. Characterization methods

As-grown WS₂ was characterized by optical microscope (OM, Nikon Eclipse ME600) and AFM (Bruker Nanoscope V). To measure room-temperature PL and Raman spectra, a confocal Raman spectroscope (Tokyo Instrument Nanofinder30) with 473 nm laser excitation was used. For measuring PL spectra at low temperatures, Nanophoton Ramantouch was used with a Helium cryostat (Janis ST-500UC) using 532 nm laser excitation.

4.2.3. DFT calculations

Theoretical calculations were performed using DFT with the generalized gradient approximation $(GGA-PBE)^{21}$ and van der Waals corrections $(vdW-DF2-C09)^{22,23}$ implemented in the STATE package.²⁴ Vanderbilt ultrasoft pseudopotentials were used to describe electron-ion interactions.²⁵ Valence wave function and deficit charge density were expanded in terms of plane-wave basis with cutoff energy of 25 and 225 Ry, respectively. Surfaces of Al₂O₃ were simulated by a repeated slab model of α -Al₂O₃ thin films with 8 atomic layers with Al-terminated Al₂O₃ surfaces. To calculate the adsorption energy of a WS₂ grain on the Al₂O₃ slab, the 4 × 4 lateral cell was adopted to exclude the interaction with the periodic images of WS₂ grain. The triangular WS₂ grain with a lateral length of 1.3 nm was constructed with 15 W and 36 S atoms. The geometric structures were optimized until the force acting on atoms was less than 0.01 Ry/Å under the fixed lattice parameter.

4.2.4. Device fabrication and measurement

For fabricating WS₂-FETs, WS₂ grains grown on sapphire was transferred onto SiO₂/Si substrate using polystyrene and KOH.²⁶ Then, electrode patterns were defined by a photolithography process, followed by deposition of Ni (5 nm)/Au (25 nm) electrodes using an EB evaporator (Kennix KB-750) and lift-off process. Prior to the measurement, the device was annealed in Ar at 200 °C for 2h to remove polymer residue. The devices were measured with a semiconductor parameter analyzer (Keysight Technologies B1500A) in vacuum (~10⁻⁴ Pa) at room temperature.

4.3. Results and discussion

4.3.1. Aligned and misaligned WS2 grains on c-plane sapphire

Figure 4.2a,b shows optical micrographs of as-grown WS₂ on c-plane sapphire. It was found that there exist two types of triangular-shaped WS₂ grains together on the single sapphire substrate. One is aligned grains whose side edge is parallel to the [1 $\overline{100}$] axis of the underlying sapphire substrate (highlighted by red triangles in Figure 4.2a,b), indicating the epitaxially grown WS₂ grains. Another is randomly oriented (misaligned) grains highlighted by green triangles in Figure 4.2a,b. Interestingly, I found that overlayer deposition of multilayer WS₂ occurs preferentially on the aligned WS₂ grains; aligned grains showed many multilayer grains on their surface. On the other hand, the misaligned grains were almost free from multilayer deposition except for a small multilayer triangle only at the center of monolayer grain which acted as a nucleation site. Such difference is clearly seen in the corresponding AFM image



Figure 4.2. (a,b) Optical images of as-grown WS_2 grains on sapphire substrate with low (a) and high (b) magnifications. The crystalline axes of sapphire are shown by white arrows. (c,d) AFM height image and schematic of the WS_2 grains shown in (b,c). The overlayer deposition occurs only on the aligned grain (left grain), while the misaligned grain is free from multilayer (right grain).

(Figure 4.2c). This AFM image signifies that many multilayer grains formed on the aligned WS₂ are oriented in the same direction with the monolayer grain underneath. Figure 4.2d displays the atomic model of the aligned (left) and misaligned (right) WS₂ grains formed on sapphire where the former has multilayer grains on its surface. To confirm this result, I performed more than 20 CVD runs. As shown in Figure 4.3, in most of the CVD runs, the same tendency was observed; aligned WS₂ grains have multilayer grains on their surfaces.

It is speculated that van der Waals interaction between WS_2 and sapphire is stronger for the aligned WS_2 than the randomly oriented WS_2 grains.^{8,27,28} This strong interaction can induce large strain in the aligned WS_2 , because the mismatch of the



Figure 4.3. Optical micrographs of as-grown WS_2 grains on sapphire substrates. (a) The preferential overlayer deposition was observed on the aligned grains. (b) In a few cases, the sapphire surface was fully covered with WS_2 together with large multilayer grains.

	Growth parameters				Growth mode	
	Gas flow rate	WO ₃	s	sapphire	Epitaxially aligned grains	Preferential multilayer growth
Α	300 ccm	1080 °C	160 °C	970 °C	Х	X
В	500 ccm				0	0
С	700 ccm				0	0
D	500 ccm 1	1075 °C	160 °C	770 °C	Х	X
Е				870 °C	Х	X
F				970 °C	0	X
G				1070 °C	0	0

Table 4.1. CVD conditions used for sample $(A) \sim (G)$ shown in Figure 4.4 and the observed growth modes.



Figure 4.4. Optical images of as-grown WS_2 on c-plane sapphire synthesized using the growth conditions shown in Table 4.1.

lattice constants of WS₂ and sapphire is relatively large (4.76 Å for sapphire and 3.15 Å for WS₂).^{29–32} This strain could induce distortion in crystal structure of WS₂, resulting in local fluctuation of electronic potential of WS₂.³³ Such specific electrical and/or geometrical distortion is expected to assist the nucleation of upper WS₂ grains because TMDs preferentially nucleates from the disordered sites.^{20,34}

Influence of other CVD parameters on the growth behavior was also investigated. The results are summarized in Table 4.1 and Figure 4.4. As the gas flow rate was increased from 300 ccm to 700 ccm, the percentage of aligned grains increased, and the density of multilayer grains also increased. The higher flow rate can increase the supply rate of WO₃ and S feedstock, thus giving a larger amount of multilayer grains. At the same time, the increase of the S can improve the WS₂ orientation, as the higher S supply is known to give epitaxially aligned MoS₂.³⁵ I also investigated the effect of the temperature of sapphire substrate by changing it from 770 °C to 1070 °C (see Table 4.1). With the increase of the substrate temperature, the lateral growth was more promoted, reflecting the enhanced surface diffusion of the WS₂ intermediate species.

Generally, monolayer WS_2 exhibits strong PL reflecting its direct band gap. It is well acknowledged that the PL is sensitive to strain, defects, and carrier concentration. Figure 4.5a-c indicates the PL mapping images (PL intensity, full-width at half maximum (FWHM), and peak position) taken at the merged grains shown in Figure 4.2b,c. The PL spectra collected at the marked positions in Figure 4.5a are also shown in Figure 4.5d. Clear differences were observed between the aligned and the misaligned WS₂ grains. Compared with the misaligned WS₂ grain, the aligned grain showed lower PL intensity (Figure 4.5a), broader linewidth (Figure 4.5b), and slight red-shift (Figure 4.5c). As seen in the original PL spectra presented in Figure 4.5d, the PL intensity is clearly reduced in the aligned grain. As the monolayer areas was


Figure 4.5. PL mapping images of the intensity (a), FWHM (b), and peak position (c). (d) PL spectra from the marked points in (a). A1, A2 are from the aligned grain and M1, M2 are from the misaligned grain.

