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INTERFACIAL BEHAVIOR IN POLYMER DERIVED CERAMICS AND SALT WATER PURIFICATION VIA 2D MOS₂

by

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A dissertation submitted in partial fulfillment of the requirements for the degree of Doctor of Philosophy in the Department of Materials Science and Engineering in the College of Engineering and Computer Science at the University of Central Florida

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ABSTRACT

In the present dissertation, the behavior of the internal potential barrier in a polymer-derived amorphous SiAlCN ceramic was studied by measuring its complex impedance spectra at various dc bias as well as different testing and annealing temperatures. The complex impedance spectra of the polymer-derived a-SiAlCN were measured under various dc bias voltages in a temperature range between 50 and 150 °C, as well as different annealing temperatures (1100-1400 °C). All spectra, regardless of temperature and bias, consist of two semi-circular arcs, corresponding to the free-carbon phase and the interface, respectively. The impedance of the free-carbon phase is independent of the bias, while that of the interface decreased significantly with increasing dc bias. It is shown that the change of the interfacial capacitance with the bias can be explained using the double Schottky barrier model. The charge-carrier concentration and potential barrier height were estimated by comparing the experimental data and the model. The results revealed that increasing testing temperature led to an increased charge-carrier concentration and a reduced barrier height, both following Arrhenius dependence, whereas the increase in annealing temperature resulted in increased charge-carrier concentration and barrier height. The phenomena were explained in terms of the unique bi-phasic microstructures of the material. The research findings reveal valuable microstructural information of temperaturedependent properties of polymer derived ceramics, and should contribute towards more precise understanding and control of the electrical as well as dielectric properties of polymer derived ceramics.

Furthermore, the desalination performances and underlying mechanisms of two-dimensional CVD-grown MoS₂ layers membranes have been experimentally assessed. Based on a successful large-area few-layer 2D materials growth, transfer and integration method, the 2D

 MoS_2 layers membranes showed preserved chemical and microstructural integrity after integration. The few-layer 2D MoS_2 layers demonstrated superior desalination capability towards various types of seawater salt solutions approaching theoretically-predicted values. Such performances are attributed to the dimensional and geometrical effect, as well as the electrostatic interaction of the inherently-present CVD-induced atomic vacancies for governing both water permeation and ionic sieving at the solution/2D-layer interfaces.

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LIST OF ABBREVIATIONS

AC Alternating Current

AFM Atomic Force Microscopy

ASB Aluminum Tri-Sec-Butoxide

CVD Chemical Vapor Deposition

DC Direct Current

MD Molecular Dynamics

MEMS Micro-Electro-Mechanical Systems

PDC Polymer Derived Ceramic

TEM Transmission Electron Microscopy

XRD X-Ray Diffraction

CHAPTER 1: INTRODUCTION

The roles of interfaces of nanomaterials have significant contributions towards the properties, performances and applications of various categories of materials, either conventional or novel. The interfacial characteristics could stem from the responses of the intrinsically-present intraphase interfaces upon external stimuli (e.g., electric field and magnetic field), or they may also arise from the interactions between the surfaces of materials with outside matters (e.g., solutions, oxygen, and irradiation).

Polymer derived ceramics (PDCs) are a novel class of amorphous materials synthesized through the controlled thermal decomposition of polymeric precursors. PDCs exhibit unique microstructural features and desirable electrical/thermal/mechanical properties with promising applications compared to conventional ceramic materials prepared via powder metallurgy route. Prominent PDCs properties include: superior thermal stability, excellent oxidation/corrosion resistance and high temperature multi-functionalities. In addition, the direct polymer-to-ceramic processing route makes PDCs particularly feasible for fabricating ceramic (micro)-components with complex structures/shapes.

Reliable and accurate real-time pressure/temperature monitoring under high temperature and harsh environments (e.g., gas turbines, nuclear reactors, and high-speed vehicles) has been on high demand for device damage prevention as well as structural performance improvement. Surviving under harsh environments (e.g., pressurized, oxidative, radiative and corrosive) demands materials that satisfy both structural stability and functionality/properties reliability, which most currently available materials fall far short of.

Recent research and development efforts have explored several types of materials for harsh-environments sensing applications. Most notably among them are semiconducting silicon and silicon carbide due to proven controllable electric properties and well-developed microfabrication capabilities. However, applications of Si-based sensors are significantly restricted to a very limited operational temperature range due to severe material degradation,⁴ not to mention when the environment involves corrosive atmospheres as well.

In the first part of this dissertation, we studied the microstructure, properties and impedance behaviors of polymer-derived SiAlCN ceramics, with a special attention on the internal potential barrier characteristics of this nano-domained two-phase material.

In a further attempt of investigating the interface/surface interactions in nanomaterials, this dissertation has also studied the interfacial interactions between the surfaces/interiors of two-dimensional (2D) MoS₂ layers upon contact with seawater solutions, with an overall goal of the development of viable 2D layers-based desalination membranes. The availability of freshwater resources has decreased with increasing potable, agricultural, and industrial water demands around the globe for the past decades. This global challenge, along with the adversely accelerating climate change, has been calling for immediate solutions for meeting the demands. Apart from the conservation of existing water resources, increasing the freshwater supply is deemed as a more sustainable and direct solution. Since seawater is readily accessible and abundant, various thermal and physical desalination technologies have been utilised worldwide to produce drinking water by removing dissolved salts and other minerals from saline water. While multi-stage flash (MSF), multi-effect distillation (MED), and reverse osmosis (RO) are the most widely used desalination technologies, RO membrane is gaining market share at a faster pace. RO is a high-pressure membrane process that utilizes a semi-permeable

polymeric membrane to retain salts and low-molecular-weight solutes while allowing the passage of water via diffusion. Despite the operational simplicity, RO membranes suffer from several technical limitations including low conversion efficiency, membrane fouling, and energy demand to pressurize the feedwater. Accordingly, developing high-performance membrane materials with improved water recovery without compromising the solute rejection remains a compelling demand.

A potentially promising solution stands with engineering the structure of traditional materials into tailored porosity or exploring a new type of intrinsically porous nanomaterials-based membranes. Recently developed two-dimensional (2D) materials present unprecedented opportunities for water desalination and molecular sieving technologies owing to their structural uniqueness. 11-13 Their extremely small thickness (i.e., a few Å for 2D monolayers) projects superior water permeation rates benefiting from small diffusion lengths for molecular transports while their high mechanical flexibility renders advantages for membrane integrations. Moreover, 2D materials produced by chemical synthetic routes possess highdensity "intrinsic" structural imperfections (e.g., near atomic vacancies) whose physical dimensions are comparable to those of salt ions and water molecules. ¹⁴ In this endeavor, a variety of 2D materials have been explored including graphene, 15-18 dichalcogenides, 19-22 and graphene oxides, ²³⁻²⁶ along with substantive efforts for the structural engineering of traditional membrane materials such as zeolites, ²⁷⁻³⁰ and metal-organic frameworks ³¹⁻³⁴. Amongst them, 2D molybdenum disulfide (MoS₂) layers have recently gained significant interests for a wide range of water and environmental technologies. They not only share the intrinsic structural advantages inherent to other 2D materials but also present additional suitability such as excellent chemical stability and non-toxicity. 35-37 Furthermore, a variety of structural variations such as point defects, grain boundaries, and van der Waals (vdW) gaps have been ubiquitously observed in 2D MoS₂ layers grown via chemical vapor deposition (CVD) process.³⁸⁻⁴⁰ Such "intrinsically" present structural imperfections of very high concentration (e.g., areal density of ~10¹³ cm⁻² for sulfur vacancies⁴¹) offer 2D MoS₂ layers high promise as an efficient membrane material for water desalination applications. Despite the theoretically-projected promise of 2D MoS₂ layers, their experimental utilization for membrane-based desalination technologies has been rarely approached. A few experimental studies have explored MoS₂ laminar membranes of very large thickness (typically, ~1-10 μm) constituting mechanically stacked individual 2D flakes.¹⁴ These laminar 2D MoS₂ membranes of large thickness have exhibited significantly limited desalination efficiencies, failing in meriting the structural uniqueness and performance advantages projected with near atom-thick membranes. Scarcity of experimental studies on the 2D MoS₂ layers-based membranes of near atomic thickness for water desalination is attributed to technical difficulties associated with integrating 2D layers preserving their structural integrity. Particularly, it has been challenging to produce large-scale (> cm²) "continuous" 2D MoS₂ layers of ~1-10 nm thickness and their seamless integration onto porous substrates, hindering their applications to desalination technologies.

In the latter part of the dissertation, we investigate the experimental realization of high-efficiency water desalination enabled by ultra-thin 2D MoS₂ layers-based membranes. We demonstrate the integration of centimeter-scale CVD-grown 2D MoS₂ layers of only ~7 nm thickness onto porous polymeric substrates and studied their desalination performances. This new type of near atom thick-membranes exhibits an excellent combination of high water permeability approaching theoretically predicated values and superior ionic sieving performance for various seawater salts. Fundamental mechanisms responsible for the observed high desalination efficiency are also discussed, with a special focus on the roles of interfaces.

CHAPTER 2: LITERATURE REVIEW

2.1 Polymer-Derived Ceramics (PDCs)

2.1.1 Introduction to Polymer Derived Ceramics

Polymer derived ceramics (PDCs) are a novel class of amorphous materials synthesised through the controlled thermal decomposition of polymeric precursors. As compared to the conventional polycrystalline ceramics, PDCs possess unique complex microstructures that comprised of a silicon-based amorphous matrix phase and a nano-domained "free-carbon" phase that contains graphitic sp^2 carbon. Silicon-based PDCs have been obtained via the pyrolysis of organo-silicon polymers as a direct synthesis route as early as the 1960s. Pioneering work by Fritz and Raabe⁴² and by Yajima *et al.*⁴³ on the synthesis of SiC ceramic fibers from thermolysis of polycarbosilanes made prominent scientific progress towards application potential of PDCs.

The PDCs processing route has drawn growing attention as an additive-free method which also yields materials with fine-tuned chemical composition as well as well-defined nanostructures. He sophisticated microstructures of PDCs would experience significant transformations under very high temperatures with a characteristic persistence of the nanodomains, which gives rise to the exceptional resistance to crystallisation. Characterisation techniques for investigating PDCs microstructures include NMR, XRD, SAXS, FTIR, and Raman spectroscopy, which give integral information, as well as TEM, SEM and EELS that provide information about local properties at the nano-scale. In addition to experimental studies, there have also been many theoretical investigations aiming to explain the transformations of nano-clustered carbons during thermal decomposition as well as the crystallisation behaviour of PDCs.

The unique yet diverse electrical and thermo-mechanical properties of PDCs such as high-temperature semiconducting behaviour⁴⁶, anomalously high piezoresistivity⁴⁷, profound doping effects⁴⁸, as well as excellent resistance towards high-temperature decomposition and creep⁴⁹ have enabled PDCs to become promising candidates for potential applications in ceramic fibres⁵⁰, ceramic matrix composites⁵¹, energy storage⁵², porous components⁵³, and high-temperature micro-sensors⁵⁴.

Through the PDCs direct polymer-to-ceramic processing route, products such as ceramic fibres and layered composite materials have been made, which the conventional powder-based technology cannot feasibly yield due to the presence of additives. Traditional polymer-forming techniques such as polymer infiltration pyrolysis (PIP), injection molding, coating from solvent, extrusion, or resin transfer molding (RTM) can also be utilised to process pre-ceramic polymers, and the formed objects then undergo further conversions to ceramic components by appropriate heat-treatment. ²

The processing procedures of PDCs comprised of the following fundamental steps (as shown in Figure 1 as a basic flow-chart):

- (a). Chemical synthesis of starting ingredients which produces polymeric precursor;
- (b). Cross-linking of polymeric precursor gives infusible pre-ceramic network;
- (c). Pyrolysis of the pre-ceramic network under high temperature yields amorphous ceramics.

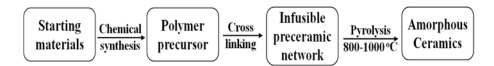


Figure 1 The processing procedure of PDCs.

2.1.2 Polymeric Precursors

As the starting materials for PDCs, pre-ceramic polymeric precursors have attracted extensive attentions, and many review articles ⁵⁵⁻⁵⁷ are dedicated to topics related to precursors synthesis. The importance of their key roles in heavily influencing the properties of PDCs stems from the fact that the chemical composition, the number of phases, the phase distribution and the microstructure of PDCs depend upon the molecular structures of polymeric precursors. Extensive research efforts have been devoted to the investigations of tailing various molecular structures to achieve optimum macroscopic chemical and physical properties of PDCs.²

Birot *et al.*⁵⁵ had proposed some key properties that effective polymeric precursors should possess in order to satisfy subsequent thermal decomposition processes:

- (a). Sufficiently high molecular weight in order to avoid volatilisation of low molecular components;
- (b). Suitable rheological properties (viscoelasticity) and solubility for the shaping process;
- (c). Latent reactivity (presence of functional groups) in order to gain thermosetting or curing properties;
- (d). Polymeric structures with cages or rings to reduce the volatilisation of the fragments due to backbone cleavage;
- (e). Low level of organic groups which lead to increased ceramic yield and decrease undesired/excessive "free-carbon" contents.

Since its inception, silicon-based polymeric precursors for PDCs have long drawn wide attention. Suitable organo-silicon PDCs precursors have a generalised molecular structure as illustrated in Figure 2. Design and tailoring of the polymeric precursors on a molecular level

mainly focuses on two aspects: a). the (X) group on the backbone, and b). the substituents groups attached to Si (i.e., R^1 and R^2). Different classes of the Si-based polymers contain different (X) group. Examples as shown in Figure 3 include: poly(organosilanes) with X=Si, poly(borosilanes) with X=B, poly(carbosilazanes) with X=C, and poly(organosilylcarbodiimides) with X=[N=C=N]. Functional side groups of R^1 and R^2 also play important roles in adjusting the solubility of the polymer, the electronic, optical, and rheological properties as well as the chemical and thermal stability of PDCs. Furthermore, the R^x groups determine the carbon content in the final PDCs products. 2

Figure 2 Generalized representation of the molecular structure of pre-ceramic organo-silicon PDCs precursors.²

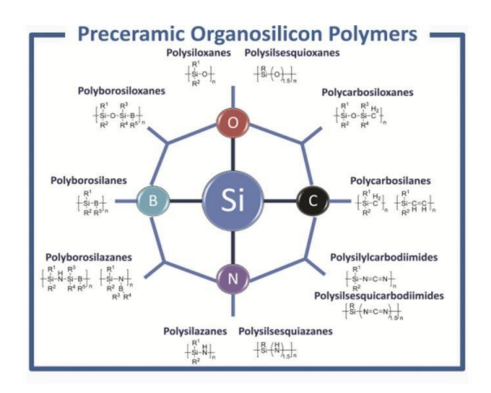


Figure 3 Main types of Si-based polymer precursors for PDCs.²

Besides from the above-illustrated Si-B, Si-C, Si-N and Si-O based polymer backbones, another promising elemental candidate is aluminium, because Al-containing materials (e.g., AlN and Al₂O₃) show desired properties such as high strength and excellent resistance to thermal shock and oxidation.⁵⁸ Research has shown that polysilazane-derived ceramic with a Si/Al/O/N system has a suppressed grain growth.⁵⁹ High-strength alkali-resistant Si/Al/C/O-fibers have been produced from polyaluminocarbosilane.⁶⁰

2.1.3 Processing of Pre-ceramic Polymers

2.1.3.1 Shaping

Since the PDCs precursors essentially remain polymeric materials during the shaping conditions/temperatures, extensive varieties of more cost-effective and industrial-friendly forming techniques can be applied than the traditional powder-/paste-based routes. Moreover,

this approach has important technological advantages in many aspects, for example: has no drying issues which has enabled bulk-fabrication; does not require long processing times for gelation and drying; no involvement of flammable solvents; can be processed in the molten state; avoids tool wear problems and brittle fracture upon finishing of the component.^{2,61}

The shaping techniques as reported in literature have been diverse: some are unique (e.g., formation of nanostructures directly by pyrolysis)⁶²; some explore polymer-processing routes (e.g., injection moulding)⁶³; while many are applicable to ceramic materials in general (i.e., casting/solidification, deformation, machining/material removal, joining, and solid free-forming).^{2,64}

2.1.3.2 Crosslinking

Since cross-linking is an essential part of the PDCs fabrication processes, a typical precursor falls into one the three categories: 1). a cross-linkable liquid; 2). a meltable and curable solid; 3). an unmeltable but soluble solid.² In addition, the shaped pre-ceramic polymers should be able to undergo a thermosetting transformation while keeping the shape unchanged during the subsequent polymer-to-ceramic conversion. Functional groups (e.g., the X group being –H or –OH) introduced to the pre-ceramic polymers can achieve a spontaneous thermosetting typically below 200 °C. It is also possible to further lower the cross-linking temperature via catalysing agents, which can also reduce oligomers evapouration hence fewer bubbles and higher ceramic yield.^{65,66} Successful laser curing has also been reported for rapid prototyping processes in fabricating complex-shaped components.⁶⁷

Various other curing methods may also be applied after-shaping. One commonly used method is oxidative curing, which would of course introduces undesired high level of oxide, thus results in ceramics with a lowered high-temperature stability. ^{68,69} Another means involve γ -radiation

or e-beam curing which can yield virtually oxygen-free ceramics, however the limited penetration depth means they are mostly suitable for producing fibres.⁷⁰ Incorporation of photosensitive functional groups onto the backbone of pre-ceramic polymers can produce UV-curable precursors, which has found desirable applications in fabricating MEMS and fibres.⁷¹⁻⁷⁴

The demands of bulk / batched productions of PDC has brought forth the introduction of various types of fillers to be incorporated with pre-ceramic polymers. Fillers serving different purposes have been employed, examples include: polymers, metals, ceramics in various formats (e.g., powders, platelets, nanotubes, and nanofibres).² Effects of fillers vary in broad ranges depending on the desired production outcomes. Inert fillers (e.g., SiC powders) help minimise the number of pore-sized defects; whereas functional fillers (e.g., MoSi₂ ⁷⁵ and FeSi have been used as electrical conductivity and magnetic functionality modifiers. Of the previously mentioned examples of fillers, metallic ones mainly contribute as to react away the gaseous byproducts during heat-treatment processes ⁷⁷; in contrast though, the polymeric fillers aim to produce micro-pores of desired sizes ⁷⁸.