measured in the aligned grain, the reduced PL intensity is not due to the formation of multilayer grains. The PL quenching occurs via non-radiative relaxation process of excitons that is formed by the photo-excitation. There are two possibilities in the reduced PL seen in the aligned WS₂ grains, low crystallinity and strong interaction with substrate (in this case sapphire).^{36–39} As will be discussed later, the interaction with substrate is the main origin of the reduced PL. The different PL FWHM and peak positions (Figure 4.5b,c) can be also explained by the interaction with the sapphire surface, because the spectral weight of neutral excitons and charged trions modifies a PL spectrum of WS₂.^{17,33,40,41}

To avoid possible influence from multilayer growth, I synthesized monolayer, aligned WS₂ grains by lowering WO₃ temperature during the CVD to reduce the supply of



Figure 4.6. PL mapping images of the peak intensity (a), FWHM (b), and position (c) of monolayer WS₂ grains. (d) AFM height image. The inset shows a magnified image of the marked area. Raman E_{2g}^{1} mappings with intensity (e) and FWHM (f), and A_{1g} mappings with intensity (g) and FWHM (h). (i) PL, (j) E_{2g}^{1} , and (k) A_{1g} spectra taken from as-grown aligned and misaligned grains.

feedstock. The absence of multilayer was confirmed by AFM and Raman measurements (Figure 4.6). Figure 4.7 shows the detailed analysis of the aligned and misaligned, monolayer WS₂ grains. Here, a misaligned WS₂ grain rotated by ~45° with respect to [1100] axis of sapphire was chosen as a representative of misaligned grains in addition to the aligned grain (Figure 4.7a). The PL peak position and linewidth of each WS₂ grain measured at more than 30 points are compared in Figure 4.7b. It was confirmed that the PL linewidth is broader for the aligned grains (58.1±1.4 meV, red plots) than the misaligned ones (54.5±1.1 meV, green plots). The



Figure 4.7. (a) Atomic models of aligned (0°) and misaligned WS_2 (~45°) grains on c-plane sapphire. (b) Distributions of PL FWHM and peak position of as-grown WS_2 grains. (c) Distribution of the PL intensity along the line shown in the inset. The x-axis is the normalized distance from the nucleation center, as illustrate in the inset.

red shift of the peak position was also seen from the misaligned grains $(2.013\pm0.002 \text{ eV})$ to the aligned grains $(2.009\pm0.003 \text{ eV})$. According to the previous literatures, this shift (~4 meV) corresponds to 0.3~0.5% of tensile strain introduced in the aligned grain.^{42,43} Figure 4.7c indicates the PL intensity change measured from the center to the edge of WS₂ grains. Again, the PL intensity of the aligned grain is weaker than the misaligned grain. Also, the Raman intensity of the aligned WS₂ grain was weaker than misaligned grain (Figure 4.6j,k). The PL intensity gradually decreases from the center to the edge of the grains. In the high temperature atmosphere, the sapphire surface below the edge area of a WS₂ grain is exposed to the high temperature

Ar flow for longer time, while the sapphire below the central area of WS_2 grain is protected by WS_2 . Thus, it is considered that the sapphire surface under the edge area of the WS_2 grain interacts more strongly with WS_2 compared with central grain area, giving reduced PL intensity toward the edges.

4.3.2. DFT calculations

In order to understand the experimental observations, DFT calculation was performed using the model of a triangular WS₂ grain adsorbed on Al-terminated, c-plane sapphire surface. Figure 4.8a,b shows the calculation model of an epitaxially aligned WS₂ grain (consisting of 15 W atoms and 36 S atoms with 1.3 nm lateral size) on the sapphire surface. Two configurations, 0° (aligned) and 20° (misaligned), with respect to the [1100] axis of the sapphire surface were studied. Figure 4.8c compares the optimized WS₂-sapphire distance and the calculated adsorption energy. It was found that the aligned WS₂ grain has a relatively shorter distance (1.65 Å) than the misaligned grain



Figure 4.8. DFT calculation of a triangular WS_2 grain adsorbed on the c-plane sapphire. (a,b) Atomic models of the calculated model used for 0° orientation (epitaxially aligned). (a) Tile view and (b) top view. (c) Comparison of optimized WS_2 -sapphire distances and adsorption energies. The positive value in the adsorption energy indicates higher adsorption energy, *i.e.* stronger WS_2 -sapphire coupling.



Figure 4.9. PL mapping images of identical WS₂ grains before (a) and after (b) transfer on SiO₂/Si substrate. The difference in PL intensity observed between aligned- and misaligned-grains in asgrown WS₂ disappeared after the transfer process. The wavy patterns observed in the PL intensity of the transferred WS₂ (b) are from wrinkles induced during transfer process.

(1.78 Å). The adsorption energy is higher for the aligned grain, probably due to the enhanced WS₂-sapphire coupling with the consistent symmetry. These results indicate that the aligned WS₂ grains are more strongly bound to the sapphire surface compared with misaligned ones. It is speculated that such strong coupling with the sapphire surface seen in the aligned WS₂ grain results in the reduction of the PL intensity and the line broadening (see Figure 4.5a-d). It is interesting to note that the graphene grains aligned on Cu (111) also suffers from strong strain,^{44,45} suggesting that the grain alignment strongly affects the physical properties of 2D materials.

The strain in the aligned WS_2 grains was found to be released when detached from the sapphire surface. Figure 4.9 displays the PL mapping images of identical WS_2 grains taken before and after the transfer process. The weaker PL seen in the aligned WS_2 grains (Figure 4.9a) showed the similar intensity with misaligned grains after the transfer (Figure 4.9b). Therefore, a different degree of coupling with sapphire substrate is considered to modulate the optical property of as-grown WS_2 .

Also, to explain the preferential multilayer growth observed on the aligned grains, DFT calculation was performed for the strained bilayer WS_2 sheets. Van der Waals energy between the stacked bilayer WS_2 sheets were calculated while applying the tension to both the upper and lower layers. By inducing 4% of the tensile strain, the total energy of the stacked bilayers decreased by 2 meV per two pairs of S-W-S units. This indicates that the tensile strain enhances the van der Waals coupling between WS_2 layers, suggesting the ease of formation of multilayer grain on the strained WS_2 grain. When the tensile strain is introduced, a WS_2 layer can be slightly flattened so that the electron density within a layer is increased, thus giving stronger van der Waals coupling between the two strained WS_2 layers.⁴³ Although the actual system is more complicated than the present theoretical model, our present result qualitatively explains our experimental result.

4.3.3. PL spectra of MoS₂ grains on sapphire

For comparison, MoS₂ grains were also synthesized on c-plane sapphire using MoO₃ and S feedstocks using the similar setup shown in Figure 4.1.⁴⁶ The growth condition was modified to obtain both aligned and misaligned MoS₂ grains on the same sapphire substrate. As seen in Figure 4.10, the reduction of the PL intensity was also observed in the aligned MoS₂. Considering the similarity between WS₂ and MoS₂, such as a similar hexagonal crystal structure with comparable lattice constant, strong influence of the sapphire is deduced on the as-grown TMD materials.



Figure 4.10. OM image and PL spectra of MoS_2 grown on c-plane sapphire. In OM image (a), preferentially oriented grains are observed along $[1\overline{1}00]$ axis of sapphire substrate. PL spectra are shown in (b) where red and green spectra correspond to PL from aligned- and misaligned-grains, respectively. Asterisk (*) indicates the peaks from a sapphire substrate. The peak at ~1.76 eV corresponds to bound exciton in MoS_2 .

4.3.4. WS₂ growth on other sapphire planes

The TMD alignment is originated from the symmetrical matching of TMD lattice and the crystalline surface of c-plane sapphire. I investigated other crystalline planes, aplane $[1\overline{1}00]$ and r-plane $[1\overline{1}02]$ sapphire as growth substrates of WS₂. These substrates were placed in the CVD chamber together with c-plane sapphire. The optical micrographs of as-grown WS₂, the surface crystal structure of each sapphire plane, grain orientation distribution, and PL spectra are compared in Figure 4.11.

Totally different growth behavior was observed depending on the sapphire substrates. Although the growth condition is optimized for c-plane sapphire, clear difference can be seen for each of growth substrate, which indicates the crystal structure of a substrate



Figure 4.11. (a) OM images of CVD-grown WS_2 on different planes of sapphire substrates. (b) Surface crystal structure of each sapphire plane. (c) Grain orientation histogram of WS_2 grown on each sapphire plane. [11 $\overline{2}0$], [1 $\overline{1}00$], and [000 $\overline{1}$] axes are set as 0° for r-, c-, and a-plane sapphire, respectively. (d) PL spectra from the as-grown WS_2 grains.

significantly influences the growth of WS₂. I observed high WS₂ coverage on r-plane sapphire, but the grains are randomly oriented (Figure 4.11a). The intense PL emission was observed from these misoriented WS₂ grains (Figure 4.11d). These results reflect weak WS₂-sapphire interaction. On the other hand, WS₂ grown on a-plane showed a low WS₂ coverage, weak PL emission, and some preferential grain orientations. Thus, the a-plane sapphire induces stronger interaction with WS₂. As seen in Figure 4.11, the PL from WS₂ grown on r- and a-plane sapphire showed blue-and red-shift, respectively, with respect to that on c-plane sapphire. This could reflect the degree of strain induced in each WS₂.

4.3.5. Low temperature PL

For further analysis of WS₂ grains grown on sapphire, low temperature PL was measured, because the low temperature measurement allows us to understand intrinsic properties of TMDs which cannot be seen at room temperature due to thermally activated decay paths of excitons. Figure 4.12a shows PL spectra of as-grown, aligned (red) and misaligned (green) WS₂ grains measured at different temperatures in vacuum. Both the aligned and misaligned grains exhibited clear blue-shift of the PL peak from ~2.01 eV (300 K) to 2.10 eV (10 K).⁴⁷ Notably, all the PL spectra (both of aligned and misaligned grains) were found to be fitted with a single peak even at 10 K. Almost no subpeaks were observed even at the low temperatures. This result indicates that relaxation of excitons via trion formation or through defect states is strongly suppressed on the sapphire surface. This suggests that the exciton lifetime is very short on the sapphire due to strong van der Waals interaction. The PL intensity is plotted in Figure 4.12c. The PL intensity of the aligned WS₂ is lower than the misaligned WS₂ at all the

measured temperatures. Thus, WS_2 -sapphire interaction is stronger for the aligned grains.⁴⁰ The linewidth decreased monotonically for both the aligned and misaligned grains (Figure 4.12d), indicating the suppression of exciton scattering by phonons.

Figure 4.12b shows the PL spectra taken after transferring WS₂ grains on SiO₂ substrate. In these transferred samples, multiple peaks were observed at all the measurement temperatures. The PL peaks of the transferred WS₂ grains consist of the subpeaks originated in neutral excitons, trions, and defect states (or less likely bi-excitons)⁵¹ which were not detected in the as-grown samples. This signifies that the WS₂-substrate interaction is weakened, and the WS₂ is heavily doped due to the transfer process and from SiO₂ surface.^{10,39} It is also noted that the PL peak shapes of the



Figure 4.12. Normalized PL spectra of as-grown (a) and transferred (b) WS_2 grains measured at different temperatures. Plots of PL intensity (c) and FWHM (d) of as-grown and transferred WS_2 . Legends used in (c) are the same in (d).

aligned and misaligned grains are almost the same, indicating the comparable crystal quality of the aligned and misaligned WS_2 grains. Thus, the possibility of low crystallinity of the aligned WS_2 grains can be excluded from the reasons of the reduced PL intensity seen in the as-grown grains (see Figures 4.5a,d, 4.7c, and 4.9a).

4.3.6. Transport property

I measured the carrier transport properties after transferring the WS₂ grains on SiO₂ substrates for back-gate device operation. I used photolithography and lift-off processes to pattern metal electrodes (Ni/Au). Figure 4.13 insets are optical micrographs of WS₂ devices. Figure 4.13a displays representative transfer curves of FETs of the aligned and misaligned WS₂ grains. Clear n-type carrier transport with a high on/off ratio (10^5 - 10^6) was observed for both the aligned and misaligned WS₂ grains. The carrier mobility was calculated by using the following equation:

$$\mu = \frac{g_{\rm m}}{c_{\rm ox}V_{\rm d}}\frac{L}{W} \tag{4.1}$$

where $g_m = dI_d/dV_g$ is the transconductance obtained from the transfer curves, $C_{ox} = 11.5$ nF/cm² is the capacitance per unit area of 300 nm-thick SiO₂, V_d is the applied drain voltage, and *L* and *W* represent the length and width of the both aligned and misaligned grains. Here, the carrier mobility was calculated by assuming a trapezoidal channel shape to take into account the geometrical effect. Figure 4.13b shows the distribution of mobility and on/off ratio measured for different grains. No clear difference was observed between the aligned and misaligned grains. This is consistent with the low-temperature PL spectra measured after the transfer (see Figure 4.13b). Thus, it can be



Figure 4.13. (a) Transfer curves of aligned and misaligned WS₂ grains which were measured after the transfer on SiO₂/Si substrate for back-gate geometry ($V_d = 1$ V). The inset shows OM images of fabricated FET devices. (b) Distribution of electron mobility and on-off ratio.

conclude the overlayer deposition as well as PL quenching in the aligned WS₂ grains is originated in the strong van der Waals interaction between WS₂ and sapphire.

This work offers several potential applications for modifying electronic and geometric structures of WS_2 when epitaxially grown on c-plane sapphire substrates. Firstly, the strong coupling with the sapphire surface induce larger strain which can be utilized to chemically functionalize or dope the TMD materials. Secondly, the preferential growth of multilayer WS_2 can be used to make high mobility transistors, because it is known that monolayer TMD suffers strong influence from the substrate and top gate insulator. Thirdly, well aligned TMD grains lead to the effective grain stitching that can be developed to wafer scale single-crystal TMD sheet. Finally, the orientation-controlled growth can be used to tune the electronic and magnetic properties by stacking different materials with controlled orientations (moiré effect).

4.4. Conclusions

This chapter presents that the aligned WS_2 grains suffer more strain due to stronger van der Waals interaction with sapphire substrate as compared with the misaligned WS_2 . This strain induces the overlayer WS_2 deposition, resulting in the formation of multilayer WS_2 grains selectively on the aligned grains. The epitaxially grown grains showed reduced PL and Raman peaks. The DFT calculations indicate the strong coupling of WS_2 grains with the sapphire surface when the grains are aligned along the sapphire lattice. It was also found that the strong coupling with sapphire does not cause severe crystal defects in the epitaxially grown grains, even though strong strain is induced. This work gives new understanding of the influence of sapphire on the growth behavior of TMDs.