2.1.3.3 Conversion from polymer precursor to ceramics

After obtaining shaped and cross-linked precursors, the final step towards PDCs products is polymer-to-ceramic transformation. Typical thermal decomposition (at approximately 1000 °C) of basic organo-silicon polymers $-[SiR^1R^2-X]_n$ —leads to amorphous SiC, $Si_xC_yO_z$, and $Si_xC_yN_z$ PDCs (as illustrated in Figure 4). The polymer-to-ceramic conversion process results in the decomposition or elimination of methyl, phenyl, vinyl groups as well as Si–H, Si–OH, or Si–NH $_x$ groups. There have been many conversion methods applied, both thermal and non-thermal, 2 yet amongst them, oven pyrolysis still remains the most extensively used coupled

with flowing inert gas continuously removing undesired decomposition residuals. One of the most routinely used conventional technique is thermal decompositions at relatively medium temperatures, followed by annealing stages at even higher temperatures. Another route explored is via photolithography, during which liquid phased mixture of pre-polymeric materials are exposed under ultra violet irradiations for a relative brief period of time. At this stage, the crosslinking processes are initiated and carried to complete, yielding pre-ceramics in solid phase with fewer residues. The resultant product then would undergo similar pyrolysis as well as annealing processes as mentioned above. The latter approach (the UV route) normally would result in fully-dense PDCs samples and shorter products turn-around times.

Various other newly-developed / implemented techniques have also been utilised to yield PDC components of complex shapes. One prominent example is 3-dimentional printing which have found applications in producing microfluidic channels via lithography ⁸⁰ as well as parts with ordered porosity based on a powdered route ⁸¹. Another promising technique for micro / nanofabrication of PDCs is focused ion beam (FIB) for micro electro mechanical systems (MEMS) applications as demonstrated by Tian *et al.*, which have been shown to yield PDCs products of more precise shape controls while retaining the superior properties of PDCs.⁸² Additive manufacturing techniques, with the help of 3-D printing and stereolithography, have also been explored to develop ceramics with greater geometrical flexibility while improving material strength and density which hold promise for potential applications in wider fields such as protective thermal shields and electronic device packaging.⁸³

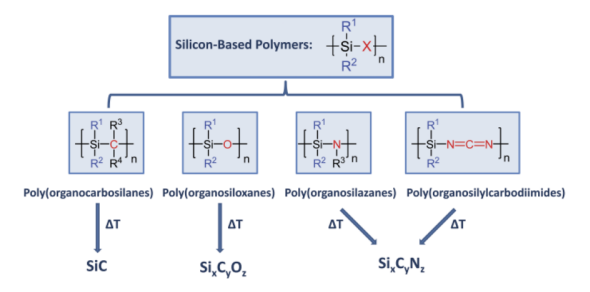


Figure 4 Typical thermal decomposition of basic organo-silicon polymers.²

Among the other polymer-to-ceramic conversion methods, chemical vapour deposition⁸⁴ and plasma spraying⁸⁵ have been used to produce ceramic coatings with the pre-ceramic polymers deposited directly on different substrates. Ceramic coatings has also been prepared via laser pyrolysis, which has the capability for selective writing.⁸⁶ In terms of non-thermal process, ion irradiation has been used to produce nanowires⁸⁷ as well as luminescent ceramic coatings/layers with high hardness.⁸⁸⁻⁹⁰

2.1.4 Microstructure of PDCs

PDCs possess sophisticated microstructures and would experience significant transformations under very high temperatures (>1000 $^{\circ}$ C) with a characteristic persistence of the nano-domains, which gives rise to the exceptional resistance to crystallisation.⁴⁵ At elevated temperatures, local crystallisation initiates in the amorphous bulk, followed by chemical bonding redistribution leading to phase separation, and finally results in nucleation and growth of nanocrystals. It has been reported that SiCN contains nano-domains of approximately 1–3 nm in

size, ⁹¹⁻⁹³ with the type, composition and shape of nano-domains identified via small-angle X-ray and neutron scattering (SAXS). ⁹⁴ A schematic illustration of the possible microstructure of Si-C-N based PDCs is given in Figure 5.

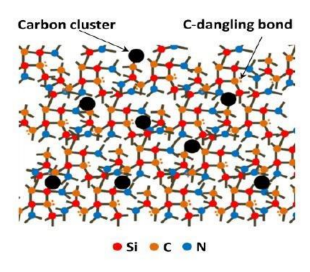


Figure 5 Schematic illustration of the possible microstructure of Si-C-N based PDCs.

For the amorphous-to-partially-crystalline transformation process, PDCs materials typically experience the following three main phases of microstructural changes: a). phase separation in the bulk of the material during annealing; b). graphitization process gives rise of the "free carbon" phase with the presence of carbon-dangling bonds; c). localised nano-crystals emerge under elevated temperatures.²

Characterisation techniques for investigating PDCs microstructures include NMR, XRD, SAXS, FTIR, and Raman spectroscopy which give integral information, as well as TEM, SEM and EELS that provide information about local properties at the nano-scale.² Various and extensive experimental characterisation efforts have been devoted to study the microstructure or microstructural evolutions of PDCs. Raman spectroscopy reveal that beside the possible presence of "free carbon", graphene sheets of relatively long-range order could exit. Raman

spectra also act as a powerful technique to conduct quantitative investigations on the evolutionary history / concentration changes of these "free carbons", which would significantly influence the electrical conduction mechanism (i.e., amorphous semiconducting behaviour for PDCs with low free carbon concentrations, whereas high free carbon concentrations yield a tunneling percolation mechanism⁹⁵). ⁹⁶

Free carbons are generally present in many PDC systems. Depending on temperatures, their distributions are different. At elevated temperatures (e.g., > 1000 °C), carbon takes the form of locally enriched regions of turbostratic graphite, whereas at lower temperatures, they normally show a homogeneous dispersion within the amorphous matrix. Although not thoroughly understood about the nature of free carbons, several reports have attributed PDCs oxidation / corrosion / crystallisation resistance to the presence of and its distribution characteristics of them. Transmission electron microscopy (TEM) have been extensively used to study the nature of the free carbons, which confirmed the presence of nano-sized carbons in turbostratic / graphitic forms.² TEM results also revealed how that the concentration of polymeric precursors as well as annealing histories could affect the ordering of carbons clusters. 97 Investigative efforts have devoted on the nano-domain-like distributions of Si-based carbon and carbon-rich clusters. One of the most valuable tools is small-angle X-ray scattering (SAXS), which proved the presence of short-range order. Together with the other auxiliary techniques such as Raman spectroscopy as well as X-ray diffraction (XRD), SAXS characterisations have confirmed to presence of nano-domains within polymer derived SiCN samples, with the increase in the nanodomain volume fraction lead to the increase in the inter-granular surface area thus the reactivity of the PDCs. 94 Experimental techniques, however, have limitations on the investigations of PDCs because of the amorphous nature of the materials. To compliment this, computational / theoretical modelling efforts have also emphasised on the microstructural changes in PDCs

systems. The microstructural evolutions of PDCs during processing / annealing as well as when under external influences (e.g., stress, temperature) can reveal both how to better tailor the properties of PDCs for targeted needs and how to understand the fundamental mechanisms of its characteristic behaviours (e.g., electronic, magnetic, optical, and chemical aspects).

As discussed above, the versatility of the potential applications of polymer derived ceramics stems from the possibility to tailor the backbone elements of the polymeric precursors (as well as a variety selections of fillers of many types), which ultimately comes from the less-sophisticated chemical synthesis processes.

Most commonly studied PDC categories are ternary ceramics as well as quaternary ones (i.e., polymeric backbones containing three or four types of main elements). Of all these PDCs, those derived from silicon (Si) containing polymeric precursors are the most promising and are the most extensively investigated.

Besides the foundational Si atom on the polymeric backbone, molecular structural differences also would influence the properties of the final PDCs products. Stoichiometric factors dictate there would be three groups attached to the Si atom (other than the Si-Si bonds themselves). These three attached groups can be classified into two main categories: organic groups of R¹ and R² as well as a functional group of X (as show below in Figure 6). The choice of different organic groups of R would mainly influence the rheology as well as solubility which would certainly affect the selection of processing techniques in subsequent steps.⁹⁸ In addition, the PDCs yield is also dependent on the amount of carbon atomic contents of the R organic groups, with higher yields correspond to increased carbons. Moreover, the functional X group is chiefly responsible for the microstructures of the resultant PDCs, where the presence of free carbons become a hindrance for diffusions thus lead to the increased resistance towards

crystallization. 99 A synergistic effect of the selection of the R and X group (as well as the Si/X ratios) will ultimately determine the properties of PDCs on the stability, electronic, and optical aspects. Mera et. al. 100 have reported nanoscale characterizations of PDC SiCN and SiCO in terms of microstructures and energetics. They found that the results are consistent with the nano-domained models as proposed by Saha et al. 101, which essentially comprised of a continuous yet inefficiently packed Si-based matrix phase, as well as entrapped within some carbon-rich clusters of varying degree of order. The illustrative depictions of Si-O-C as well as Si-C-N PDC systems are given in Figure 7 and Figure 8 respectively. The mixed bonds (Si-C and Si-O bonds) sandwiched between the Si-O-C and carbon-rich domains (as shown in Figure 7) are attributed to the high thermodynamic stability according to calorimetric results. Additionally, carbon contents in SiCN PDCs significantly affect the microstructure in terms of the nano-domain dimension and distributions. At lower carbon contents, polymer derived SiCN ceramics are constituted of Si₃N₄ and free carbon (mainly in sp²) nano-domains (which are inherently amorphous), with the inter-domain regions consist of mixed bonds (Si-N and Si-C bonds) which is likely stabilised due to the H atoms. Whereas for higher carbon contents. SiCN PDCs witness free carbon nano-domains of larger sizes, which are semi-continuously interconnected within the matrix (depicted below in Figure 8). One noteworthy feature of PDCs is that the annealing temperatures would cause further microstructural changes in terms of the loss H as well as lowered amount of mixed bonds (refer to Figure 9).

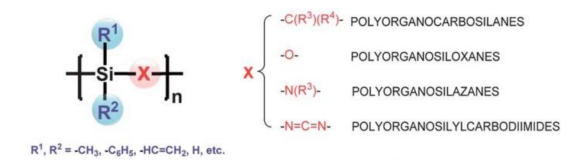


Figure 6 A generalized formula of Si-containing polymeric PDC precursors. 100

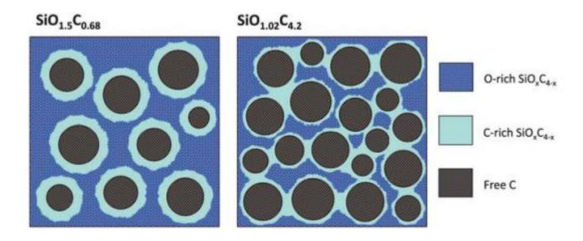


Figure 7 Nano-scale structural model of low carbon (left) and high carbon (right) containing SiOC ceramics at $1100~^{\circ}$ C. 100

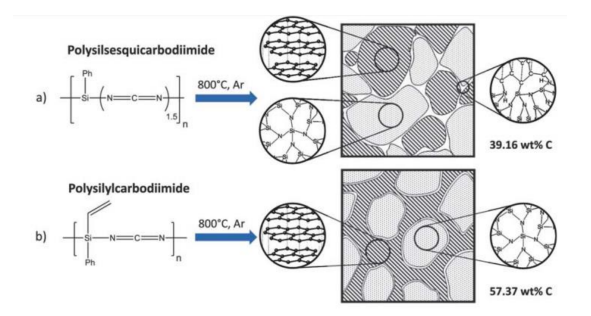


Figure 8 Carbon-rich precursor system comprised of Si_3N_4 clusters embedded within a continuous amorphous / turbostratic carbon matrix. 100

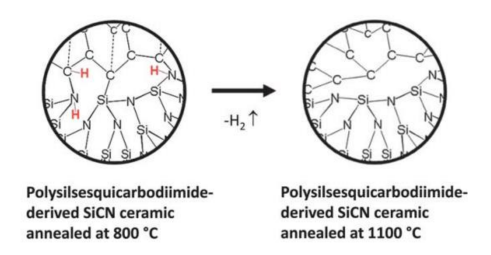


Figure 9 Mixed bonding in the interfacial region between Si₃N₄ and carbon nano-domains in a samples synthesized at 800 °C and (right) loss of hydrogen and of mixed bonding post 1100 °C pyrolysis.¹⁰⁰

Microstructural changes in PDCs under external stress / strains have also been reported, both for SiOC ¹⁰² as well as SiCN ⁴⁷. Both teams observed piezoresistive effect for respective PDCs samples with the latter team ascribed this phenomenon to the unique self-assembled nano-

domain microstructures discussed above following a tunneling-percolation model. Whilst the former team studying polymer derived SiOC discovered similar mechanism due to percolation effects owing to the segregated (electrically conductive) carbon clusters within the Si-based matrix. Specifically in terms of the origin of piezoresistivity, a tunneling-percolation network within the Si-based matrix stem from the conducting graphene sheets, with the tunneling process (which PDCs resistivity is dependent of) become ultra-sensitive to the distance between the graphene sheets. Such model therefore connects the piezoresistivity to the distance changes between graphene sheets due to external pressure.

In addition to the influence of external pressure on microstructural changes in PDCs, temperature could also influence the microstructures of PDCs especially causing changes in free carbons thus further determine the electrical properties. The pyrolysis temperatures could cause the ratio of sp^2/sp^3 carbon to increase thus lead to a rise in electrical conductivity owing to various mechanisms: a band-tail hopping conduction mechanism for SiAlCN 104 , a caused by the conductivity increase in the carbon phase for SiBCN 105 , as well as C-O bonds re-arrangement in the free carbon phase (for SiOCN) 106 .

2.1.5 Properties of PDCs

2.1.5.1 Thermal Properties

Riedel *et al.*⁴⁹ have reported a polymer-derived SiBCN ceramic that can withstand thermal degradation up to ~ 2000 °C, much higher than that of SiCN and Si₃N₄. The composition of SiBCN annealed at 2000 °C almost remains the same as the as-synthesised specimen, indicating its excellent high-temperature stability.

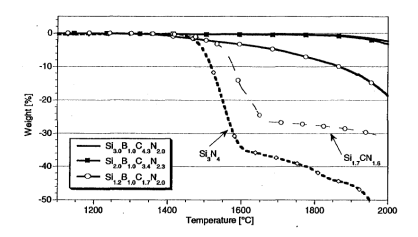


Figure 10 Thermal gravimetric analysis of polymer-derived SiCN, SiBCN and commercially available Si3N4.⁴⁹

As shown in Figure 11 of the XRD results, SiBCN remains amorphous up to 1700 °C, and when annealing temperature reaches 2100 °C, β -SiC is found as the only crystalline phase, showing excellent resistance to crystallisation.⁴⁹

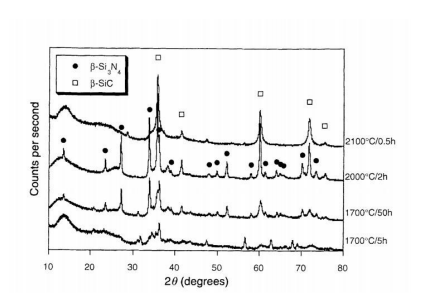


Figure 11 The XRD results of SiBCN annealed at different temperatures.⁴⁹

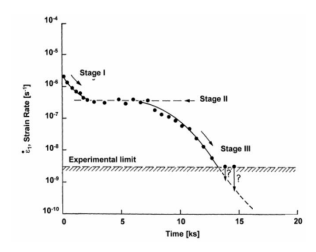


Figure 12 Typical creep observed in polymer-derived SiCN and SiBCN materials. 108

Polymer-derived SiAlCN ceramics have been reported to exhibit superior oxidation and hot-corrosion resistance than that of SiCN (as shown in Figure 13). 109-111 Comparison of SiAlCN and SiCN surfaces underwent hot-corrosion in NaCl saturated solution shows virtually no cracks on SiAlCN (Figure 14). Due to the excellent oxidation/corrosion resistance of SiAlCN ceramics, they are promising candidates for high-temperature and harsh-environment applications.