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Chemical Control of Electrical Transport Polarity of Transition Metal Dichalcogenides for Advanced Electronics

Abstract

Monolayers of TMDs have attracted a great interest for post-silicon electronics and photonics due to their high carrier mobility, tunable band gap, and atom-thick two-With the analogy to conventional silicon electronics, dimensional structure. establishing a method to convert TMD to p- and n-type semiconductors is essential for various device applications, such as CMOS circuits and photovoltaics. In this chapter, a successful control of the electrical polarity of monolayer WSe₂ by chemical doping is demonstrated. Two different molecules, 4-NBD and DETA, were utilized to convert ambipolar WSe₂ FETs to p- and n-type, respectively. Moreover, the chemically doped WSe_2 showed increased effective carrier mobilities of 82 and 25 cm²V⁻¹s⁻¹ for holes and electrons, respectively, which are much higher than those of the pristine WSe₂. The doping effects were studied by PL, Raman, XPS, and DFT calculations. Chemically tuned WSe₂ FETs were integrated into CMOS inverters, exhibiting extremely low power consumption (~0.17 nW). Furthermore, a p-n junction within a single WSe₂ grain was realized via spatially-controlled chemical doping. The chemical doping method reported in this chapter for controlling the transport properties of WSe₂ is expected to contribute to the development of TMD-based advanced electronics.

5.1. Introduction

Silicon forms the basis of the modern electronics, and controlled doping techniques have been widely used to fabricate both p- and n-type semiconductors made from silicon. Such spatially controlled doping techniques greatly contributed to the prevalence of Sibased CMOS circuits, photovoltaic cells, and optical sensors. Atomically thin TMD semiconductors, a part of the family of 2D materials, have attracted a great interest as a new alternative to silicon. This is because the monolayer TMDs show excellent electrical transport properties with high on/off ratios, strong PL in the visible range, and tunable band gap ranging from 1.5 to 2 eV depending on the combination of transition metals and chalcogenides.^{1–8} Recent development of the CVD method for the waferscale growth of TMDs have accelerated the research on their electronic and optoelectronic devices, including logic circuits, photodetectors, solar cells, and light emitting diodes.^{7,9–13}

However, the precise control of the polarity in TMD devices is still difficult, thus hindering practical applications. This is because most of TMDs, such as MoS₂ and WS₂, are n-type semiconductors due to the unavoidable presence of S vacancies.^{14,15} Attempts to convert n-type MoS₂ or WS₂ to p-type semiconductors have been reported. For example, partial substitution of Mo (or W) atoms with Nb and other metal atoms changed MoS₂ (or WS₂) to p-type character. However, the Mo_xNb_{1-x}S₂ devices show very weak gate dependence, reflecting a heavily degenerated state originated in the inserted Nb atoms.¹⁶ Different types of dopants, such as thiol-based molecules and Zn atoms, were also used to dope MoS₂.^{17,18} In these works, the threshold voltage (*V*_{th}) of devices shifts to higher gate voltages reflecting hole doping, but the main carriers are still electrons as observed from the corresponding transfer characteristics.^{17,18}

Being different from MoS₂ and WS₂, pristine WSe₂ is known to show ambipolar behavior. Chen et al. reported the p-doping effect of Au nanoparticles deposited on WSe_2 .¹⁹ However, the Au-coated WSe₂ still shows ambipolar behavior rather than ptype transport. Moreover, the PL from WSe₂ was strongly quenched by the nanoparticles, which is a disadvantage for optical applications. In ambient condition, p-type transport characteristics were observed for WSe₂.^{20,21} Different metal contacts, Pd and Ag were used to operate p-type and n-type WSe₂, respectively,^{22,23} though multiple electrode deposition processes are needed for the device fabrication. Partial substitution of Se atom with P was reported to induce p-type doping on WSe₂, but its influence on the electrical transport property was not studied.²⁴ In contrast, a mild plasma treatment was introduced to generate Se defect on WSe₂ to achieve n-type doping.²⁵ The defect engineering was utilized only for n-type doping but not for ptype doping and, more importantly, Se defects generally deteriorate the physical properties.

Consequently, it is difficult to obtain both p- and n-type transport properties using an identical TMD without changing the contact metal. Therefore, previous works combined different types of TMDs, such as p-type WSe₂ (or p-type MoTe₂) and n-type MoS₂, to fabricate CMOS circuits.^{13,26,27} Alternatively, very high effective gate voltages were applied to operate p- and n-type WSe₂ channels with the aid of ionic liquid.²⁸ Recently molecular doping technique was applied to various exfoliated TMD flakes, and both p- and n-type doping effects were observed. In this work, however, different metals, Pd and Ti, were used for p- and n-doping, and the use of exfoliated TMD flakes limited the practical applications.²⁹

In this chapter, the chemical doping of large-area CVD-grown WSe₂ monolayers was performed for selective conversion from ambipolar to p- and n-type semiconductors without changing the contact electrodes. The doping treatments were found to increase the effective carrier mobilities up to four orders of magnitude above the threshold region with respect to the pristine WSe₂, obtaining maximum values of 82 cm²V⁻¹s⁻¹ and 25 cm²V⁻¹s⁻¹ for holes and electrons, respectively. The chemically doped WSe₂ monolayers were integrated into CMOS circuits that showed extremely low power consumption (~ 170 pW with V_{gain} of ~10). Moreover, spatially controlled doping was also achieved within a single WSe₂ grain to obtain separated p- and n-doped regions. The WSe₂ grain with the artificial p-n junction showed clear rectification behavior and optical response to visible light, implying its potential for photodetection applications.^{30–34}

5.2. Experimental method

5.2.1. CVD growth of WSe₂ and WS₂

Monolayer WSe₂ was synthesize by Prof. Miyata's laboratory in Tokyo Metropolitan University using an improved salt-assisted CVD method with WO₃ and Se powders. KBr powder was used as a growth promoter.²² In a tubular furnace, WO₃ and KBr were placed 2 cm and 9 cm upstream of the SiO₂/Si substrate. Temperatures of the substrate and the WO₃ were gradually increased to the growth temperature (780 °C) with a N₂/H₂ gas mixture at a flow rate of 400/1.2 ccm. After the substrate reached the growth temperature, Se was heated to 300~370 °C by using another furnace to supply Se vapor to grow WSe₂. After 10 min reaction, the sample was cooled down immediately.

In our laboratory, WS_2 was synthesized by ambient pressure CVD method using WO_3 powder and S power. Growth promoter was not used for WS_2 growth and c-plane sapphire was used as a growth substrate. The details of the WS_2 synthesis are reported elsewhere.³²

5.2.2. Chemical doping

For p-type doping, a WSe₂ sample was immersed in aqueous solution of 4-NBD with concentrations of 10~100 mM and left in dark environment. The sample was kept in the solution at 60 °C for 30 minutes. For n-type doping, a WSe₂ sample was exposed to DETA vapor (~460 Pa vapor pressure) at 70 °C for 30 minutes.³⁵ In the closed petri dish with ~100 cm³ volume, 300 μ L of DETA was placed for applying DETA vapor to WSe₂.

5.2.3. Characterization

Optical micrographs were taken using an optical microscope (Nikon Eclipse ME 600), and the AFM height images were collected using a Nanoscope V Multimode SPM (Bruker Co.). PL and Raman spectra were measured using a confocal Raman spectroscope (Tokyo Instruments Inc., Nanofinder 30) under 532 nm laser excitation. XPS measurements were conducted in the beamline 10 (BL10) of SAGA Light Source (SAGA-LS) at 10⁻⁸ Pa with the incident energy of 600 eV. For the XPS measurement, WSe₂ grains were transferred to an Au-coated SiO₂/Si substrate, followed by respective doping processes.