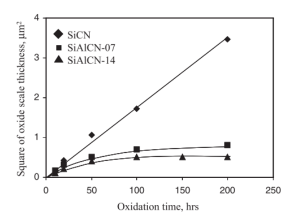


Figure 13 A plot of the square of the oxide scale thickness as a function of annealing time for both SiCN and SiAlCN at 1200°C in dry air.⁵⁸

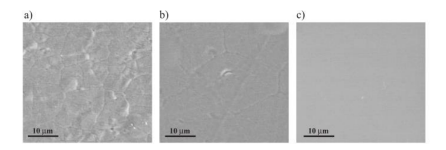


Figure 14 SEM micrographs of the surfaces of (a) SiCN, (b) SiAlCN-07 and (c) SiAlCN-14. The specimens first immersed in de-ionized water saturated with NaCl at 100°C, and then annealed at 1200°C for 50h in air.⁵⁸

2.1.5.2 Mechanical Properties

Apart from PDCs fibres, the Si–C–O, and the Si–C–N systems have attracted detailed investigations in terms of bulk mechanical properties of PDCs. For both systems, mechanical properties have been measured at different pyrolysis temperatures, with the summarised reported results presented below in Table 1.² In addition to the excellent room-temperature mechanical properties, it is expected that such superior mechanical behaviours can be retained at high temperatures due to the absence of grain boundary phases.

Table 1 Summary of the Mechanical and Thermal Properties Measured on Bulk Samples of the Si–C–N and Si–C–O Systems. ²

System	Bulk density , , , , , , , , , , , , , , , , , , ,	Elastic modulus , , E (GPa)	Vickers hardness , HV (GPa)	Fractur e strength σ_F (MPa)	Poisson's ratio,	Fracture toughness, K _{IC} (MPa · m ^{1/2})	Thermal expansion , α (×10 ⁶ °C ⁻¹)
SiCN (Ceraset)	1.85- 2.16	82-140	8.3-11.3	<1100	0.21- 0.24	0.56-1.3	
SiCN (NVP200)	2.32	110-130	11-15	<118	_		_
SiCN (VT50)	1.95	105	6.1	88-146		2.1	3.08-3.96
SiCN (HVNG/HPS)	2.6	109-118	7.9-12.8	80-235			
Si(O)C (AHPCS)	2.3-2.9		9.13			1.4-1.67	
SiCO (sol- gel)	1.7- 2.28	57-113	4.7-9.3	70-900		0.57-0.77	3.21
SiOC (MK)	2.23	101	6.4		0.11		
SiOC (SR350)	2.35	97.9	8.5	133- 612		1.8	3.14

In general, the elastic moduli of PDCs systems of S-C-N and S-C-O is lower than that of the crystalline silicon carbides or silicon nitrides, due to the less densely packed systems of networks, which also lead to low densities.² For the conventional thermal pyrolysis routes of producing PDCs, higher annealing temperatures typically result in increased elastic modulus

as well as density because of the loss of H atoms in C-H bonds leading to more closely connected networks. Similar trends are observed for the hardness values of PDCs, where higher pyrolysis temperatures normally yield increased hardness, since more carbon atoms migrated into the silica networks, thus impose more restrictions on dislocations. Extraordinary creep resistance is another interesting feature of (Si-based) PDCs, which have typical viscosities of three orders of magnitudes better than vitreous silica. 108

2.1.5.3 Electrical Properties

Electronic behaviours of amorphous PDCs have gained extensive attentions because: a). they provide a unique model system for studying the structure-properties-performance relationships of complex amorphous covalent materials, and b). they could find the promising applications in harsh-environment sensing as well as MEMS.^{54,113}

2.1.5.3.1 DC Conductivity

Haluschka *et al.*¹⁰³ have shown that the electrical properties of amorphous Si–C–N ceramics are in the range between the electrical properties of semi-conducting SiC and that of insulating Si₃N₄. Their results revealed that the DC conductivity is influenced by the conditions of the final heat treatment, namely the annealing time and temperature. Haluschka *et al.* identified three distinctive regimes: a) due to the loss of residual hydrogen and an increased sp^2 -/ sp^3 -ratio of the carbon atoms, the conductivity of the amorphous SiCN matrix would experience a rise up to three orders of magnitude (up to 1300 °C); b) due to the formation of SiC and the loss of nitrogen of the remaining amorphous a-SiC(N) matrix, the material had a pronounced conductivity increase (between 1300 and 1600 °C); and c) lastly the nano-crystalline SiC particles were responsible for the conductivity increase by formation of percolation paths throughout the sample (above 1600 °C).

2.1.5.3.2 AC-conductivity

Studies on AC (alternative current) behaviour (as shown below in Figure 15) of a PDC material (SiCN) show that at low frequencies the AC conductivity remains stable (attributed mainly to the long-range transport of charge carriers through the matrix-to-free-carbon path), while at high-frequency regime it rises with frequency (proposed to be caused by interfacial polarisation due to charge carrier relaxation within the free-carbon phase).¹¹⁴

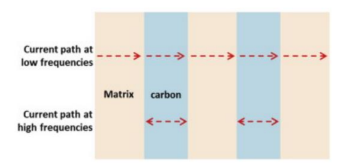


Figure 15 Schematic showing the conduction mechanisms in the a-SiCN at low and high frequency regions.¹¹⁴

2.1.5.4 Impedance

Complex impedance behaviors of polymer-derived amorphous SiCO and SiAlCO are reported. 115,116 Equivalent circuit models have been applied and showed fitting correspondence with the measured data. The conduction of current in both materials is found to pass through the matrix and free carbon phases in series, with the conductivity of both phase increases with increasing synthesis temperature, whereas the matrix has a significantly shorter relaxation time than that of the free carbon. The relaxation process of polymer-derived SiAlCO is found to move to higher frequency with increasing pyrolysis temperature.

Despite the investigative efforts made as discussed above, little attention has been focused on the interfacial areas sandwiched between the matrix and carbon phases. The role and effect of the interfaces deserve more in-depth investigations as the electrical behaviors may experience great influence from the interfaces between two phases with vastly different conductivities. ¹¹⁷ However, this has started to attract research efforts with some recent reports investigating mainly Si-C-N PDCs systems.

Strong interfacial charge polarisation has been reported to present, leading to giant relative permittivity. ¹¹⁸ The interfacial polarisation underwent relaxation processes (from -50 to 300 °C), following a 3D random hopping mechanism, depend on the capacitance and resistance of the interfacial areas. ¹¹⁹ Employing complex impedance spectroscopy with superimposed direct current (dc) bias, Li *et al.* have applied a double Schottky barrier model to help determine the room temperature interfacial characteristics. The authors have revealed that the capacitance and resistance appeared to be independent of dc bias, whereas that of the interface is suppressed with increased dc bias voltage, with such effect attributed to bias influence on potential barrier height and barrier region width. ¹²⁰

Due to intense research interests on PDC as a promising candidate particularly for high-temperature sensing applications, it would provide more in-depth revelations to further study the interracial behaviours under elevated temperatures, and impedance spectroscopy with superimposed dc current bias on ac current remains a powerful pathway towards this aim.

2.1.5.5 Contacts between Metallic-Conducting Phase and Semi-Conducting Phase

Due to the bi-phasic nature of most Si-based PDCs, the interface sandwiched between the two phase becomes an accumulation venue for charge carriers when PDCs are under external AC current. The interface is therefore a highly conductive region, which may be treated as

possessing metallic-conducting characteristics. Furthermore, comparative to the highly-conductive interfacial phase, the free carbon phase as well as the ceramics matrix phase have semi-conducting behaviours. Therefore, the contact between the (highly-conductive) interface phase and the semi-conducting phase (either the free carbon phase or the ceramics matrix phase) have similar electrical conductive characteristics of the metal-semiconductor contacts. Here in the following contents, we will review the fundamental theories of the Schottky contact, which shows the non-linear, rectifying current voltage characteristics. ¹²¹

For the energy band structures of either the interfaces or the (n-type) semi-conducting phase, their respective work functions are inherently different. With the work function of the interfaces being the minimum energy required to eject one electron (i.e., interfacial Fermi energy of $E_{F_{Inter}}$), whilst that of the semi-conducting phase has a work function of E_{F_C} (energy needed to "kick" an electron from the Fermi level at the conduction band). For discussion convenience purposes, we consider the contact between an n-type semi-conducting phase and the highly-conductive interfacial phase so that the Fermi level of the former phase is larger than that of the latter (i.e., $E_{F_{Inter}} < E_{F_{C}}$). Because of such energy level difference, when the interfaces phase makes contact with the semi-conducting phase (i.e., when the bi-phasic PDCs is placed under AC electric field forming the interfacial phase filled with accumulated charge carriers), the energy levels tend to align to reach thermodynamic equilibrium via the migration of charge carriers from the (high-energy level) conduction band to the (low energy level) interfacial regions. As the electrons depart from the semi-conducting phase, they also leave behind certain donor atoms with positive charges. Such process creates a region depleted of electrons (i.e., depletion zone). When thermodynamic equilibrium is achieved after contact has been fully established, the Fermi energy levels reach an aligned value in addition to the

formation of the depletion region. Therefore, the process results in a curved energy band in the semi-conducting phase. Such curved energy band acts as a potential barrier, impeding the flow of electrons from the semi-conducting phase to the interfaces.

Under complete thermodynamic equilibrium and without any external electric field, electrical neutrality dictates that the current flow from the semi-conducting phase to the interfaces is exactly matched by the flow of the reverse direction. However, when the AC electric field is biased due to a DC voltage, the current flows of the two directions becomes different. Under a forward DC bias (*V*), the potential barrier (at the semi-conductor side) is lowered (by a level of *V*). This eases the restrictions of electron migrations and therefore increased electron flow from the semi-conducting phase to the interfaces. Furthermore, since the potential barrier from the interfaces side remains the same, the level of electrons crossed into the semi-conducting phase is still unchanged. Such imbalance of the bi-direction electron flows effectively creates current from the interfaces to the semi-conducting phase. On the other hand, when under a reverse bias (of *V*), the potential barrier witnesses a rise (of *V*). Similar to the above discussions, the electron flows from the semi-conducting phase are reduced because of the heightened potential barrier. Since the interface-to-semiconductor current still remains unchanged, the resultant net current flows from the interfaces to the semi-conducting phase.

2.1.5.6 Piezo-resistivity

Some amorphous PDCs have been reported to demonstrate significant piezoresistivity, with a polymer-derived SiCN showed piezoresistive coefficients of an unprecedented 4000 ⁴⁷ following a tunneling-percolation model based on Saha *et al.*¹⁰¹ who proposed the formation of conductive graphene sheets within the framework of PDCs nano-domains. Similarly, the tunneling-percolation mechanism has been attributed to a reported high strain sensitivity of 145

of a PDC SiCO owing to the presence of a percolating network of electrical conductive free carbon. ¹⁰² Interestingly, the electrical resistance of SiCO rises with increased compressive stress, contrary to that of SiCN.

2.1.6 Magnetic Properties

Polymer derived ceramics containing ferrous elements have attracted research efforts investigating its magnetic behaviours. Since PDCs themselves are not inherently magnetic, one common approach to impart such properties is to incorporate PDCs with other ferrous materials or particles to make essentially PDCs composites. One less sophisticated method is to simply disperse Fe₃O₄ powders into liquid-phased polymeric precursors before undergoing a reduction process in graphite, yielding a high M_s of around 57 emu·g⁻¹. 122 Iron silicide fillers have also been incorporate into preceramics producing PDCs with cubic Fe₃Si particles embedded the matrix, showing an M_s of up to 17 emu·g⁻¹. A more fundamental approach to synthesise magnetic PDCs is to modify the design at the molecular / polymeric backbone level, which is to add metallic iron atoms (e.g., via ferrocene).² Sun et al. have obtained phosphoruscontaining PDCs with good magnetisation (M_s up to 50 emu·g⁻¹) exhibiting very low remanence and coercivity. 123 Higher (more than doubled magnetisation of 118 emu·g-1) have been detected from Cobalt-containing PDCs which also retained similarly low remanence and coercivity. 124 In addition, the observed magnetic behaviours also agree with the evidence of the existence of domain-like regimes within PDCs matrix. And such functionalised PDCs with tunable electromagnetic properties may find promising applications as regenerable absorbers or regenerable diesel particulate traps.⁷⁶

2.1.7 Optical Properties

Polymer derived ceramics did not initially attract much attention upon investigating its optical properties since most PDCs are not optically transparent. The blackish appearance of many PDCs is chiefly believed to have cause by the presence of sp^2 carbons forming light-absorbing graphitic thin layers.² However, research efforts have yielded largely transparent SiCO PDC discs (due to the lowered presence of sp^2 carbons). ¹²⁵ These transparent PDCs show a luminescence band centering around 500 nm which the authors ascribed still to the sp^2 carbons. Produce via a sol-gel route, SiCO PDCs doped with Eu²⁺ proved to be a luminescent glass, with an emission band exhibit a peak in 450–480 nm. 126 With the aim of producing high-temperature stable materials for white light applications, an Italian team has pyrolyzed sol-gel derived SiOC PDCs films showing visible luminescence. 127 They show that different pyrolysis temperatures yielded luminescence of varying colours: bluish emitting from low-temperature annealed samples, whereas their high-temperature analogues exhibit green-yellow. Besides SiOC PDCs, polymer derived SiCN have also been reported with regard to its photo-luminescence behaviours and mechanisms. 128 In contrast to their SiCO counterparts, these N-containing PDCs show prominent luminescence at much lowered annealing temperatures (500 – 600 °C), which the authors attributed to the microstructural re-arrangement as well as the presence of sp^2 carbon along with the formation of radicalized species. In addition, SiOCN PDCs have also been shown to exhibit intense photo-luminescence in the range of 500 - 800 nm. ¹²⁹ The author also employed the previously-discussed nano-domain model to explain the luminescence phenomena, which they ascribed to the mixed bonds of Si-C-N-O tetrahedral.

2.1.8 Chemical Properties

Chemical stability of PDCs is the reason they attract research interest to develop and study in the first place, with the additional characteristics of superior oxidation resistance properties.² There have been extensive research reports with regard to the chemical durability of PDCs. Polymer derived SiOC ceramics have been put to the test of highly corrosive acidic and basic solutions to examine its chemical stability.¹³⁰ Compared to its silica counterparts, SiOC PDCs show much greater endurance because of the presence of Si-C bonds, and the cross-linked carbons as well as the highly disordered composition, which jointly retard the transport of nucleophilic species.

The oxidation behaviors of PDCs have been widely studied (with focus on systems of SiC, SiCO, SiCN, and SiBCN). Here, the pyrolysis processes play a vital role, since the elimination of H atoms is key to achieve parabolic oxidation rates.² Most systems investigated reveal a dense, continuous (defect-free) oxide layer above the ceramic interfacial areas. Efforts have also been devoted to the effects of additional elements incorporated with Si-C-N PDC systems (e.g., B, Al, as well as Zr), ^{109,131,132} relying on the blockage effect of these added atoms towards a dropped O₂ diffusion rates.

2.1.9 Applications of Polymer Derived Ceramics

As the above-discussed superior properties of polymer derived ceramics indicate, they could find numerous potential applications in various technical fields which this section aims to review and correlate the structure, processing and properties relationships.

The fundamental mechanism of application / properties tailoring lies with the versatile capability of designing the microstructures of the pre-ceramic polymers at a molecular level.

When combined with various types of fillers as previously discussed (e.g., metallic and polymeric), the properties of PDCs can be tuned to meet or more often exceed conventional ceramics in terms of mechanical strength, electrical / dielectric behaviours, and oxidation / corrosion resistance. It is therefore promising both in terms of fundamental research as well practical applications to study the properties of PDCs more in-depth.

2.1.9.1 PDCs Protective Coatings

Since the processing of polymer derived ceramics is based on liquid phased polymeric precursors, coatings / thin membranes could be produced while retaining the exceptional wear / oxidation / corrosion resistance capabilities. SiCO coatings have been applied on to fibres acting as flame / temperature shields. SiCN has also been spray coated on conventional substrates against oxidising environments and mechanical wears. Similar techniques have produced porous SiCN membranes aimed for gas (e.g., hydrogens) separation under high temperatures.

2.1.9.2 Micro-electro-mechanical Systems (MEMS)

The successful development of micro- or nano- electro mechanical systems relies on the interconnectivity of different components of various sizes. This again depends on the capability of
producing parts with complex shapes / dimensions. Conventional methods, besides the lack of
high accuracy, are limited in terms of this aspect, since molds of pre-determined corresponding
shapes would need to be manufactured first, which is also less cost-effective. The (high
temperature as well as harsh environment) mechanical strength of polymer derived ceramics
makes them a very promising candidate for MEMS components with desired features can be
as small as less than one micron achievable via lithographic techniques. Typical examples
of polymer derived ceramics MEMS include: electrostatic actuators (which could also find use

in high power laser systems)⁵⁴, photonic crystals¹³⁷, as well as embossed micro-fluidic channels¹³⁸.

2.1.9.3 PDCs Fibres

The fibres, which are composite in nature, made from polymer derived ceramics combines both the strength of carbon fibres and oxidation / corrosion resistance of ceramics. These PDC fibres have found wide applications as ceramics reinforcements and have attracted extensive further research interests for performance optimisations. Till today, three generations of PDC (mainly non-oxide) fibres have been developed, with the latest ones have already been successfully commercialised possessing exceptional thermal stability as well as very high Young's modulus. Flores *et al.* have reported polymer derived SiCN fibres which can retain a Young's modulus of over 300 GPa under temperatures as high as 1400 degrees Celsius. ¹³⁹

2.1.9.4 Polymer Derived Composites

Though polymer derived ceramics themselves already possess composites-like properties, tremendous interests and efforts have still been devoted to the further development of PDC-based composites combining the superior properties of many other newly-discovered nano materials. Carbon nanotubes / PDC composites have been reported with the modulus, hardness, as well as damage resistance of the reinforced composite product greatly improved even with just a small amount of multi-walled carbon nanotubes incorporated.⁵¹

2.1.9.5 Sensing Materials / Devices based on Polymer Derived Ceramics

As discussed in previous sections, one of the most unique aspects of PDCs are that their microstructures could change with external parameters (e.g., temperature, stress, and strain),

which lead to detectable changes in intrinsic bulk properties (such as: thermal coefficient, electrical conductivity, as well as relative permittivity).

Heat flux sensor for gas turbine environment under high temperatures have been reported based on polymer derived SiCN via lithography microfabrication techniques. ¹⁴⁰ Temperature sensors have also been variously reported take advantage of the "tunneling percolation" effect due to microstructural changes. One example demonstrated a wireless PDC (SiCN) sensor based on cavity radio frequency resonator. ¹⁴¹ Another showed a SiAlCN-based high temperature sensor with retained good accuracy even after many repetitive cycles. ¹⁴²

Pressure sensor, with desired shapes and very high hardness (23 GPa), for high temperatures have also been reported employing the drag effect. Gas sensing capabilities of PDCs have also been investigated, with hydrogen sensors being explored owing to the semiconducting behavior of polymer derived SiCN which were reported to show good performance up to 785 K. 444

2.2 Polymer-Derived SiAlCN Ceramics

Due to excellent oxidation and corrosion resistance¹¹¹, anomalously high piezoresistivity⁴⁷ as well as its capability for temperature measurement in harsh environment¹⁴², SiAlCN has become a novel promising candidate of high-temperature multi-functional material. It is particularly attractive to apply and integrate the PDCs into wireless sensing systems and embedded within the interior regions of moving parts of turbine engines, highly irradiative surfaces of gasifiers / reactors to detect the real-time temperatures and pressures to gain reliable and accurate feedbacks for system operation optimisations. Its excellent mechanical strength

means it could also find potential future applications in civil engineering field for structural health monitoring (for example: high-rise buildings or long-span bridges).

Polymer-derived SiAlCN ceramics has been found to have more superior oxidation resistance than other silicon-based materials, with the oxidation rate for the SiAlCN is about an order of magnitude lower than that for pure silicon carbide/nitride.^{58,110} It was also observed that SiAlCN exhibits excellent water-vapour corrosion resistance under 1100 ℃, which was attributed to the formation of aluminum-doped cristobalite oxide scale.¹¹¹

Electric properties characterisations of PDC SiAlCN show a smooth increase of DC conductivity with elevated sintering temperatures.³ The dielectric measurements revealed that the dielectric constant and dielectric loss rise with increasing temperature.³

Temperature sensor design based on polymer-derived SiAlCN for high-temperature applications has been reported. 142 The sensor resistance showed a monotonous change with ambient temperature (up to 830 °C), and has demonstrated excellent accuracy repeatability to both uni-directional and bi-directional temperature variations.

2.3 Desalination Membranes

The imminent risks of global water shortages pose as threats not only to those directly impacted due to the lack of clean water for everyday usage, industrial waste processing and treatment, as well as agricultural irrigations, but also cause collateral damages to the society since the imbalance of such basic human needs of accessing clean water would induce social unrest and conflicts. ⁶

As a readily accessible and abundant resource, seawater has the promising potential of greatly reducing the risks of the fresh water shortage. However, as a complex mixture of various types

of salt, water, and dissolved solvents, seawater is not directly potable or useful for industrial or agricultural demands. For decades, researchers and engineers have been exploring materials and techniques for removing the mineral components within seawater in an efficient and costeffective manner. Durable and selective-filtering membranes have been the dominant candidates for such applications, with the polymeric membranes still account for the majority of the commercially available products. ⁸ With respect to the most widely-adopted desalination techniques, reverse-osmosis (RO) have been the dominant method for producing large amounts of fresh water, with multistage flash (MSF), multi-effect distillation (MED) are also attracting increasing attentions. RO is a high-pressure membrane process that utilises a semipermeable polymeric membrane to retain salts and low-molecular-weight solutes while allowing the passage of water via diffusion. However, a major drawback of RO remains, which is its high energy demands due to the pressurisation process, which in turn contributes to the release of greenhouse gases from fueling burning. The polymeric membranes developed decades ago have underwent significant modifications in terms of increasing desalination performances for example new polymerisation routes, developing thin-film composite membranes, as well as post-synthesis surface modifications to obtain membranes with high permeability, salt rejection rates, fouling-resistant capability, and chlorine-tolerance. In addition, future challenges for desalination technique also come from integrating renewable energy source, as well as stringent water standard and brine management.9

In addition to the polymeric-based membranes, there have been many recently developed materials for desalination applications, although not fully capable of large-scale industrial usage. For example, ceramics-based membranes have been used in harsh environmental situations (e.g., high-temperature, highly-radioactive feeds, and highly-reactive components) where polymeric membranes could not endure. ¹⁴⁵ Furthermore, mixed matrix membranes (e.g.,

incorporating zeolite nano-particles) have been developed with smoother, more hydrophilic surface exhibiting improvement in salt rejection rates. Another category of novel desalination membrane involves the incorporation of carbon nanotubes (CNTs) which have been excellent water transport. However, although CNTs-integrated membranes show great promise in flux enhancement, efficient synthesis methods to produce membranes with well-aligned CNTs, as well as the development of surface modifications for more efficient salt rejection remain major obstacles for future applications. 147

Nano-structured RO membranes offer inherently high permeability as well as salt ions rejection, while the development of nanotechnologies could bring forth multitudes of innovative progresses towards high-performing desalination membranes. However, practical issues of cost-effectiveness, scaling-up, and potential health risks still demand extensive investigations.

2.4 Two-Dimensional Materials for Desalination Applications

As a promising alternative to the conventional polymeric desalination membranes, a viable potential solution stands with engineering the structure of traditional materials into tailored porosity or exploring a new type of intrinsically porous nanomaterials-based membranes. Recently developed two-dimensional (2D) materials present unprecedented opportunities for water desalination and molecular sieving technologies owing to their structural uniqueness. ¹¹⁻¹³ Their extremely small thickness (i.e., a few Å for 2D monolayers) projects superior water permeation rates benefiting from small diffusion lengths for molecular transports while their high mechanical flexibility renders advantages for membrane integrations. Moreover, 2D materials produced by chemical synthetic routes possess high-density "intrinsic" structural imperfections (e.g., near atomic vacancies) whose physical dimensions are comparable to those

of salt ions and water molecules.¹⁴ In this endeavor, a variety of 2D materials have been explored including graphene,¹⁵⁻¹⁸ dichalcogenides,¹⁹⁻²² and graphene oxides,²³⁻²⁶ along with substantive efforts for the structural engineering of traditional membrane materials such as zeolites,²⁷⁻³⁰ and metal-organic frameworks ³¹⁻³⁴.

Although for graphene in its pristine form is impervious to water molecules, techniques to generate nanopores on graphene sheets have been developed, and the nanoporous graphene membranes have shown great potential in desalination performances, with some investigations have demonstrated their capability of withstanding high pressures associated with RO. For example, Suk *et al.* have shown that graphene-based membranes show water permeation rates of 2–3 orders of magnitude higher compared to conventional RO membranes, with salt rejections approaching 100%. However, mature techniques for producing large-area graphene with well-defined distribution of nanopores are still absent at this stage of research. In addition to graphene-based membranes, graphene-oxide-based membranes offer many desiring features for potential water treatment technologies including distinct microstructural design, hydrophilicity, excellent mechanical strength, as well as good anti-fouling performances.

In addition to graphene and GOs based desalination membranes, many other types of 2D nanoporous membranes have been investigated for their potential for filtration applications owing to their ultra-thinness as well as inherent nano-sized pores for impeding salt ion translocations. For example, zeolites are composed of three-dimensionally interconnected nanopores of 0.3-0.8 nm, and have been shown to be able to effectively reject salt ions, while retaining a water permeability of 40 L/(cm² day MPa), similar to that of graphene. ¹⁴⁹ Moreover, covalent organic frameworks (COFs) consisting of tunable and precisely controlled nanopores

have been proposed as a promising candidate for filtration ultrathin membranes. For instance, Triazine framework nanoporous membranes have demonstrated a water permeation of 24.6 L/(cm² day MPa), and a salt rejections rate of nearly 100%. 150 Furthermore, owing to their high flexibility, mechanical strength, and hydrophilicity, MXenes with controllable thickness can also yield suitable membranes for separation applications with ultra-fast water flux. Ti₃C₂T_x MXene membranes have been shown to have a water flux of 37.4 L/(m² h bar), due to the hydrophilic nature of the material and the presence of intra-layer water molecules. ¹⁵¹ 2D materials of mono-to-few layers (e.g., h-BN, phosphorene, and MoS₂) have attracted extensive investigations towards their potentials for ionic sieving. A prominent example is 2D MoS₂ which have been fabricated for nanoporous membranes for desalination applications, in addition to many theoretical projections of their performances. MoS₂ also demonstrates excellent mechanical properties (e.g., MoS₂ nanoporous 1 nm-thick membrane possesses a Young's modulus of 270 GPa), as well as good flexibility. 2D molybdenum disulfide (MoS₂) layers have recently gained significant interests for a wide range of water and environmental technologies. They not only share the intrinsic structural advantages inherent to other 2D materials but also present additional suitability such as excellent chemical stability and nontoxicity.35-37 Furthermore, a variety of structural variations such as point defects, grain boundaries, and van der Waals (vdW) gaps have been ubiquitously observed in 2D MoS₂ layers grown via chemical vapour deposition (CVD) process. 38-40 Such "intrinsically" present structural imperfections of very high concentration (e.g., areal density of ~10¹³ cm⁻² for sulphur vacancies⁴¹) offer 2D MoS₂ layers high promise as an efficient membrane material for water desalination applications. In fact, several studies based on theoretical calculations and molecular dynamics simulations have predicted superior desalination performances of monolayer 2D MoS₂. ^{14,19,41,152,153} Heiranian et al. have predicted that 2D MoS₂ monolayers of

enriched defective sites can exhibit ~2-5 orders-of-magnitude higher water fluxes than traditional filtration membranes, which is even 70% greater than the performance of graphene nanopores. 154 Köhler et al. have projected that the ion rejection capability of 2D MoS₂ membranes is superior to graphene under a wide range of test conditions including pressure, nanopore size, and cation valences. ¹⁵² Li *et al.* have simulated the desalination performance of 2D MoS₂ layers, demonstrating high water transport rate (355.3 L m⁻² h⁻¹ bar⁻¹) and excellent salt rejection capability. 155 Despite the theoretically-projected promise of 2D MoS₂ layers, their experimental utilisation for membrane-based desalination technologies has been rarely approached. A few experimental studies have explored MoS₂ laminar membranes of very large thickness (typically, ~1-10 µm) constituting mechanically stacked individual 2D flakes. 14 Sun et al. studied ~1.7 µm thick MoS₂ membrane films prepared via chemical exfoliation and subsequent filtration of aqueous dispersion. 156 Hirunpinyopas et al. demonstrated chemically functionalised ~5 µm thick laminar membranes composed of exfoliated 2D MoS₂ flakes employing the similar filtration process.¹⁵⁷ These laminar 2D MoS₂ membranes of large thickness have exhibited significantly limited desalination efficiencies, failing in meriting the structural uniqueness and performance advantages projected with near atom-thick membranes. For example, their water permeation rates were experimentally identified to be in a range of ~245-275 L m⁻² h⁻¹ bar⁻¹, 156,157</sup> significantly falling short of theoretical values predicted by simulation studies. 154 Moreover, ion rejection rates of mere ~15-30% were reported for pristine laminar 2D MoS₂ membranes of ~1-5 µm thickness. Scarcity of experimental studies on the 2D MoS₂ layers-based membranes of near atomic thickness for water desalination is attributed to technical difficulties associated with integrating 2D layers preserving their structural integrity. Particularly, it has been challenging to produce large-scale (> cm²) "continuous" 2D

 MoS_2 layers of ~1-10 nm thickness and their seamless integration onto porous substrates, hindering their applications to desalination technologies.

CHAPTER 3: INTERFACIAL CHARACTERISTICS OF POLYMER DERIVED SIALCN CERAMICS¹

3.1 Introduction

The bi-phasic nature of PDCs features two phases of different conductivities, with the free-carbon phase exhibited higher conductivity than the amorphous matrix phase. The difference in conductivity between the two phases gives the materials many unique electrical properties, notably high-temperature semiconducting behaviour ^{46,158} and anomalous piezoresistivity. ^{47,159} It was shown that the dc (direct current) conductive behaviour of PDCs was determined by the concentration of the free-carbon phase. ⁹⁵ When the free-carbon content is high, a tunneling percolation network is formed by the free-carbon clusters, leading to high piezoresistivity. ¹⁶⁰ On the other hand, when the free-carbon content is low, the materials exhibit amorphous semiconducting behavior. ¹⁶¹

Previous studies also demonstrated that the electric heterogeneity resulted in an internal potential barrier at the interface between the matrix and the free-carbon clusters. This barrier caused the transition of the ac (alternating current) conduction mechanism from the long-distance transport of the charge carriers to the interfacial polarisation. The barrier also led to a very high permittivity in PDCs. Given its importance in determining the dielectric behaviour of PDCs, the behavior of the barrier itself has yet been explored.

In this study, the behaviour of the internal potential barrier is studied by using a polymerderived amorphous silicoaluminum carbonitride (a-SiAlCN) as the model material, which

analyzed the data.

¹ Part of contents in this Chapter contains published works adapted with permission from: Hao Li, Baisheng Ma, & Linan An. Effect of temperature on internal potential barriers in polymer-derived amorphous ceramics. *Ceramics International* **45**, 13575-13578 (2019). Copyright 2019 Elsevier Ltd and Techna Group. Published on March 30th 2019. Hao Li prepared the samples and conducted part of materials characterization, as well as

exhibited excellent oxidation and corrosion resistance.^{58,109} It is revealed that the interfacial capacitance of the material can be described by a double Schottky barrier model. The height of the barrier and the charge-carrier concentration exhibit Arrhenius dependence on testing temperature and annealing temperature respectively. The results are discussed in terms of the electronic structures of the material.

3.2 Experimental Methods

The a-SiAlCN studied here was prepared from a polyaluminasilazane precursor following the procedure as previously reported. 163 First, the precursor was synthesized by mixing/reacting liquid polysilazane (VL20, Kion, Huntingdon Valley, PA) as the major precursor with 20 wt% poly (melamine-co-formaldehyde) acrylated solution (PVN, Sigma-Aldrich, St. Louis, MO) as the N source and 5 wt% aluminumtri-sec-butoxide (ASB, Sigma-Aldrich, St. Louis, MO) as the Al source. The mixing/reacting was carried out at 80 °C for 12 h under continuous stirring to ensure the completion of the reaction. The synthesized precursor was then photo-cured under ultraviolet light, followed by heat-treatment at 120 °C for 12 h to enhance cross-link. The crosslinked samples were finally annealed at different temperatures (i.e., 1100, 1300, and 1400 °C) for 4 h to obtain a-SiAlCN ceramic specimens of 10 mm in diameter and 1 mm in thickness. The entire procedure was protected by flowing ultrahigh-purity argon. The obtained samples were characterised via X-Ray diffraction utilising a Rigaku D/MAX Xray Diffractometer (XRD Rigaku, Tokyo, Japan) with a monochromatic Cu-K α radiation (λ =0.154 nm), which confirmed that they are amorphous without detectable crystalline phase. Furthermore, Raman spectroscopy was also used to examine the microstructural evolution of the free carbon phase in the material. Raman spectra of samples prepared under different pyrolysis temperatures were obtained from the Renishaw inVia Raman microscopy (Renishaw Inc., Gloucestershire, UK).

The excitation source was a 532 nm line of silicon-solid laser. For each sample, 25 Raman spectra were obtained by a 10 μ m \times 10 μ m mapping acquisition for measurement accuracy.

To obtain the complex impedance spectra of the samples, silver paste was painted on their surfaces as the electrodes. The spectra were acquired in a frequency range of 4 Hz–10 MHz on an Agilent 4294 A precision impedance analyzer with a 16065 A external voltage bias fixture. The measurement was performed at different temperatures (i.e., testing temperature range of 50-150 $^{\circ}$ C and on samples underwent annealing treatment from 1100 to 1400 $^{\circ}$ C), and dc bias (0-10 V).

3.3 Results and Discussions

3.3.1 Raman Spectroscopy and XRD

Raman spectroscopy is a useful characterisation tool for investigating the formation and evolution of free carbon phase in PDCs.¹⁶⁴ As shown in the Raman spectra (Figure 16), two prominent bands at approximately 1350 cm⁻¹ ("D-band" resulting from breathing modes of sp^2 carbon atoms in rings) and 1582 cm⁻¹ ("G-band" due to in-plane bond stretching of sp^2 carbon) can be observed, which feature the presence of free carbon.^{165,166} As shown in Figure 16, both the D and G peaks exhibit shifting trend as annealing temperature increases, with the G peaks showing an up-shift, whereas the D peaks exhibit an initial up-shift before remained stable around 1350 cm⁻¹. Such shifts stemmed from microstructural changes: the up-shift of D peak reflects the increase in the number of ordered three-fold aromatic rings, whereas that of the G peak indicates the transition from disordered carbon to nanocrystalline carbon.¹⁶⁷ Such shifting patterns along with the decrease in the widths of D and G peaks indicate the graphitisation process of the free carbon phase as it transitions from amorphous carbon to nanocrystalline

graphite as annealing temperature increases. ^{166,167} An increase of the D-to-G peak intensity ratio was observed as annealing temperature increased, resulting from the in-plane growth of nanopolycrystalline graphite. ^{166,167} In addition, it has been revealed that during the pyrolysis stage of PDCs, large amounts of unpaired electrons (i.e., dangling bonds) are formed owing to the release of hydrogen and methane, ¹⁶⁸⁻¹⁷⁰ where the unpaired electrons are related to the carbon-dangling bond in the free carbon phase. ^{3,170} The presence of these unpaired electrons facilitates charge accumulation under AC field at the interface between the matrix and free carbon phases which exhibit different conductivities. ¹¹⁸

XRD analysis revealed that samples pyrolyzed at up to 1400 °C are amorphous exhibiting no obvious diffraction peaks, indicating that they are all amorphous without significant indications of the formation of crystalline carbon, which is expected to appear at higher temperatures (Figure 17). The amorphous feature is common to most polymer derived ceramics with comparable annealing history.^{114,171}

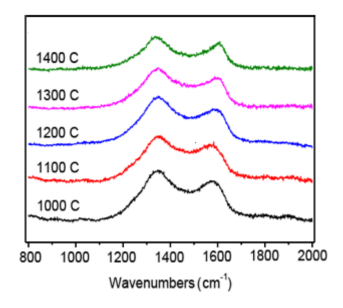


Figure 16 Raman spectra of SiAlCN samples pyrolyzed at different temperatures.