5.2.4. DFT calculation

Electronic structure calculations of DETA, 4-NBD, and WSe₂ were conducted using the DFT implemented in the STATE code.³⁶ The exchange correlation interaction among interacting electrons is expressed by the generalized gradient approximation (GGA) with the PBE functional form. For the electron-ion interaction, ultrasoft pseudopotentials generated by the Vanderbilt scheme was adopted. The valence wave functions and deficit charge density were expanded in conjunction with a plane-wave basis set, applying cutoff energies of 25 and 225 Ryd, respectively. To obtain qualitative insight into the possibility of the charge transfer from the dopant molecules, 4-NBD and DETA, to WSe₂, the absolute values of the electron energy levels of isolated 4-NBD, DETA, and WSe₂ was evaluated by DFT calculations. For the evaluation of the energy level with respect to the vacuum, super cell calculations were conducted for isolated molecules and WSe₂ in which the molecules or WSe₂ are separated their periodic images by at least 1.5 nm vacuum spacing. For WSe₂, a 4×4 lateral periodicity was considered to evaluate the effect of chalcogen defects on the electronic energy band. The geometric structures were optimized until the remaining force acting on the atoms becomes smaller than 0.005 Ryd/Å.

5.2.5. Device fabrication and measurement

As the CVD process slightly damages the surface SiO₂ layer and gave gate leakage, the as-grown WSe₂ grains were transferred onto a fresh SiO₂(90 nm)/Si substrate using polystyrene as a support layer.³⁷ Prior to the transfer, the SiO₂ /Si substrates were cleaned by sonicating in acetone and isopropyl alcohol for 10 minutes each. For the fabrication of FETs, the transferred WSe₂ was spin-coated by photoresist and patterned using photolithography, followed by electron beam evaporation of Au(30 nm)/Ni(1 nm) electrodes (Kennix KB-750) and lift-off process. Carrier transport properties were measured using a semiconductor parameter analyzer (Keysight Technologies B1500A) in vacuum (~10⁻⁴ Pa) at room temperature. To fabricate the inverter circuit, two WSe₂ FETs were externally connected by Au wire (99.95 %, 300 μ m in diameter). The inverter performance was also measured in a vacuum (~10⁻⁴ Pa) at room temperature. For measuring the optical response of a p-n device, a 532 nm laser (Showa Optronics Co.) was used together with ND filters for controlling the laser power.

5.3. Results and Discussion

5.3.1. CVD growth and chemical doping of WSe₂

Figure 5.1 shows the experimental procedure employed to prepare p- and n-doped WSe₂. As written in the experimental section, triangular grains of monolayer WSe₂ were synthesized on SiO₂/Si substrates by thermal CVD using WO₃ and Se precursors. An optical microscope image of the as-grown WSe₂ is shown in Figure 5.1b. To avoid gate leakage, the as-grown WSe₂ was transferred to a fresh SiO₂ (90 nm)/Si substrate.³⁷ After the transfer, FETs with WSe₂ channels were fabricated by photolithography, Ni/Au electrode deposition, and lift-off processes.

For the p-type doping, WSe₂ was immersed in aqueous solution of 4-NBD (Figure 5.1c), while WSe₂ was exposed to DETA vapor for the n-type doping (Figure 5.1d). AFM images of pristine, 4-NBD and DETA doped WSe₂ grains are shown in Figure 5.2, where the monolayer thickness (~1 nm) and dopant deposition can be seen from the height profile. The as-grown, pristine WSe₂ had a thickness of ~ 1 nm and showed an intense PL at 760 nm, reflecting the direct band gap nature. These results confirm the



Figure 5.1. (a) Schematic of CVD growth of WSe₂ on SiO₂/Si substrate. (b) Optical micrograph of as-grown triangular WSe₂ grains. (c,d) Chemical doping processes using 4-NBD and DETA molecules for p- and n-type doping, respectively. (e) Transfer curves of pristine (black) and 4-NBD doped (red) and DEA doped (blue) WSe₂ measured with $V_d = 5$ V.

growth of monolayer WSe₂. Figure 5.1e shows the device structure of WSe₂-FET and the corresponding transfer characteristics measured before and after the chemical doping. Drain current (I_d) was measured as a function of gate voltage (V_g) applied from back side of the device through a SiO₂ layer. The pristine WSe₂ (black curve) showed ambipolar behavior with a slightly n-doped character. After the contact with 4-NBD and DETA dopants, the WSe₂ device changed to p- (red curve) and n-type (blue curve), respectively. The V_{th} shifted to more positive (p-doped) and more negative (n-doped) voltages after each doping. This implies that holes and electrons are



Figure 5.2. AFM height images of pristine (a), 4-NBD doped (b), and DETA doped (c) WSe₂ grains. (d-f) Magnified images of the marked positions. (g-i) Three-dimensional AFM images of those shown in (d-f).



Figure 5.3. Transfer characteristics of WSe₂ FETs before doping (b) and after doping with 4-NBD (a) and DETA (c) with y-axis linear scale. Drain voltage is 5 V, and SiO₂ thickness is 90 nm.

accumulated not only by the gate voltage but also by the chemical doping effect. The transfer curves with linear scale of the drain current are shown in Figure 5.3.

5.3.2. Electrical transport properties after doping

It is noted that the slopes of the transfer curve after the doping are much steeper than those of the undoped devices (see Figure 5.1e, in which *y*-axis scale is logarithmic). Figure 5.4a,b shows more detailed analysis of the carrier transport of the doped WSe₂ monolayers. To avoid device-to-device variations, I measured the electrical transport of the same WSe₂ channels before and after the doping process. Because the V_{th} are different for the pristine and doped WSe₂ samples, the overdrive gate voltage (V_{g} - V_{th}) was used in the graphs for better comparison of the gate voltage dependence. The effective field-effect carrier mobility (μ) was calculated by using the following equation:

$$\mu = \frac{g_{\rm m}}{C_{\rm ox}V_{\rm d}}\frac{L}{W} \tag{5.1}$$

where $g_m = dI_d/dV_g$ is the transconductance obtained from the transfer curve; $C_{ox} = 38.3$ nFcm⁻² is the capacitance per unit area of the 90 nm-thick SiO₂; V_d is the applied drain voltage; *L* and *W* represent the effective length and width of the channel. The effective carrier mobility was calculated by assuming a trapezoidal channel shape to take into account the geometrical effect. It is seen that the carrier mobility after the doping with both 4-NBD and DETA increased by more than 3-4 orders of magnitude.

Figure 5.4c compares the effective carrier mobilities of WSe₂ devices (20 devices were measured for each dopant). The measured hole and electron mobilities of the pristine WSe₂ were found to be rather low. This can originate in intrinsic defects in



Figure 5.4. (a,b) Carrier mobilities of pristine (black) and doped (red and blue) WSe₂ devices plotted as a function of the overdrive gate voltage. The doping-induced change of the carrier mobility is shown by green curve. Insets show the optical images of the devices (scale bar: $20 \ \mu m$). (c) Distributions of maximum hole and electron mobilities of pristine and doped WSe₂.

the as-grown WSe₂ grains, degradation due to air exposure, and/or possible damages induced by the device fabrication process, as based on our experience WSe₂ is easier to be oxidized than MoS₂ and WS₂. Interestingly, both the hole and electron mobilities drastically increased after the chemical doping, as can be seen in Figure 5.4c. The maximum carrier mobilities measured after the p- and n-type doping were 82 cm² V⁻¹s⁻ ¹ (for holes) and 25 cm² V⁻¹s⁻¹ (for electrons), respectively. It is speculated that influence of defects existing in WSe₂ is reduced by the chemical doping due to increased carrier concentration, contributing to the increase of the measured carrier mobilities. In addition, lowering the Schottky barrier height due to the chemical doping-induced Fermi level shift should be considered as another source of the increased effective carrier mobility. In previous literatures, 23,38-41 it has been proposed that the interface between the 2D material and the electrode plays an important role in the carrier transport due to strong modification of the Fermi level of the 2D material. Thus, it is highly likely that reduction of the contact resistance induced by the chemical doping largely contributes to the increased carrier mobility of WSe₂ FETs. The enhanced carrier mobilities together with the polarity control would allow prospective use for high mobility switching devices.

5.3.3. Spectroscopy and DFT calculations

Figures 5.5a,b compare the PL spectra of monolayer WSe₂ before and after the doping with 4-NBD and DETA, respectively. The 4-NBD doping was found to increase the PL intensity and decrease the FWHM from 64 to 53 meV. The PL peak from the pristine WSe₂ can be deconvoluted with exciton and trion peaks located at ~1.64 eV and ~1.59 eV, respectively, as shown in Figure 5.6. As reported previously, it is considered that the trion in the pristine WSe₂ is a negative trion (A⁻),⁴² being consistent with the experimental observation that electrons are main charge carriers in the absence of a gate voltage (see Figure 5.1e). The 4-NBD doping increased the intensity of the exciton peak while reduced that of the trion (Figures 5.5a and 5.6a). The increased exciton peak can be accounted for by the reduction of the negative charges due to the supply of holes from 4-NBD, resulting in the decreased ratio of negative trions. The trion peak in the doped WSe₂ can be either remaining negative trions or newly formed positive



Figure 5.5. (a,b) PL spectra of pristine and doped WSe₂ measured on SiO₂/Si substrates. The identical WS₂ grains were used to monitor the doping-induced spectrum change. (c) Raman spectra of pristine and doped WSe₂.



Figure 5.6. Peak fitting analysis of the PL spectra of the pristine and doped WSe₂. The original PL spectra are displayed in Figure 5.5a,b. Orange and magenta curves correspond to exciton and trion peaks, respectively.

trions. Although the origin of the trion peak is unclear, the PL spectra supports the hole doping to the WSe₂ induced by 4-NBD molecules.

The DETA doping gave a completely different change in the PL spectrum (Figure 5.5b). The PL peak was strongly suppressed and red-shifted by \sim 50 meV. The fitting analysis indicates that the spectral weight of the trion peak increased, while the exciton peak decreased significantly (see Figure 5.6). This can be explained by the increased concentration of negative trions, while preventing the formation of neutral excitons as a result of electron doping.^{29,43} In this case, the observed trion peak should be originated

in the negative trions. The observed peak shift of ~ 50 meV after the DETA doping corresponds to the trion binding energy of WSe₂.^{44,45}

Effects of the chemical doping on the Raman spectrum were also studied, as shown in Figure 5.5c. The E^{1}_{2g} band which corresponds to the in-plane vibration mode clearly shifted to higher and lower wavenumbers upon 4-NBD and DETA doping, respectively. The position of the out-of-plane mode (A_{1g} band) exhibited a similar tendency. Such doping-induced Raman shift is consistent with previous studies,^{46,47} and can be explained by the change of the electron density in monolayer WSe₂; the n-type doping increases the electron-phonon scattering due to a higher electron concentration, while the p-type doping decreases the electron-phonon scattering.^{46,47} The decrease of the A_{1g} peak intensity is also in accordance with the existing literature.^{19,46} Thus, the Raman spectroscopy also supports the p/n selective doping in our CVD-grown WSe₂.

For further understanding of the changes of the WSe₂ electronic structure induced by 4-NBD and DETA molecules as well as the charge transfer phenomena, XPS measurement and DFT calculations were performed. Figure 5.7a,b shows XPS W_{4f} and Se_{3d} spectra of the pristine and the doped WSe₂. The chemical doping with 4-NBD molecule shifted the binding energies of both W_{4f} and Se_{3d} peaks to lower energies (see Figure 5.7c for the XPS peak positions). This shift implies that the Fermi level of the WSe₂ is moved downward to the valence band edge. On the other hand, the DETA doping increased the binding energies of both W_{4f} and Se_{3d} peaks, suggesting the shifting of the Fermi energy upward to the conduction band edge. These observed energy shifts are consistent with the previous literatures reporting the selective p/ndoping to MoS₂.^{29,48,49} Therefore, the changes of the binding energies after the 4-NBD and DETA adsorption qualitatively supports the p-/n-selective doping by tuning the Fermi energy of the monolayer WSe₂.



Figure 5.7. (a,b) XPS spectra (W_{4f} and Se_{3d}) of pristine and doped WSe₂. (c) XPS peak positions of pristine and doped WSe₂ (in eV). The number in the parenthesis indicates the relative peak position with respect to the pristine WSe₂. Note that the energy shift is dependent on the degree of doping (or amount of dopant).

Figures 5.8a,b display the energy levels of WSe₂, 4-NBD, and DETA, as determined by the DFT calculations. In the case of 4-NBD, the diazonium molecule forms a complex with BF₄⁻ anion.⁵⁰ Therefore, in the present calculation, the diazonium molecule is assumed to possess a singly occupied molecular orbital (SOMO) located between the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO). As seen in Figure 5.8a, the relative position of SOMO of 4-NBD and valence band edge (VBE) of WSe₂ supports the hole doping in WSe₂ via the electrons transfer to the SOMO.

DETA molecule has a closed-shell electronic structure and its HOMO locates within the band gap of pristine, monolayer WSe_2 (see Figure 5.8b for the band energy of the pristine WSe_2). However, the HOMO level is ~1.59 eV below the conduction band edge (CBE) of WSe_2 (Figure 5.8b). It is thought that there are several mechanisms



Figure 5.8. (a) Band diagram of pristine WSe₂, 4-NBD and DETA molecules. The energy levels were calculated by DFT. (b) Molecular orbital levels of 4-NBD and DETA molecules, and band structures of pristine WSe₂, Se-deficient WSe₂, and S-deficient WS₂ monolayers. Broken lines show newly appeared defect states. The energy levels were calculated by the DFT.



Figure 5.9. Atomic model of WSe₂ monolayer with single Se vacancy (indicated by the arrow) in 4×4 cell, which is used to calculate the defect-related band structure shown in Figures 5.8a,b. The Se vacancy is introduced to the upper Se site of the monolayer WSe₂. Green and gray atoms represent Se and W, respectively.
which assist the electron transfer from DETA to WSe₂. First, Se vacancies are frequently observed in WSe₂ samples,⁵¹ which can be the origin of the relatively low effective carrier mobility observed in our pristine WSe₂ (see Figure 5.4c). Therefore, in Figure 5.8a, the band structure of the monolayer WSe₂ containing Se vacancies (one Se atom was removed from 4×4 unit cell, which corresponds to ~3% Se vacancy (see Figure 5.9 for the atomic structure)) is displayed. As seen in Figure 5.8a, the introduction of Se vacancies produced a new state, which is associated with the defect, at 0.61 eV below the CBE. This defect state is expected to act as an intermediate trap state through which the electrons transfer from DETA to WSe₂, thus assisting the n-type doping. Although the exact density of structural defects including Se vacancies is not clear, the post-growth treatment, such as exposure to ambient and the device fabrication process, may give more defects with variety of defect structures. It is noted that a STEM study revealed the presence of various crystal defects in monolayer MoS₂, suggesting the formation of localized electronic states within the band gap.⁵² Other DFT calculations also indicated the formation of a defect state (or in-gap state) within the band gap when chalcogen vacancies are introduced in MoS₂ and MoSe₂.^{53,54} Therefore, it is speculated that there is a possibility that the defect-related bands appear just above the HOMO level of DETA or close to the HOMO level, which can account for the experimentally observed n-type doping. The second possibility is the coordination of paired electrons of N atoms in DETA to W atoms of WSe₂, introducing the charge transfer to WSe₂. As this model is also based on the presence of Se vacancies to access a W metal, we speculate that the imperfect WSe₂ structure is related to the observed n-type doping induced by DETA molecules.