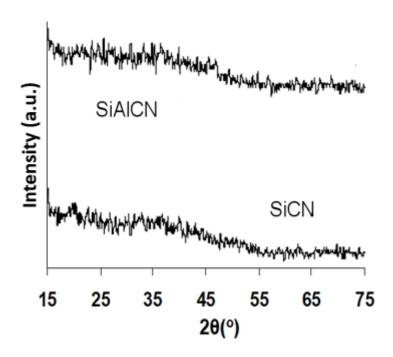


Figure 17 XRD of SiAlCN sample at 1400 °C, with that of SiCN for reference.

3.3.2 Effect of Testing Temperature

Figure 18 shows the typical complex impedance spectra of the *a*-SiAlCN, obtained under various testing temperatures without the effect of bias. It is seen that all the spectra contain two semi-circular arcs. Previous study demonstrated that the one in high-frequency region (left) corresponds to the free-carbon phase, and the other in low-frequency region (right) corresponds to the amorphous matrix. In addition, only moderate decrease in impedance was observed as the temperature increased, since such modest change in testing temperature would not cause microstructural rearrangements or any changes in chemical compositions in PDCs due to its superior thermal stability.

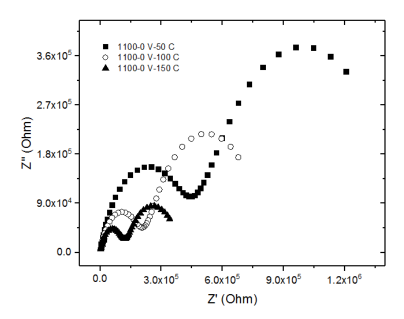


Figure 18 Influence of testing temperature on the complex impedance of the *a*-SiAlCN ceramics, without bias.

Typical impedance spectra of the *a*-SiAlCN under different bias are shown in Figure 19, with the specimen kept under 150 °C testing temperature. It is interesting to note that the response of the free-carbon phase is not affected by the dc bias; but the response of the amorphous matrix is greatly held down as the dc bias increases. This indicates that the arc in low-frequency area has significant contribution from the interfacial polarisation.¹⁷²

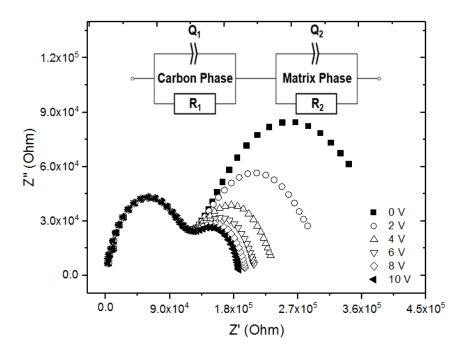


Figure 19 The complex impedance spectra of the a-SiAlCN, measured at 150 °C under different dc bias as labeled. The inset is the equivalent circuit used to analyze the spectra.

The spectra were further analysed by curve-fitting them with a modified-Debye equivalent circuit comprised of two parallel constant phase element (CPE or Q)-resistance (R) circuits connected in series (the inset in Figure 19). The parameters obtained from the best fitting are summarized in Table 2. As can be seen, the capacitance (C_2) of the amorphous matrix is about three orders of magnitude higher than that (C_I) of the free-carbon phase, suggesting that the capacitance of the amorphous matrix is mainly due to the contribution from the internal barriers. 172 The applied bias has negligible effect on the resistance and capacitance of the freecarbon phase. In contrast, the resistance of the amorphous matrix and the capacitance of the interface decrease with increasing dc bias levels. Similar trends were observed at all testing temperatures.

Table 2 Parameters obtained by curve-fitting the impedance spectra in Figure 19

Bias (V)	R ₁ (ohm)	C ₁ (nF)	R ₂ (ohm)	C ₂ (nF)
0	1.18×10^5	4.46×10^{-3}	2.89×10^{5}	2.69
2	1.17× 10 ⁵	4.42×10^{-3}	1.88×10^{5}	1.49
4	1.16×10^5	4.36×10^{-3}	1.23×10^5	1.15
6	1.15×10^5	4.33×10^{-3}	9.68×10^4	0.97
8	1.15×10^{5}	4.31×10^{-3}	8.33×10^4	0.88
10	1.15×10^5	4.31×10^{-3}	7.50×10^4	0.81

Previous studies suggested that in heterogeneity materials, a Schottky barrier could be formed at the interface between the phases with different conductivities.¹⁷³ This barrier provides extrinsic capacitance. The dependence of the interfacial capacitance on the applied dc bias can be described by a double Schottky barrier model.^{172,173} According to the model, the interfacial capacitance can be related to the dc bias by the following equation¹⁷⁴

$$\left(\frac{1}{C} - \frac{1}{2C_0}\right)^2 = \frac{2}{\varepsilon' q N_d} (\phi_b - V) \tag{1}$$

where C and C_o are capacitances of the interface with and without the applied dc bias, respectively; ε' is the permittivity of the material; q is the electronic charge; N_d is the charge carrier concentration in the more conducting phase; ϕ_b is the height of the potential barrier; and V is the applied dc bias. This equation suggests that the modified capacitance $((\frac{1}{c} - \frac{1}{2c_o})^2)$ should exhibit a linear relationship with the bias.

Figure 20 is a typical plot of $(\frac{1}{C} - \frac{1}{2C_0})^2$ as a function of V, using the data obtained from the a-SiAlCN at 150 °C (Table 2). The good linear trend indicates that the double Schottky barrier model well represents the behaviour of the a-SiAlCN, similar to that reported previously. This result is not surprising since the microstructure of the a-SiAlCN is comparable to that of the inter barrier layer capacitor (IBLC), 173,175 with the free-carbon clusters analogous to the relatively conducting grains and the amorphous matrix analogous to the relative insulating grain boundaries.

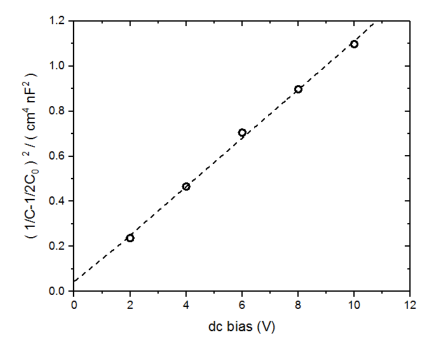


Figure 20 Plot of the modified interfacial capacitance vs. dc bias, using the data obtained at 150°C.

The interfacial behaviour was then studied by estimating the charge-carrier concentration and the potential barrier height of the *a*-SiAlCN from the intercept and the slope of the fitting lines obtained at various testing temperatures. Figure 21 shows the dependence of the charge-carrier concentration and the potential barrier height on testing temperature. It is seen that the charge-

carrier concentration increases with increasing testing temperature, but the barrier height decreases with increasing testing temperature. The figure also reveals that both the charge-carrier concentration and the barrier height exhibit Arrhenius dependence on testing temperature, indicating that they are dominated by thermally active processes. The estimated activation energies are 0.07 eV and 0.054 eV for charge-carrier concentration and barrier height, respectively. The difference in the activation energy suggests that they are governed by different processes.

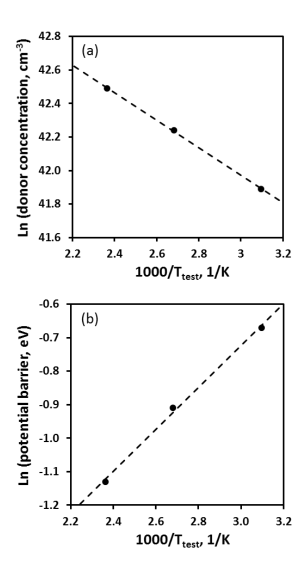


Figure 21 The influence of testing temperature on the charge-carrier concentration (a) and potential barrier (b).

The above phenomena can be correlated to the unique heterogeneous structure of the *a*-SiAlCN. Previous studies demonstrated that both the free-carbon phase and the amorphous ceramic matrix are *n*-type semiconductors with significantly different conductivities. The gaps between Fermi energy and conduction band are different for the amorphous Si-based ceramics and the free-carbon phases. A Schottky barrier can then be established between the two phases with the potential height approximating to the difference between their Fermi levels. As

a result, double (back-to-back) Schottky barriers are created at the interfaces between *n*-type free-carbon clusters (Figure 22).

The formation of the internal potential barrier is discussed below. At the point immediately after the bi-phasic PDC is under external AC electric field, the free carbon phase and the ceramic matrix phase establish close electrical contact where the respective Fermi energy levels (E_F) are still different and not in thermal equilibrium (i.e., as shown in the flatband diagram in Figure 22a, where E_{BT} and E_F represent the mobility edge for band-tail and Fermi level at this point, respectively). Because of the Fermi energy levels difference between the free carbon and the ceramics matrix phase, the electrons in the free carbon phase tend to transverse through the junction to lower their energy. The departed electrons leave behind, due to the ionised donor atoms, a positive charge, which then causes the formation of a negative field and lowers the band edges of free carbon. Thermal equilibrium will finally be achieved (i.e., no net electrons flow) when the free-carbon-to-matrix electrons diffusion equals the electrons-drifting stemming from the formed internal electric field.¹⁷⁷ A leveled Fermi energy is thus reached at thermal equilibrium, with the band tail being bent therefore a potential barrier to overcome (as shown in Figure 22b).

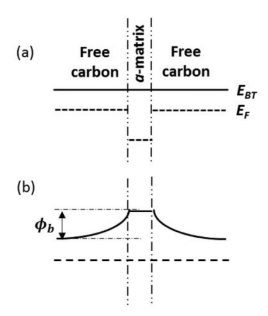


Figure 22 Schematic illustrations of the energy structure of the a-SiAlCN, (a) flatband diagram and (b) thermal equilibrium diagram. A back-to-back double Schottky barrier is formed as the gap between the band tail (E_{BT}) and Fermi level (E_F) for the free-carbon phase is different from that of the matrix phase.

According to this model, the increase in charge carrier concentration with temperature (Figure 21a) suggests that more electrons are excited to the band tail in the free-carbon phase. This can be understood as follows. For crystalline semiconductors, the charge-carrier concentration is fixed by the doping concentration and is temperature independent within a reasonable temperature range. However, for amorphous semiconductors, since chemical environment is different from defect to defect, the energy level of defects (dangling bonds) is varied in a range to form a narrow defect band within the band gap. Accordingly, increasing temperature will activate more defects, leading to more charge carriers. The activation energy estimated from the plot in Figure 21a is likely corresponds to the difference between the band-tail and defect levels. This is consistent with the fact that the material exhibited the band-tail hopping conduction behavior in the temperature range used here. 158

The cause for the change in barrier height with temperature (i.e., a 0.19 eV change within a temperature range of 100 °C) is rather complex. Two possible factors could lead to an increase in Fermi level: (1) increasing temperature (increasing temperature could reduce the band gap); and (2) increasing concentrations of active defects. In the present study, the first factor (i.e., band gap drop due to increasing temperature) would play only a negligible role, since the coefficient for the temperature-dependent band gap change would only appear at a miniscule level, ¹⁷⁸ and therefore would not be able to cause any significant effect over a moderate (100 °C) temperature increase. On the other hand, increasing temperature leads to higher levels of active dopants (dangling defects) concentrations for both the free carbon and the amorphous matrix, which lead to an increased Fermi level in both phases. However, the two phases experience different degrees of the Fermi level changes: the amorphous matrix would witness a more substantial rise (exhibiting stronger temperature dependence), because of the deeper defect levels in the amorphous matrix compared to its free carbon counterparts. Such variation in the magnitudes of Fermi level rise in the two phases would therefore result in a reduced difference between the Fermi energy in the free carbon and amorphous matrix phases, which leads to the decrease in the potential barrier height. Since the charge-carrier concentration has Arrhenius temperature dependence for both the free carbon and the amorphous, the potential barrier should exhibit similar Arrhenius temperature-dependence, which is consistent with the experimental results shown in Figure 21b.

3.3.3 Effect of Annealing Temperature

As a more systematic approach towards studying the interfacial behaviors of PDCs, the influence of annealing temperatures on the internal potential barrier was also assessed in addition to that of the testing temperature. As discussed in the above section, the range of

variation in testing temperature did not cause microstructural rearrangements or band gap drop.

Instead, an increased Fermi level in both phases was induced by increased concentrations of thermally-activated defects.

In contrast, variations in annealing temperatures of PDCs would have profound effects in both microstructural evolutions and electronic structure changes. 171,179 It is well-established that both the free carbon phase and the amorphous matrix phase experience a gradual increase in the degree of order with increasing annealing temperature. 115 On one hand for the free carbon phase, increasing annealing temperature induces a transformation from amorphous carbon to nanocrystalline graphite, 105 accompanied by the sp^3 – sp^2 transition. 180 One the other hand for the amorphous matrix phase, it has been revealed that as annealing temperature increases, the major structural evolutions involve the de-mixing of the SiC_xN_{4-x} mixed bonds, resulting in the shrinkage of the SiC_xN_{4-x} area, thus a reduced number of the mix-bonded silicon tetrahedral, and therefore a more ordered structure. 170,171,181,182

Figure 23 shows the typical complex impedance spectra of the *a*-SiAlCN, obtained under various annealing temperatures, under the same testing conditions without the effect of dc bias. It shows more drastic changes in impedance due to increased annealing temperatures in comparison with the influence of varying testing temperature (e.g., as shown in Figure 18).

Superimposing bias voltage at different levels has induced very similar effects in the impedance behaviours (as observed in Figure 19), which also exhibit two semi-circular arcs corresponding to the amorphous matrix and the free carbon phase, with the former showed significant diminishing features as bias level rose and negligible influence on the free carbon phase. Such features again indicate significant contribution from the interfacial polarisation to the amorphous matrix phase (represented by the low-frequency semi-arc). The spectra were

analysed by curve-fitting them with the modified-Debye equivalent circuit (the same as shown in the inset in Figure 19). Comparable trends in the changes in the resistance and capacitance of the amorphous matrix and free carbon phases were observed across all samples under similar measurement conditions as discussed in Section 3.3.2. Similar to that discussed above, the contribution from the internal barriers accounts for the larger capacitance of the amorphous matrix.

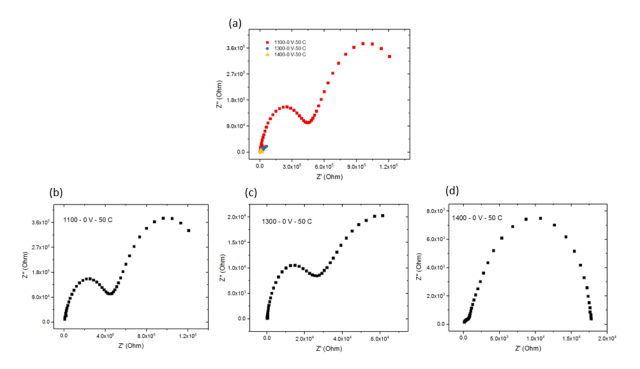


Figure 23 Influence of annealing temperature on the complex impedance of the *a*-SiAlCN ceramics, under 50 °C without dc bias voltage. Combined plots of different annealing temperatures are shown in (a) in the same scale, and the individual plots corresponding to 1100 °C, 1300 °C and 1400 °C are shown in (b), (c), and (d) respectively.

The double Schottky barrier model (as discussed in detail in Section 3.3.2) was similarly applied to describe the bias dependent interfacial capacitance to account for the influence of annealing temperature. Good linear fitting trend was observed in the modified capacitance

 $((\frac{1}{c} - \frac{1}{2C_o})^2)$ versus bias voltage (*V*) plots, similar to that shown in Figure 20. The charge carrier concentration and the potential barrier height of the *a*-SiAlCN are estimated from the intercept and the slope of fitting lines for samples obtained under different annealing temperatures. Figure 24 shows the annealing temperature dependence of the charge-carrier concentration and the potential barrier height, with both parameters increase with elevated annealing temperature.

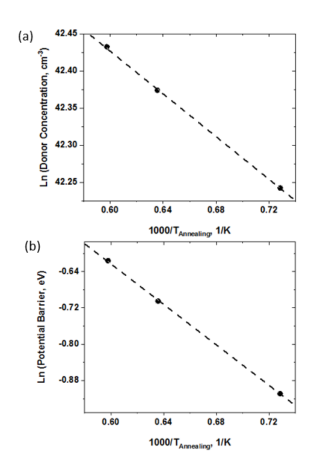


Figure 24 The influence of annealing temperature on the charge-carrier concentration (a) and potential barrier (b).

Such phenomena are also correlated to the heterogeneous structure of the *a*-SiAlCN (i.e., biphasic material comprised of a free carbon phase and an amorphous ceramic matrix phase possessing significantly different conductivities). As discussed above, double (back-to-back)

Schottky barriers are created at the interfaces between n-type free carbon clusters, since the gaps between Fermi energy and band tail are different for the amorphous Si-based ceramics and the free carbon phases (Figure 22). 161,176

Previous studies have suggested that in the free carbon phase, an increased amount of C-dangling bonds would lead to the increase in charge carrier concentration with rising annealing temperature (Figure 24a). As discussed above, a defect band is formed within the band gap, with the defects are predominantly the C-dangling bonds, which act as donors to provide electrons. Previous studies have revealed that the concentration of the C-dangling bonds increases with increasing annealing temperature. Therefore, the increase in C-dangling bonds leads to an increase in the density of the defect states, resulting in more defects, leading to more charge carriers, thus higher donor concentrations with increasing annealing temperature.

With respect to the increase of potential barrier height with increasing annealing temperature (Figure 24b), it can be understood as follows considering the changes in the respective Fermi levels in the free carbon and amorphous matrix phases. It has been revealed that C-dangling bonds are present as defects in the free carbon phase, and also within the bulk of the amorphous ceramic network, with the free carbon contains much more defects than the amorphous matrix (i.e., higher defect concentration). In addition, investigations have shown that the concentration of the C-dangling bonds increases with increasing annealing temperature, leading to an increase in the density of the defect states. Therefore, due to the increase in annealing temperature, both the free carbon and the amorphous matrix witness higher levels of active dopants (dangling defects) concentrations, which lead to an increased Fermi level in both phases, both moving towards the band tail, which is consistent with previous reports. The state of the phases, both moving towards the band tail, which is consistent with previous reports.

However, the two phases experience different degrees of the Fermi level increase: the free carbon would witness a more significant rise, because of the higher defect concentration compared to that of the amorphous matrix. Such variation in the magnitudes of Fermi level rise in the two phases would therefore result in an enlarged difference between the Fermi energy in the free carbon and amorphous matrix phases, which leads to the increase in the potential barrier height.

3.4 Conclusions

The complex impedance spectra of the polymer-derived *a*-SiAlCN were measured under various dc bias voltages in a temperature range between 50 and 150 °C, on the samples annealed at different temperatures (1100-1400 °C). All spectra, regardless of temperature and bias, consist of two semi-circular arcs, corresponding to the free-carbon phase and the interface, respectively. The impedance of the free-carbon phase is independent of the bias, while that of the interface decreased significantly with increasing dc bias. It is shown that the change of the interfacial capacitance with the bias can be explained using the double Schottky barrier model. The charge-carrier concentration and potential barrier height were estimated by comparing the experimental data and the model. The results revealed that increasing testing temperature led to an increased charge-carrier concentration and a reduced barrier height, both following Arrhenius dependence, whereas the increase in annealing temperature resulted in increased charge-carrier concentration and barrier height. The phenomena were explained in terms of the unique bi-phasic microstructures of the material.

CHAPTER 4: FEW LAYER 2D MOS₂ MEMBRANES OF NEAR ATOMIC THICKNESS FOR HIGH EFFICIENCY WATER DESALINATION²

4.1 Introduction

The availability of freshwater resources has decreased with increasing potable, agricultural, and industrial water demands around the globe for the past decades.⁵ This global challenge, along with the adversely accelerating climate change, has been calling for immediate solutions for meeting the demands. Apart from the conservation of existing water resources, increasing the freshwater supply is deemed as a more sustainable and direct solution. Since seawater is readily accessible and abundant, various thermal and physical desalination technologies have been utilised worldwide to produce drinking water by removing dissolved salts and other minerals from saline water. While multi-stage flash (MSF), multi-effect distillation (MED), and reverse osmosis (RO) are the most widely used desalination technologies, RO membrane is gaining market share at a faster pace.^{7,8} RO is a high-pressure membrane process that utilises a semi-permeable polymeric membrane to retain salts and low-molecular-weight solutes while allowing the passage of water via diffusion. Despite the operational simplicity, RO membranes suffer from several technical limitations including low conversion efficiency, membrane fouling, and energy demand to pressurise the feedwater.^{9,10} Accordingly, developing highperformance membrane materials with improved water recovery without compromising the solute rejection remains a compelling demand.

² Part of contents in this Chapter contains published works adapted with permission from: Hao Li, Tae-Jun Ko, Myeongsang Lee, Hee-Suk Chung, Sang Sub Han, Kyu Hwan Oh, Anwar Sadmani, Hyeran Kang, and Yeonwoong Jung. Experimental Realization of Few Layer 2D MoS₂ Membranes of Near Atomic Thickness for High Efficiency Water Desalination. *Nano Letters* 19, 5194–5204 (2019). Copyright 2019 American Chemical Society. Published on July 1st 2019. Hao Li co-conceived the project, prepared the membrane materials and characterized them, performed the desalination tests, and analyzed the data.

A potentially promising solution stands with engineering the structure of traditional materials into tailored porosity or exploring a new type of intrinsically porous nanomaterials-based membranes. Recently developed two-dimensional (2D) materials present unprecedented opportunities for water desalination and molecular sieving technologies owing to their structural uniqueness. 11-13 Their extremely small thickness (i.e., a few Å for 2D monolayers) projects superior water permeation rates benefiting from small diffusion lengths for molecular transports while their high mechanical flexibility renders advantages for membrane integrations. Moreover, 2D materials produced by chemical synthetic routes possess highdensity "intrinsic" structural imperfections (e.g., near atomic vacancies) whose physical dimensions are comparable to those of salt ions and water molecules. 14 In this endeavour, a variety of 2D materials have been explored including graphene, 15-18 dichalcogenides, 19-22 and graphene oxides,²³⁻²⁶ along with substantive efforts for the structural engineering of traditional membrane materials such as zeolites, ²⁷⁻³⁰ and metal-organic frameworks ³¹⁻³⁴. Amongst them, 2D molybdenum disulfide (MoS₂) layers have recently gained significant interests for a wide range of water and environmental technologies. They not only share the intrinsic structural advantages inherent to other 2D materials but also present additional suitability such as excellent chemical stability and non-toxicity. 35-37 Furthermore, a variety of structural variations such as point defects, grain boundaries, and van der Waals (vdW) gaps have been ubiquitously observed in 2D MoS₂ layers grown via chemical vapour deposition (CVD) process.³⁸⁻⁴⁰ Such "intrinsically" present structural imperfections of very high concentration (e.g., areal density of ~10¹³ cm⁻² for sulphur vacancies⁴¹) offer 2D MoS₂ layers high promise as an efficient membrane material for water desalination applications. In fact, several studies based on theoretical calculations and molecular dynamics simulations have predicted superior desalination performances of monolayer 2D MoS₂. ^{14,19,41,152,153} Heiranian et al. have predicted

that 2D MoS₂ monolayers of enriched defective sites can exhibit ~2-5 orders-of-magnitude higher water fluxes than traditional filtration membranes, which is even 70% greater than the performance of graphene nanopores. 154 Köhler et al. have projected that the ion rejection capability of 2D MoS₂ membranes is superior to graphene under a wide range of test conditions including pressure, nanopore size, and cation valences. 152 Li et al. have simulated the desalination performance of 2D MoS₂ layers, demonstrating high water transport rate (355.3 L m⁻² h⁻¹ bar⁻¹) and excellent salt rejection capability. ¹⁵⁵ Despite the theoretically-projected promise of 2D MoS₂ layers, their experimental utilisation for membrane-based desalination technologies has been rarely approached. A few experimental studies have explored MoS₂ laminar membranes of very large thickness (typically, ~1-10 µm) constituting mechanically stacked individual 2D flakes. 14 Sun et al. studied ~1.7 µm thick MoS₂ membrane films prepared via chemical exfoliation and subsequent filtration of aqueous dispersion. 156 Hirunpinyopas et al. demonstrated chemically functionalized ~5 µm thick laminar membranes composed of exfoliated 2D MoS₂ flakes employing the similar filtration process. ¹⁵⁷ These laminar 2D MoS₂ membranes of large thickness have exhibited significantly limited desalination efficiencies, failing in meriting the structural uniqueness and performance advantages projected with near atom-thick membranes. For example, their water permeation rates were experimentally identified to be in a range of ~245-275 L m⁻² h⁻¹ bar⁻¹, ^{156,157} significantly falling short of theoretical values predicted by simulation studies.¹⁵⁴ Moreover, ion rejection rates of mere ~15-30% were reported for pristine laminar 2D MoS₂ membranes of ~1-5 µm thickness. 157 Scarcity of experimental studies on the 2D MoS₂ layers-based membranes of near atomic thickness for water desalination is attributed to technical difficulties associated with integrating 2D layers preserving their structural integrity. Particularly, it has been challenging to produce

large-scale ($> cm^2$) "continuous" 2D MoS₂ layers of ~1-10 nm thickness and their seamless integration onto porous substrates, hindering their applications to desalination technologies.

In this Chapter, we discuss the experimental realisation of high-efficiency water desalination enabled by ultra-thin 2D MoS₂ layers-based membranes. We demonstrate the integration of centimetre-scale CVD-grown 2D MoS₂ layers of only ~7 nm thickness onto porous polymeric substrates and studied their desalination performances. This new type of near atom thickmembranes exhibits an excellent combination of high water permeability approaching theoretically predicated values and superior ionic sieving performance for various seawater salts. Fundamental mechanisms responsible for the observed high desalination efficiency are also discussed.

4.2 Experimental Method

CVD growth of 2D MoS₂ layers: Very thin films of Mo (typically, ~2-2.5 nm thickness) were deposited onto SiO₂/Si substrates via electron beam evapouration to ensure the growth of horizontally-oriented 2D MoS₂ layers avoiding their unwanted vertical alignment. As-deposited Mo-SiO₂/Si samples were then placed inside a 1 inch quartz tube located in the middle of a horizontal tube furnace (Lindberg Blue M, ThermoScientific). Subsequently, sulphur powders (99.5%, Sigma-Aldrich, St. Louis, MO) were loaded within the quartz tube at the furnace upstream side. The furnace was pumped down to a base pressure of 5 mTorr following purging with high-purity argon (Ar) gas. It was then ramped up to 780 °C and was held for 50 mins at a ramping rate of 15 °C/min under a constant flow of Ar gas (100 SCCM). After the reaction, the furnace was naturally cooled down to room temperature under Ar supply.

Water permeation rate measurement: A typical external pressure method was used to assess the water permeation rate by employing a common dead-end filtration setup. The sandwiched membrane containing 2D MoS₂ layers was placed inside a leak-free syringe filter holder. The water permeation rate was calculated by weighing the volume of water passing through the 2D MoS₂/PES membrane area under a constant pressure of 1 bar as a function of time.

Salt ion rejection rate measurement: A conductivity probe (ThermoScientific, model: Orion 013605MD) connected to a solute conductivity meter (ThermoScientific, model: Orion 5 Star) was used to monitor conductivities in respective containers. The probe was thoroughly rinsed each time before each measurement. The salt rejection rate was calculated by $\left(1 - \frac{C_P}{C_F}\right) \times 100\%$, where C_P and C_F are the conductivity of the permeate and feed side, respectively. Seawater salts of NaCl (99%, Sigma-Aldrich, St. Louis, MO), KCl (99%, Sigma-Aldrich, St. Louis, MO), MgCl₂ (98%, Sigma-Aldrich, St. Louis, MO), and CaCl₂ (93%, Sigma-Aldrich, St. Louis, MO) of various concentrations were employed for rejection rates measurements. Each measurement was repeated for three times to improve consistency and averaged values are presented.

TEM/STEM characterisation: The crystalline structure of 2D MoS₂ layers was characterised using a JEOL ARM200F FEG-TEM/STEM with a Cs-corrector. Plane-view TEM samples were prepared by delaminating the 2D MoS₂ layers from SiO₂/Si substrates inside water and by transferring them to holey carbon TEM grids. Cross-sectional TEM samples were prepared by focused ion beam (FIB)-based milling and lift-out techniques. As-grown 2D MoS₂ layers-SiO₂/Si substrates were coated with a carbon film of ~100 nm thickness and were subsequently cross-sectioned inside a FIB (Quanta 2D FEG, FEI). Ga ion milling (30 keV) was executed until the target area became electron transparent and the prepared

specimen was transferred to a TEM grid using a micromanipulator (Omniprobe). All TEM operations were performed at an accelerating voltage of 200 kV.

AFM characterisation: Topography images and height profiles of the 2D MoS₂ layers integrated on PES supporters were obtained with AFM (nanoIR2 by Anasys Instruments, Santa Barbara, CA) operated in a tapping mode at a scan rate of 1 Hz. AFM probes (PR-EX-T125-10, Anasys Instruments) with a nominal spring constant of 30 N/m were employed.

MD simulation: MD simulations were performed using a large-scale atomic/molecular massively parallel simulator (LAMMPS) package 188 in a similar way to a recent computational study. 154 We generated a 2D MoS₂ layer (4.1 × 3.5 nm²) with a nanopore (1.6 nm in diameter) and applied a constant pressure (250 atm) to the graphene piston using VMD 1.9.1. 189 The simulation system is $6.0 \times 5.6 \times 11.5$ nm³ in size containing ~6,600 atoms. Lennard-Jones (LJ) parameters for 2D MoS₂ layer, 154 graphene sheet, 190 and NaCl 191 were obtained from previous studies. The mixing rule was applied to obtain the remaining LJ parameters following the recent computational study. 154 Specifically, TIP3P water model 192,193 and SHAKE algorithm were employed to maintain water molecules in rigid states. The cut-off distance for LJ was set to 10 Å and long-range electrostatic interactions were calculated using the Ewald method. 194

4.3 Results and Discussion

4.3.1 2D MoS₂ Layers Membrane Preparations and Characterisations

Centimetre-scale 2D MoS₂ layers were grown by the modified CVD process developed in our lab.¹⁹⁵⁻¹⁹⁷ A thin film of Mo (typically, ~2-2.5 nm thickness) was deposited onto a SiO₂/Si substrate via electron beam evapouration. The as-deposited Mo-SiO₂/Si sample was thermally sulphurised inside a CVD chamber under the controlled flow of vapourised sulphur (S), which

yields the conversion of Mo to 2D MoS₂ layers. Detailed conditions for 2D MoS₂ layers growth are described in Method section. After the CVD reaction, as-prepared 2D MoS₂-SiO₂/Si sample was gently immersed into de-ionised water until the 2D MoS₂ layers were spontaneously delaminated. The separated 2D MoS₂ layers floating on the water surface were then gently scooped up with a polyethersulfone (PES) membrane substrate (~0.03 µm pore size and ~100 μm thickness; commercial products from Sterlitech, Kent, WA). Note that PES has been extensively employed as a substrate for conventional desalination ¹⁹⁸ as well as 2D materialsbased membranes. 199 Following the integration of 2D MoS₂ layers on PES, a thin film of polydimethylsiloxane (PDMS) was selectively deposited on the 2D MoS₂/PES surface leaving a small opened area which is to function as a membrane directly exposed to water. Figure 25(a) illustrates the membrane preparation procedure, from the large-area CVD growth of 2D MoS₂ layers and their integration onto PES substrate as well as selective PDMS sealing. Given that the porosity of PES substrates is much larger than the intrinsic near-atomic vacancies inherent to 2D MoS₂ layers (to be verified in the later section), water permeation and ionic sieving are exclusively determined by 2D MoS₂ layers which are the integral part of the membrane. Prior to the integration of 2D MoS₂ layers, the PES surface can be treated with oxygen plasma to ensure the uniform adhesion of large-area 2D MoS₂ layers to it. Figure 25(b) demonstrates the water-assisted delamination of 2D MoS₂ layers from a SiO₂/Si growth substrate, revealing that only 2D MoS₂ layers become spontaneously detached upon immersion in water. The success of this facile 2D layer separation is attributed to the combined effect of the high surface energy contrast between 2D MoS₂ basal planes and SiO₂ surface as well as the weak vdW molecular attraction in between them as clarified in our previous reports.²⁰⁰

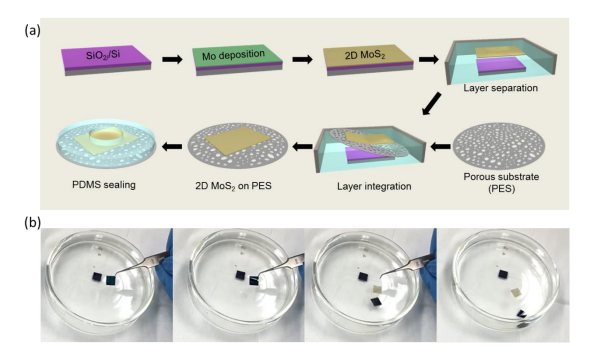


Figure 25 (a) Schematic procedures for few-layer 2D MoS₂ membranes preparation, transfer, and integration. (b) Sequential images showing the detachment process of 2D MoS₂ layers from a SiO₂/Si substrate upon water immersion.