To support our interpretation of the doping mechanism in WSe₂, I also treated monolayer WS₂ with 4-NBD and DETA for the sake of comparison. The PL spectral



Figure 5.10. Investigation of chemical doping effect on WS_2 by PL analysis. (a) An optical micrograph of CVD-grown WS_2 on sapphire substrate. PL spectra of 4-NBD doped (b), pristine (c), and DETA doped (d) WS_2 . Black curves are measured spectra, while orange and magenta curves are fitted curves which correspond to exciton and trion peaks.



Figure 5.11. Transfer characteristics of WS₂ FETs before and after doping with 4-NBD (a) and DETA (b). $V_d = 1V$, SiO₂ thickness is 300 nm.

change is presented in Figure 5.10. A similar tendency was observed for monolayer WS₂, with an increased ratio of the exciton peak after 4-NBD doping, and the significant increase of the trion peak after DETA doping.^{55,56} As shown in Figure 5.11a, the WS₂ FET showed the shift of V_{th} to positive gate voltages after dipping in 4-NBD solution, but it did not show p-type transport character. According to DFT calculations (see Figure 5.8b), the VBE of WS₂ is located below the SOMO of 4-NBD, making it difficult to inject holes from 4-NBD to WS₂. This is consistent with the FET result in which clear p-type transfer characteristics was not observed in 4-NBD doped WS₂ system (see Figure 5.11a). The exposure to DETA vapor shifted the V_{th} of WS₂ to negative voltages, indicating the n-type doping (Figure 5.11b). This agrees with the calculated band structure, as the defect state of WS₂ is closer to the HOMO level of DETA than the defect state of WSe₂ is suitable for obtaining clear p/n selective doping with the employed dopant molecules.

Previously, formation of a covalent bond was proposed between a 4-NBD molecule and monolayer TMD,^{57,58} similar to the case of graphene and 4-NBD.⁵⁰ In the case of graphene, the covalent bond formation can be recognized from the appearance of defectrelated Raman D-band. However, as the Raman spectrum of TMDs is not so sensitive to defects or covalent bond formation, it is difficult to experimentally discuss how the dopant molecules attach on the WSe₂. The p-doping effect in the 4-NBD doped WSe₂ was still observed after thorough washing with water, while n-type doping effect by DETA disappeared easily after washing with water. Thus, there is a possibility of chemisorption and physisorption for 4-NBD and DETA doping, respectively. However, the XPS spectra of 4-NBD doped WSe₂ (Figure 5.7a,b) did not show any apparent changes in the peak shapes except for shift of the peak positions to lower binding energies. There is no solid evidence for the possible chemisorption of 4-NBD molecules on WSe₂ surface. Therefore, further study is necessary to understand the detailed doping mechanism by employing atomic scale imaging techniques, such as transmission electron microscope (TEM) and scanning tunneling microscope (STM).

5.3.4. Fabrication of a CMOS inverter

The controlled p- and n-type doping in WSe₂ was developed to fabricate a CMOS inverter. Figure 5.12a shows the input voltage (V_{in}) dependence of the output voltage (V_{out}) collected with different supply voltages (V_d). Here, the p- and n-FETs were externally connected in series by an Au wire to form the inverter, with the common back-gate being the input (V_{in}) (See Figure 5.12a inset). As can be seen in Figure 5.12a, the V_{in} signal was properly inverted. The V_{out} was kept at a value of V_d for V_{in} lower than ~ -28 V and dropped to ~0 V for V_{in} higher than ~-25 V, with a sharp inversion. Figure 5.12b shows the voltage gain, defined as $V_{gain} = -dV_{out}/dV_{in}$, which is one of the most important performance parameters of CMOS inverters. As can be expected from the sharp transition of V_{out} (Figure 5.12a), a maximum V_{gain} as high as 10 was achieved.



Figure 5.12. CMOS inverter operated with p- and n-doped WSe₂. (a) V_{out} plotted as a function of V_{in} . Inset shows the circuit diagram of the inverter. (b,c) Voltage gain and power consumption of the CMOS inverter.



Figure 5.13. Detailed study of inverter performance. (a-c) Transfer characteristics of 4-NBD and DETA doped WSe₂. The gap between V_{th} of two transfer characteristics is the largest in (a) and smallest in (c). (d-f) Inverter performance corresponding to (a-c).

To the best of my knowledge, this is the first report of a CVD-grown WSe₂ CMOS inverter where both p- and n-channels are formed by respective doping treatments.^{13,26,59–63} Considering that this CMOS circuit was fabricated on a SiO₂ (90 nm)/Si substrate with a back-gate geometry ($C_{ox} = 38.3 \text{ nF cm}^{-2}$), the obtained gain can be further improved by increasing the capacitance via top-gate operation for a more effective switching control.^{13,23}

One of the biggest benefits of CMOS circuits is their low power operation. Figure 5.12c shows the power consumption of the WSe₂-based inverter as a function of the input voltage. I found that my inverter can operate with an extremely low power consumption of ~170 pW. This value is three orders of magnitude lower than those from previous references using TMD channels.^{13,26,59} I attribute this low power consumption to the optimized overlap of the transfer curves of p- and n-FETs, allowing to keep the current flow through the inverter low. As shown in Figure 5.13, the power consumption of my inverters increased with increasing the overlap of the transfer curves of p- and n-FETs. Liu et al. reported 1 µW power consumptions when the transfer curves of the individual FETs overlap at a drain current of ~1000 nA.⁶² Zhang et al. reported 3.1 nW power consumptions when the overlap of the individual FETs transfer curves occurs at 1 nA.⁵⁹ In contrast, my p- and n-FETs overlapped at ~0.1 nA, thus efficiently minimizing the leakage current to prevent the power loss. Therefore, controlled doping through the optimization of the exposure time and dopant concentration is required for both p- and n-type WSe₂ to achieve low power CMOS inverters.

5.3.5. Fabrication of a p-n junction

For practical applications of TMD materials in electronic and optoelectronic devices, the p-n junction is the most important and basic working unit. Therefore, I fabricated a p-n junction within a single WSe₂ grain through spatially controlled p- and n-doping. The fabrication process is depicted in Figure 5.14. In brief, 4-NBD doping was performed through a photoresist window, followed by deposition of SiO₂ as a protective layer. Then, another window is opened in the SiO₂ layer for the successive DETA doping. Figures 5.15a,b show a false colored optical image and the corresponding



Figure 5.14. Fabrication process of the spatially controlled p-n junction in single WSe₂ grain. PR is photoresist.