Prior to investigating the water permeation and ion rejection capability of 2D MoS₂ layers integrated on PES substrates, we first verified the material quality of CVD-grown 2D MoS₂ layers in their as-grown state. Figure 26(a) shows an image of 2D MoS₂ layers grown on a SiO₂/Si substrate of > 2 cm² in size. The chemical and structural integrity of the as-grown 2D MoS₂ layers were characterised via Raman spectroscopy and transmission electron microscopy (TEM). Figure 26(b) presents a typical Raman spectrum of the CVD-grown 2D MoS₂ layers revealing two distinctive peaks. The peak positions correspond to the in-plane (E_{2g}^1) and out-of-plane (A_{1g}) vibration modes of 2D MoS₂ layers, confirming their presence on the growth substrate. Figure 26(c) shows a TEM image of the same sample in Figure 26(a) in a cross-sectional view, revealing the presence of continuous 2D MoS₂ layers of uniform thickness. Figure 26(d) shows an enlarged view of the red box in Figure 26(c), revealing the

crystalline structure of 2D MoS₂ layers. The image clarifies that they are composed of horizontally-oriented individual layers of uniform ~7 nm thickness. Moreover, the individual 2D layers are separated by the physical gaps formed by vdW interaction with an average interlayer spacing of ~0.65 nm. Figure 26(e) illustrates the atomic schematics of the 2D MoS₂ layers corresponding to the red box in Figure 26(d). Although the vdW gap denotes the interlayer spacing corresponding to Mo-to-Mo distance across adjacent 2D layers, the actual "physical gap" which serves as a transport channel for molecular diffusion in this case should be close to ~0.3 nm determined by the interlayer S-to-S distance. Note that the dimension of this transport channel is larger than that of a single water molecule, i.e., ~0.275 nm in a diagonal length. 203 Figure 26(f) shows a plane-view high-resolution TEM (HRTEM) image of the horizontally-oriented 2D MoS₂ layers. The image clearly reveals the Moir é fringes of hexagonal MoS₂ basal planes, reflecting the crystallographic stacking of mis-oriented individual 2D layer grain as well as indicating the presence of grain boundaries. 195 These comprehensive characterisation results conclude that our CVD-grown 2D MoS₂ layers possess high spatial continuity and uniform thickness (~7 nm in Figure 26(c), (d)) while containing a large degree of intrinsic structural porosity (e.g., ~0.65 nm vdW gaps). Such combined attributes of macroscopic structural homogeneity and near atomic scale heterogeneity render the CVD-grown 2D MoS₂ layers high suitability as desalination membranes, as verified below.

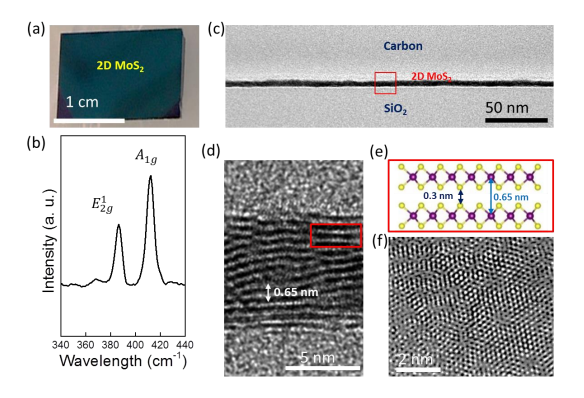


Figure 26 (a) Optical image of as-grown 2D MoS₂ layers on a SiO₂/Si substrate. (b) Raman spectrum of the corresponding CVD-grown 2D MoS₂ layers. (c) TEM image of the same sample in Figure 26(a) in a cross-sectional view. (d) Enlarged view of the red box in Figure 26(c). (e) Atomic schematics of the few-layer 2D MoS₂ corresponding to the red box in Figure 26(d). (f) Plane-view HRTEM image of horizontally-oriented 2D MoS₂ layers showing Moir éfringes.

Having identified the intrinsic structural suitability of the CVD-grown 2D MoS₂ layers for desalination applications, we then characterised the morphologies of 2D MoS₂ layers integrated on PES supporting substrates. As depicted in Figure 25(a), complete membranes are in a configuration such that the 2D MoS₂ layers are selectively covered by an impermeable PDMS film, which defines a specific region for water penetration by sealing the surrounding surface. Figure 27(a) illustrates a schematic set-up for water permeation measurements. The 2D MoS₂ layers-integrated PES substrate was covered with another bare PES substrate on the back side.

It was subsequently sandwiched in between two PDMS sealing films defining a membrane area of ~5 mm on the centre region. Note that this sandwich-like assembly has been extensively employed in configuring nanomaterials-based membranes including 2D materials of large thickness (> 1 µm). 157 This layered membrane was then loaded inside a syringe filter holder in a typical dead-end filtration setup. The water permeation rate was then measured via an external pressure method by weighing the volume of water passing through the membrane area (i.e., ~5 mm central hole composed of 2D MoS₂/PES only) as a function of time under a constant pressure. Details are presented in the *Method* section. Figure 27(b) shows a representative image of a 2D MoS₂ layers-integrated PES supporter achieved following the procedures in Figure 25(a). On top of the sample surface, PDMS was selectively deposited (denoted by yellow false colour) leaving an open membrane hole at the center while sealing the surrounding area. The morphology of the 2D MoS₂ layers integrated on PES was characterised by atomic force microscopy (AFM) and Raman spectroscopy. Figure 27(c) shows an AFM topography image of 2D MoS₂ layers whose thickness was determined by obtaining the height profile across their interface with PES (denoted by the yellow line). Figure 27(d) shows the corresponding AFM height profile revealing that the integrated 2D MoS₂ layers are ~7 nm thick which corresponds to ~10-11 layers given the interlayer spacing of ~0.65 nm. ^{204,205} The result is quite consistent with the TEM characterisation (Figure 26(d)) obtained from the sample in a pristine state before integration onto PES, indicating the high fidelity of our 2D layer integration process. Figure 27(e) shows a Raman profile of the 2D MoS₂ layers on PES, revealing two characteristic peaks of E_{2g}^1 and A_{1g} vibration modes consistent with the observation prior to their separation and integration (Figure 26(b)). This analysis confirms that the 2D MoS₂ layers integrated on PES maintain their original structural and chemical integrity, which were not altered through the membrane preparation procedures.

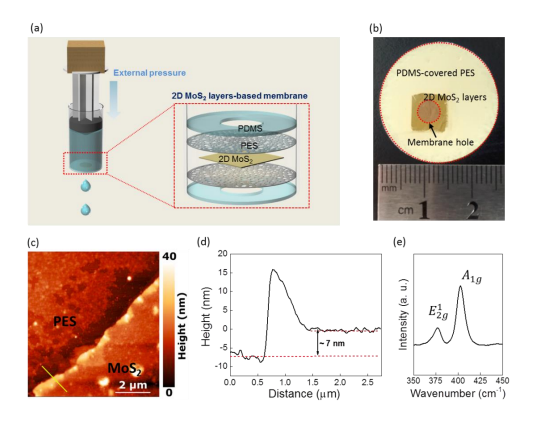


Figure 27 (a) Schematic set-up for water permeation measurement. (b) Representative image of a 2D MoS₂ layers-integrated PES supporter with PDMS sealing. 2D MoS₂ layers-membrane hole is denoted by the inner dotted circle. (c) AFM topography image of a 2D MoS₂ layers/PES interface. (d) AFM height profile corresponding to the yellow line in Figure 27(c). (e) Raman spectrum profile obtained from 2D MoS₂ layers integrated on a PES supporter.

4.3.2 Desalination Performances Evaluations and Comparisons

Desalination efficacies of 2D MoS₂ layers on PES supporters were investigated by measuring their water permeation and salt ion rejection rates. The water permeation rate was determined via the external pressure method employing the equipment described in Figure 27(a). Figure 28(a) compares the water permeation rates of bare PES vs. 2D MoS₂ layers (~7 nm thick) integrated on PES, revealing the exclusive role of 2D MoS₂ layers on controlling the

permeability of water molecules. The bare PES substrate exhibits a high water flux of 2735 (\pm 139) L m^{-2} h^{-1} bar⁻¹, reflecting its intrinsic porosity of large size and high density (~0.03 μm pore size). Integrating 2D MoS₂ layers suppresses the permeability to 322 (± 27.2) L m⁻² h⁻¹ bar⁻¹ owing to their intrinsic porosity of much smaller dimension - typically, atomic vacancies of ~1-1.5 nm in size which is to be verified in the later section. It is noteworthy that this permeability value achieved with ~7 nm thick 2D MoS₂ layers is significantly larger (up to > 30%) than those realised with the 2D MoS₂ laminar membranes of ~1-10 µm thickness, ^{156,157} indicating their suitability for desalination. Such micrometer-thick 2D MoS₂ laminar membranes have been developed by mechanically stacking up individual 2D flakes dispersed in aqueous solvents using filtration processes. Owing to the nature of high pressure-demanding methods in their preparation, the assembled 2D laminar MoS₂ layers often tend to exhibit uncontrolled physical gaps in between constituent layers. 199,206,207 For example, an interlayer spacing of ~0.3 nm has been reported, ¹⁹⁹ which is much smaller than the intrinsic vdW gap distance of ~0.65 nm observed with our CVD-grown 2D MoS₂ layers (Figure 26(d), (e)). Accordingly, the transport of water molecules in between 2D layers becomes interfered, resulting in lower water permeation rates given the diameter of a single water molecule is \sim 0.275 nm.²⁰³ Moreover, the 2D MoS₂ laminar membranes assembled by the high-pressure liquid filtration processes exhibit significantly larger thickness of ~1-10 µm compared to our CVD-2D MoS₂ layers, which further contributes to slowing down the transport kinetics of water molecules. The proficiency of our CVD-2D MoS₂ layers on PES supports for rejecting a variety of salt ions was determined using a measurement set-up whose schematic is presented in Figure 28(b). A custom-built apparatus composed of a pair of one-litre water containers and a conductivity probe was assembled with a leakage-proof pathway. The sandwiched membrane containing 2D MoS₂ layers was then placed inside the water pathway which allows for the

access of ions only through the 2D MoS₂ layers-membrane hole. The container at the permeate side was filled with pure de-ionised water while the feed side maintained an equal volume of de-ionised water mixed with salt ions to be tested. Measurements were carried out at room temperature under continuous stirring of a magnet in the feed container to minimise the concentration gradient of test salt ions. We have quantified the movement of ions by employing the conductivity measurement method well established in the field; 157 i.e., the salt rejection rate was calculated by $\left(1 - \frac{C_P}{C_F}\right) \times 100\%$, where C_P and C_F are the conductivity of the permeate and feed side, respectively.²⁰⁸ Measurement conditions are detailed in the *Method* section. Various seawater salts of diverse concentrations were tested: (1) sodium chloride (NaCl) of 0.6 M (equivalent to seawater salinity concentration²⁰⁹), (2) NaCl of 0.469 M (typical seawater Na⁺ concentration²¹⁰), (3) potassium chloride (KCl) of 0.1 M, (4) magnesium chloride (MgCl₂) of 0.1 M, and (5) calcium chloride (CaCl₂) of 0.1 M. The concentration of 0.1 M for KCl, MgCl₂ and CaCl₂ was selected for measurements consistency although the typical seawater concentrations of K^+ , Mg^{2+} and Ca^{2+} are below 0.1 $M.^{210,211}$ Figure 28(c) presents the plots of seawater ion rejection rates obtained at identical interval times during the course of three hours. Remarkably, the 2D MoS₂ layers-integrated membrane exhibits very high salt rejection capability (> 99.5%) for all the tested species at their respective concentrations. Moreover, the durability of the membrane in retaining desalination capability was evaluated by using 0.1 M NaCl solute as a testing agent over a continuous time period of > 24 hours. Figure 28(d) confirms the superior performance of the 2D MoS₂ layers-integrated membrane which retains ~ 98.3% rejection rate even after 24 hours. Apart from the rejection rate tests with single-ionic solutions, two different types of seawater containing multiple ions were employed to further assess the desalination efficacies of our CVD-2D MoS₂ layers. One was actual seawater obtained from the Atlantic coast and the other was synthetic seawater prepared to mimic the

concentrations of typical ion species present in standard seawater, i.e., Na⁺ of 0.469 mol/L ²¹⁰, K^+ of 0.01 mol/L 210 , Mg^{2+} of 0.0528 mol/L 210 , and Ca^{2+} of 0.01 mol/L 210 . Figure 28(e) presents salt rejection rates for both cases, confirming the excellent desalination capability of near 100% rejection rates. It is noteworthy that the overall desalination efficiencies of our CVD grown-2D MoS₂ layers of ~7 nm thickness are significantly higher than those of previously developed micrometre-thick 2D MoS₂ laminar membranes assembled via liquid filtration processes. 157 Detailed comparisons will be presented in the later section. Figure 28(f) presents pressure-dependent rejection rates for various salt ions of diverse concentrations. The plots reveal a slight decrease of the rejection rates with increasing applied pressure which exerts higher mechanical forces causing more ion translocation events, consistent with numerical predictions with monolayer 2D MoS_2 . Moreover, the high rejection rate of > 98.5 % is well retained for all salt ions irrespective of applied pressure within the tested range. We believe that the superior salt rejection performance observed with our CVD-2D MoS₂ layers is attributed to their ionic sieving capability owing to the intrinsically present vdW gaps of uniform spacing (Figure 26(e)). Meanwhile, the micrometer-thick 2D MoS₂ laminar membranes inevitably contain a high density of uncontrolled physical gaps in between the edges of adjoining 2D flakes owing to the intrinsic randomness associated with stacking them up in solutions. Discussion will follow in the later section of this Chapter.

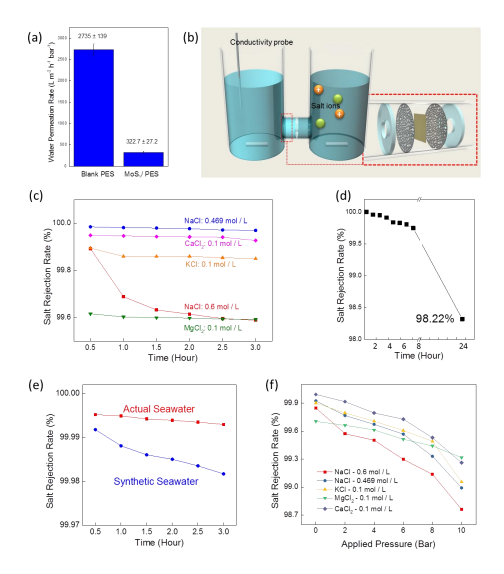


Figure 28 (a) Comparison of water permeation rates for 2D MoS₂ layers/PES vs. bare PES. (b) Schematics of salt rejection rates measurement set-up. (c) Time-dependent rejection rates for typical seawater salt ions of various species. (d) Rejection rate for 0.1 M NaCl over a continuous 24-hour period. (e) Salt rejection performances of 2D MoS₂ layers tested with actual and synthetic seawater. (f) Pressure-dependent rejection rates for various salt ions.

We compare the desalination performances of our CVD-2D MoS₂ layers with previously developed thin membrane materials. Figure 29(a) presents comparisons with a variety of systems ranging from conventional porous membrane materials to recently developed

nanostructured materials. The materials include MFI zeolite, 212 seawater reduced reverse osmosis (RO),²¹³ brackish RO,²¹³ nanofiltration,²¹³ high-flux RO,²¹³ polysulfone²¹⁴ as well as graphene oxide enhanced polyamide thin-film nanocomposite membrane (GO/PA),²¹⁵ and multi-walled carbon nanotubes incorporated polyamide (MWNTs-PA)²¹⁶ materials. The comparison plot is presented in terms of water permeability vs. salt rejection rate – two most decisive indicators for defining the performances of desalination membranes. This survey result reveals that our CVD-2D MoS₂ layers present an outstanding combination of high permeability and strong ionic sieving capability. Despite the similar ion rejection rates noted for many of the compared materials, our CVD-2D MoS₂ layers of extremely small (~7 nm) thickness exhibit orders-of-magnitude higher water permeability. We note that monolayer graphene membranes in recent experimental studies exhibit much higher water permeability than our CVD-2D MoS₂ layers, which must be attributed to their extremely small one-atom thickness. 217,218 However, it is worth mentioning that their areal dimensions are very limited, i.e., in a range of $\sim 20~\mu m^2~^{217}$ to $\sim\!10^4~\mu m^2~^{218},$ much smaller than our "centimetre-scale" membrane areas. Moreover, the fabrication of these monolayer graphene membranes has relied on top-down lithographic patterning processes, much more complicated than our facile manual integration approach. Furthermore, we compare the desalination performances of the CVD-2D MoS₂ layers with previously developed 2D MoS₂ laminar membranes of micrometre thickness prepared via liquid exfoliation/filtration processes (Figure 29(b)). Compared to the traditional membrane materials (Figure 29(a)), we note that decently higher water permeability was achieved with these micrometre-thick 2D MoS₂ laminar membranes. 157 However, they significantly fall short of our CVD-2D MoS₂ layers of ~ 7 nm thickness in terms of ionic sieving capability. Particularly, pristine 2D MoS₂ laminar membranes of ~1-5 µm thickness exhibit ion rejection rates of only ~15-30%, 157 in a sharp contrast to the performance (i.e., rejection rate > 99%) achieved with our CVD-2D MoS₂ layers which were not even functionalised for further property improvement.

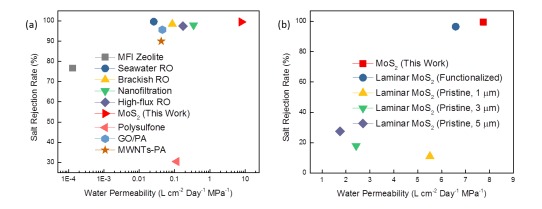


Figure 29 (a) Desalination performances of various membranes regarding salt rejection and water permeation rates. (b) Desalination performances of the CVD-2D MoS₂ layers of ~7 nm thickness developed in this study vs. 2D MoS₂ laminar membranes of micrometer thicknesses prepared via liquid exfoliation/filtration processes.