Figure 5.15. (a,b) Optical micrograph and schematic of a p-n junction made by the spatially controlled p/n doping within single WSe₂ grain. (c) Output curves of the pristine and p/n doped WSe₂ monolayer device. The device with p-n junction shows clear rectification behavior. (d) Light response of the WSe₂ device with p-n junction.

schematic model of the p-n doped single grain device fabricated in this work. Output characteristics of the WSe₂ before and after the formation of the p-n junction are shown in Figure 5.15c. While the pristine WSe₂ grain exhibited an almost linear I_d - V_d curve with high resistivity (black curve), after the spatially controlled p-n doping, the device showed a clear rectifying behavior (red curve). Such asymmetric transport behavior is a strong evidence of the formation of a p-n junction.^{30,31,64} Under illumination with visible light (532 nm, 6.4 mWcm⁻²), I observed a clear photoresponse (Figure 5.15d), demonstrating its potential in photodetection applications.

5.4. Conclusions

In this chapter, the effective and controlled chemical doping of CVD-grown WSe₂ was demonstrated. Spectroscopic analyses and DFT calculation proved that 4-NBD and DETA molecules allow efficient charge transfer to WSe₂, changing the ambipolar WSe₂ to p- and n-type semiconductors, respectively. The chemical doping with 4-NBD and DETA was found to significantly improve the effective carrier mobilities of both holes and electrons. The integration of the p- and n-type WSe₂ was also demonstrated for the operation of a low power CMOS inverter, originating from the high control of the degree of the doping. Moreover, the position selective doping has been developed to obtain the p-n junction within a single WSe₂ grain. The p-n doped WSe₂ grain showed clear rectification behavior and optical response, as required for optoelectronics applications. This chemical doping strategy can offer a versatile approach to modify the electrical property of TMD materials for various device applications.

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Chapter 6

Conclusions and Future Outlook

6.1. Conclusions

In this thesis, synthesis and applications of 2D TMD materials are demonstrated. In Chapter 3, epitaxial growth of WS_2 on c-plane sapphire substrate using H_2 carrier gas is demonstrated. By strong reducing effect of H_2 on the reaction of precursors, crystal quality of WS_2 is significantly improved. H_2 gas also changes the surface termination of c-plane sapphire, resulting in epitaxial growth of WS_2 on sapphire. Epitaxially grown WS_2 shows atomically smooth stitching between merging grains, which is revealed by detailed STEM study, and they show improved chemical stability against oxidation as well as good electrical transport properties across seamlessly stitched grain boundaries.

In Chapter 4, influence of c-plane sapphire substrate on the growth behavior of WS₂ is investigated. Due to similar hexagonal lattice structures of c-plane sapphire and WS₂, larger lattice strain is induced in the epitaxially aligned grains by strong van der Waals interaction with underlying sapphire. It causes PL quenching and preferential overlayer deposition on the epitaxially aligned WS₂ grains. Low-temperature optical study and FET measurement reveal that van der Waals interaction-induced lattice strain does not cause a degradation of crystal quality. DFT calculations on lattice orientation-dependent relaxation energies on sapphire well support the experimental data.

In Chapter 5, chemical tuning of electrical transport polarity of CVD grown WSe₂ is demonstrated. By using 4-NBD and DETA molecules as p- and n-type dopants, electrical polarity is successfully controlled to p- and n-type, respectively. In addition, the effective carrier mobility significantly increases after each doping treatment, and DFT calculation supports the observed doping. Using chemically doped WSe₂ FET devices, a CMOS inverter circuit is fabricated, which shows high voltage gain as well as extremely low power consumption. Furthermore, by combining chemical doping and lithography techniques, a p- n junction is fabricated within a single triangle grain of WSe₂, being half of the grain selectively p-doped and another half n-doped. This p-n junction shows clear rectification behavior together with light response.

6.2. Future outlook

As a result of devoted effort by many researchers, TMD research has dramatically advanced for last several years. Recent development of CVD technology realized wafer-scale synthesis of uniform TMD films.^{1,2} Also new physical properties have been continuously discovered. Especially, optical and electronic properties of twisted TMD heterostructures with unique superlattice formation have been on the focus of many researchers.^{3–5} As already known, atomically thin TMD monolayers possess large spin-orbit splitting in their valence and conduction bands.⁶ Existence of degenerate energy states in a *K* valley makes TMDs have an additional spin degree of freedom. Furthermore, TMDs have degenerate inequivalent *K* points (*K* and *K*' or K^+ and K^-) in their hexagonal Brillouin zone which differently respond to circularly polarized light.⁷ Thus, this valley-dependent optical selectivity lets TMDs have an additional valley degree of freedom. Consequently, the spin and valley degree of freedoms observed in TMDs cause unique optical and electronic properties, such as formation of spin,

valley polarized exciton and spin, valley hall effect.⁸ Additional degree of freedoms in carriers make it possible to realize new paradigm of data manipulation with orders of magnitude higher processing capacity by using combination of electrical charge, spin, and valley. To realize this, further research is necessary. First, the crystal quality of CVD grown TMDs must be improved. In spite of the development of CVD technology, still the defect density in CVD-grown TMD crystals is high. In carrier dynamics in conduction and valence bands, these structural defects act as scattering sites and degrade the polarization controllability. Thus, more study is needed to further improve the crystal quality of CVD TMDs. Second, good insulating materials are necessary. As proved by many literatures, TMDs show more intrinsic and improved physical properties when encapsulated between h-BN layers.^{9–11} Due to the subnanometer thickness of monolayer TMDs and their resultant sensitivity to the dielectric surrounding, it is very important to protect TMDs using an insulator. For this, h-BN, which is a 2D insulator without dangling bond, is being widely used. In spite of the development of h-BN CVD technique¹²⁻¹⁴, it is still difficult to grow large, uniform, and highly crystalline h-BN so that most of people are using the small flakes of exfoliated h-BN. Therefore, further development of h-BN synthesis technique is highly required for continuing advance of 2D research. As insulating materials, there are additional choices other than h-BN, such as large band gap 2D materials, Cu(OH)₂, Ni(OH)₂, Mg(OH)₂, TaIc, GaS,¹⁵ covalent organic frameworks,¹⁶ and insulating 2D polymers.¹⁷ Third, for continuing thrive of 2D research, it will be also important to keep looking for new physical properties from known materials and new materials with interesting properties. Research on the stacking of various 2D materials to investigate emergence of new physical property (Twistronics) is just on the beginning stage. There are many on-going works in this field so that significant advance is expected in the near future. There are other emerging 2D materials, for example, Nb_3X_8 family (X = Cl, Be, or I).¹⁸ Similar to TMDs, Nb₃X₈ also have a two-dimensional layered

structure. Theoretical studies have expected variable band gap and ferromagnetism depending on their composition.¹⁸ Also 2D perovskites have attracted attention. When thinned down to nanometer thickness, perovskites show different physical properties from the bulk because of the quantum confinement effect.^{19–21} From this, we can expect thinning of various materials to see their property change in quantum confined condition.²² Furthermore, there are many other promising 2D candidates, such as synthetic 2D polymer,¹⁷ graphyne,^{23,24} borophene,^{25–27} germanene,^{28,29} phosphorene,^{30,31} and MXenes,^{32,33} which have been proved to possess promising physical properties although most of them need more detailed experimental investigations.

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List of publications

• Chapter 3

Ji, H. G.; Lin, Y.-C.; Nagashio, K.; Maruyama, M.; Solís-Fernández, P.; Sukma Aji, A.; Panchal, V.; Okada, S.; Suenaga, K.; Ago, H. Hydrogen-Assisted Epitaxial Growth of Monolayer Tungsten Disulfide and Seamless Grain Stitching. *Chem. Mater.* **2018**, *30*, 403–411.

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