Due to the imperfect nature of CVD process optimisation, intrinsic structural variations such as atomic vacancies and grain boundaries have been regarded inevitably present in CVD-grown 2D MoS₂ layers. ^{38,40,199,219-221} For instance, various types of near atomic-scale vacancies have been identified to exist within CVD-grown 2D MoS₂ basal planes, including mono-sulphur vacancy (V_S), di-sulphur vacancy (V_{S2}), vacancy complex of Mo with three surrounding sulphur (V_{MoS3}), and vacancy complex of Mo with three surrounding pairs of di-sulphur (V_{MoS6}). ³⁸ Water permeability achieved with CVD-2D MoS₂ layers can be understood by considering the dimensional effect of this intrinsic porosity as well as its areal density; i.e., the size of these vacancies should be large enough to allow for the facile translocation of water molecules and their areal density should be large to ensure high probability for the translocation event. Amongst the above listed vacancies, it has been known that sulphur vacancies of various

sizes yield a large areal density of up to ~10¹³ cm⁻², ⁴¹ rendering CVD-grown 2D MoS₂ layers sulphur-deficient in general. 220,222 However, it is known that the areal density of large-sized (~ 1 nm) intrinsic sulphur vacancies such as V_{MoS6} is much smaller than that of small-sized vacancies, e.g., V_S of < 0.3 nm. ²²⁰ This competing situation makes it difficult to experimentally quantify and identify the kind of specific vacancies which most govern water permeability. In addition to the intrinsic atomic vacancies within basal planes, CVD-grown 2D MoS₂ layers contain a large density of intrinsic "nanopores" present along the grain boundaries formed by individually stitching 2D grains of distinct crystallographic orientation. The size of these nanopores is larger than that of the atomic vacancies within basal planes as they are generally composed of multiple un-coordinated atoms. For instance, nanopores of ~0.6–0.82 nm² in size can be naturally formed by adjoining 2D layers in a binary stack while their size can further vary with grain boundary angles. 223,224 Accordingly, their role on governing water permeation and ion rejection must become pronounced as far as they are abundantly present within grain boundaries, which must be the case for our CVD-2D MoS₂ layers as verified in the next section. Indeed, theory predicts that nanopore sizes of ~0.74–0.98 nm would yield the optimal combination of high salt rejection rate and high water permeation flux for 2D MoS₂ layers.⁴¹ In addition to the contributions from atomic vacancies and nanopores to water permeation, the uniformly spaced vdW gaps inherent to our CVD-2D MoS₂ layers (Figure 26(e)) are believed to play a significant role as well, particularly, for achieving high ionic sieving capability. In fact, the physical sizes of the hydrated ions tested in this experiment are identified to be larger than the interlayer vdW gap size of our horizontally-aligned 2D MoS₂ layers. The interlayer vdW gap which functions as a channel for molecular/ionic transport is ~0.65 nm wide accompanying the actual S-to-S distance of ~0.3 nm, as shown in Figure 26(d) and Figure 26(e). Meanwhile, diameters of the tested cations in their hydrated states are as follows: 0.66 nm (K⁺),

0.72 nm (Na⁺), and 0.86 nm (Mg²⁺).^{225,226} This observation suggests that the transportation of the hydrated ions through the vdW gap within our material is efficiently impeded, which well explains the observed high rejection rates. In addition to the above accounted dimensional/geometrical effect, electrostatic interaction of atomic vacancies is considered as another major factor for governing both water permeation and ionic sieving. Theory suggests that salt ions encounter significantly high energetic and steric barriers when approaching sulphur vacancies while water molecules are relatively unaffected.¹⁵⁵ For example, cations (e.g., Na⁺) experience high Coulombic barrier since the sulphur vacancies are positively charged due to the surplus amount of un-bonded Mo atoms.¹⁵⁵ Moreover, these atomic vacancies expose hydrophilic Mo-rich sites which can readily attract water molecules allowing for their efficient permeation while impeding ion movements, yielding high water flux and ion rejection rates.^{154,} Molecular dynamics (MD) simulation studies by Köhler *et al.* further justify this analysis, identifying that the water-accessible nanopores of up to ~1 nm present strong ion rejection capability even at a high pressure of 100 MPa.¹⁵²

The proposed desalination mechanism based on the intrinsic nanopores and vdW gaps of CVD-2D MoS₂ layers is elucidated in Figure 30. The schematic in Figure 30(a) illustrates that the nanopores primarily present within the grain boundaries of poly-crystalline 2D MoS₂ layers allow for the efficient permeation of water molecules. The illustration in Figure 30(b) shows a cross-sectional view of water permeation and ion rejection events through the interlayer vdW gaps within horizontally-aligned multilayer 2D MoS₂. While water molecules easily translocate through the nanopores as well as transport within the interlayer vdW gaps, the transportation of hydrated ions is efficiently interfered owing to their relatively large sizes. Accordingly, high water fluxes and ion rejection rates should be resultant, as above experimentally verified. The presence of such nanopores localised within grain boundaries has

been directly verified by aberration-corrected TEM characterization. Figure 30(c)-Figure 30(g) present representative TEM images of the CVD 2D MoS₂ layers of ~ 7 nm thickness, highlighting their intrinsic structural imperfections. Figure 30(c) and Figure 30(d) reveal their typical poly-crystalline structure with ample crystalline grains and grain boundaries. The red arrows in Figure 30(c) denote the areas where multiple nanopores are observed, typically across the grain boundaries. Figure 30(d) shows one large (~ 1.5 nm) localised nanopore (blue dotted circle) observed in the area different from Figure 30(c). Figure 30(e)-Figure 30(g) present HRTEM images of localised grain boundaries and their detailed crystalline structures. Figure 30(e) reveals a large-sized (~ 2 nm) highly disordered region (red dotted circle) constituting multiple nanopores of various sizes on the interfacial junction formed by the grains of two distinct crystallographic orientations. Note that the yellow triangle and the green hexagon clarify two different Moir éfringes evidencing the distinct orientation of each grain. Figure 30(f) and Figure 30(g) present zoomed-in HRTEM images of individual nanopores on the grain boundaries of the same sample - areas different from Figure 30(e). The images clearly evidence the presence of multiple nanopores the size of which varies in a range of ~0.5 nm to ~ 1 nm (denoted by the yellow circles), suitable for the permeation of water molecules. Although it is experimentally challenging to precisely quantify the areal density of such vacancies in a statistical manner, their presence is strongly believed to be responsible for the overall desalination performances.

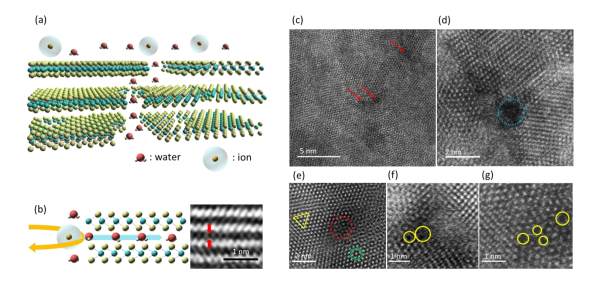


Figure 30 (a) Schematic illustration to visualize the translocation of water molecules through nanopores within the grain boundaries of horizontally-aligned multilayer 2D MoS₂. Note that the size of the nanopores is large enough to allow for water permeation. (b) Side-view illustration to show that water molecules easily transport through the interlayer vdW gaps while hydrated ions are efficiently rejected due to their larger sizes. The inset HRTEM image visualizes the vdW gap (red arrows) which functions as a water transporting channel. (c)-(g) TEM images to confirm the presence of intrinsic nanopores within the grain boundaries of CVD-2D MoS₂ layers revealing: (c) multiple nanopores, (d) a localized nanopore of ~1.5 nm in size, (e) distinct grain orientations, and (f)-(g) nanopores of various sizes.

Lastly, we further justify the exclusive role of nanopores within 2D MoS₂ layers on the experimentally observed water permeation and ionic sieving effects. We have carried out non-equilibrium molecular dynamics (MD) simulations and monitored the filtration of saline water through a 2D MoS₂ layer containing a near-atomic scale vacancy. A schematic of the simulation system is shown in Figure 31(a) which illustrates NaCl-containing saline water at the onset of a filtration process. A NaCl concentration of 0.6 M was chosen to mimic seawater salt concentration conditions. A graphene sheet placed on top of the saline water box acts as a

rigid piston and the permeation of water molecules is directly visualised using visual molecular dynamics (VMD) 1.9.1.¹⁸⁹ Upon the application of constant pressure to the piston, water molecules as well as Na⁺ and Cl⁻ start to move toward the 2D MoS₂ layer. Figure 31(b) illustrates the time-lapse snapshot of this molecular and ionic movement at varying timeframes. It is apparent that a majority of water molecules facilely pass through the pore, which becomes more pronounced as the timeframe extends from 5 ps to 40 ps, while both Na⁺ and Cl⁻ barely penetrate it. This MD simulation result well agrees with our experimental observation of high water permeability and ion rejection rates and is also consistent with the recent computational study.¹⁵⁴

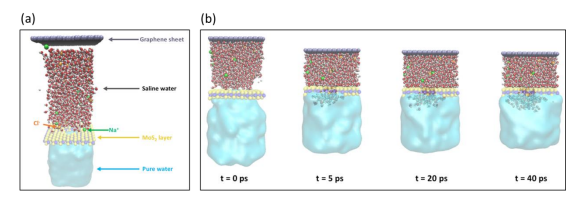


Figure 31 (a) Schematic of MD simulation system. Saline water is located between the graphene sheet which acts as a rigid piston and the 2D MoS₂ layer. Green and orange spheres indicate Na⁺ and Cl⁻ ions, respectively. Pure water is located underneath the 2D MoS₂ layer to prevent the dispersion of filtered water. (b) Time-lapse snapshot images of water desalination through the near atomic vacancy of the 2D MoS₂ layer during 40 ps MD simulations.

4.4 Conclusion

In summary, we experimentally evaluated the performances of CVD-grown few-layer 2D MoS₂ of ~7 nm thickness as water permeable membranes for desalination and molecular sieving. This near atom-thick membrane presented an excellent combination of high water permeability and high ionic sieving capability for a variety of tested ions, outperforming previously developed 2D MoS₂ layer membranes of much larger thickness. This experimentally confirmed superiority is believed to be a combined result of small thickness and near atomic vacancies inherent to CVD-grown 2D MoS₂ layers, verified by STEM characterization and MD simulation. The present study merits further investigations towards the potential exploration of 2D materials for high-efficiency membrane and desalination technologies.

CHAPTER 5: WATER DESALINATION VIA VERTICALLY-ALIGNED 2D MOS₂ LAYERS MEMBRANES

5.1. Introduction

The past decades have seen an escalating global shortage of freshwater resources due to the spiraling rise of water demands from the agricultural and industrial sectors.⁵ Apart from the management and conservation of existing water resources, more environmentally and technologically sustainable solutions that facilitate increasing freshwater supplies are urgently demanded around the globe in mitigating the challenging impacts of the looming water crisis, particularly during the rapidly evolving effects of climate change. Significant technological and research efforts have been devoted towards seawater desalination techniques, due to the abundance and accessibility of oceans and seas, aiming to produce freshwater in a more time and energy efficient manner. 10 Amongst the most widely adopted desalination technologies are the multi-stage flash (MSF), multi-effect distillation (MED), and reverse osmosis (RO), with RO increasingly attaining more market share. 7,8 As a pressure-driven process, the RO polymeric membranes still face unfavourable technical drawbacks in particular low conversion efficiency, and the high energy demand for pressurising the feedwater. Therefore, the development and adoption of high-performance membrane materials still present compelling demands, especially materials armed with an inherently balanced combination of optimal water recovery efficiency and maximised solute rejection rates.

Two-dimensional (2D) materials based membranes have rapidly emerged as a promising candidate for addressing and overcoming the undesirable trade-offs between permeability and selectivity for improved desalination performances. 11-13,149

Their desirable unique suitability stems from a combination of the ultra-thinness of only one-to-few molecular layers which allows for rapid water pathways as well as superior ionic/molecular sieving capabilities due to versatile mechanisms (e.g., in-plane nanopores, interfacial voids, and inter-gallery channels). In addition, many other attributes of 2D materials including their high mechanical flexibility and chemical functionality further qualify them as a favourable alternative towards high-performance desalination. Moreover, the chemical synthetic processing routes can impart intrinsic sieving/filtration nano-channels which can be even further tailored towards desired channel sizes, distribution density, and orientation configurations for optimal desalination capabilities. I4,228,229

In this endeavour, the ionic sieving and water permeation performance of various types of 2D materials have been explored, with the most notable categories of examples include graphene, ^{15-18,230} dichalcogenides, ^{19-22,231} and graphene oxides. ^{23-26,232}

Amongst a host of 2D materials based membranes, 2D molybdenum disulfide (MoS₂) layers attracted newly gained research efforts for their promise in water and environmental fields, ^{149,153} along with their demonstrated superior properties in various other fields of applications including electronics, ²³³ catalysis, ²³⁴ biomedical, ²³⁵ and energy conversion and storage ²³⁶. In addition to their inherent structural advantages, other attributes such as excellent chemical stability and non-toxicity further make 2D MoS₂ layers a fitting candidate for high-performance desalination applications. ³⁵⁻³⁷ Despite their promising potential as an efficient membrane material for water desalination applications, only limited experimental research efforts have been devoted towards the investigation of 2D MoS₂ layers for their ionic sieving and water permeation capabilities, whilst extensive theoretical projections based on molecular dynamics simulations have already revealed their promising outlooks in this

regard. ^{19,149,154,155,237} Recent experimental desalination assessments of 2D MoS₂ flakes/layers based membranes include chemically functionalised ~5 µm thick laminar membranes composed of exfoliated 2D MoS₂ flakes via a filtration process. ¹⁵⁷ Moreover, a recent notable work by Li *et al.* reported the experimental demonstration of high-efficiency water desalination membranes based on horizontally-aligned few-layer 2D MoS₂ of only ~7 nm thickness, attributing the high performance to the small thickness and near atomic vacancies inherent to the CVD-grown 2D MoS₂ layers. ²³⁸

Compared to their horizontally-aligned counterparts, vertically-aligned 2D / nano materials have received even more scarce research efforts for their potential roles in environmental applications, both in terms of theoretical and experimental investigations. Amongst the limited reported results, few explored the theoretical performance of vertical graphene-based membranes, whilst the majority of the efforts have been devoted to vertically-aligned carbon nanotubes (VA-CNTs). ^{229,239-242} In addition, a recent MD simulations work by K öthler *et al.* has suggested that theoretically the transport of water through MoS₂ nanotubes can be as anomalous as in CNTs, potentially have promising applications such as nano-filtration. ²⁴³ In this Chapter we present results and discussions on the superior ionic sieving capability of a vertically-aligned 2D MoS₂ layers based membrane, both the experimentally demonstrated capabilities, with a special attention on the comparisons with its horizontal counterparts ²³⁸ with regards to the differences in performances and the respective underlying mechanisms.

5.2. Experimental Methods

CVD Growth of 2D VA-MoS₂ Layers: Ultra thin films of Mo (6 nm thickness) were deposited onto SiO₂/Si substrates via electron beam evapouration (Thermionics VE-100 evapourator) at a deposition rate of 0.15 Å s⁻¹. As-deposited Mo-SiO₂/Si samples were then centrally-placed inside a 1 inch quartz tube located in the middle of a horizontal tube furnace (Lindberg Blue M, ThermoScientific). Sulphur powders (99.5%, Sigma-Aldrich, St. Louis, MO) placed in a ceramic boat were then loaded within the quartz tube at the furnace upstream side. The furnace was then pumped down to a base pressure of 1 mTorr following purging with high-purity argon (Ar) gas. The tube system was then ramped up to 780 °C at a ramping rate of 15 °C/min for 50 mins and was held for another 50 mins under the constant flow of Ar gas (100 SCCM), with a working pressure of 75 mTorr. After the CVD reaction, the furnace was naturally cooled down to room temperature under continuous Ar supply.

Water Permeation Rate Measurement: A typical external pressure method was used for the water permeation rate assessments via a common dead-end filtration setup. The sandwiched membrane with exposing holes over the vertically-aligned 2D MoS₂ layers was secured inside a leak-free syringe filter holder. The water permeation rate was calculated by weighing the volume of water passing through the 2D VA-MoS₂/PES membrane area under a constant pressure of 1 bar as a function of time.

Salt Ion Rejection Rate Measurement: A conductivity probe (ThermoScientific, model: Orion 013605MD) connected to a solute conductivity meter (ThermoScientific, model: Orion 5 Star) was used to monitor conductivities in respective containers, in order to quantify salt ion concentration variations. The conductivity probe was calibrated on a routine basis, and was thoroughly rinsed with de-ionised water each before measurements. The salt rejection rate was

calculated via $\left(1 - \frac{C_P}{C_F}\right) \times 100\%$, where C_P and C_F are the conductivity of the permeate and feed side, respectively. Common seawater salt types of sodium chloride (NaCl) (99%, Sigma-Aldrich, St. Louis, MO), potassium chloride (KCl) (99%, Sigma-Aldrich, St. Louis, MO), and calcium chloride (CaCl₂) (93%, Sigma-Aldrich, St. Louis, MO) were employed for salt rejection rates measurements. The seawater salt solutions tested were as follows: (1) NaCl of 0.6 M (equivalent to seawater salinity concentration²⁰⁹), (2) NaCl of 0.469 M (typical seawater Na⁺ concentration²¹⁰), (3) KCl of 0.1 M, (4) MgCl₂ of 0.1 M, (5) CaCl₂ of 0.1 M, (6) actual seawater sampled from the Florida Atlantic coast, and (7) synthetic seawater. The concentration of 0.1 M for KCl, MgCl₂ and CaCl₂ was selected for measurements consistency, although the typical seawater concentrations of K⁺, Mg²⁺ and Ca²⁺ are below 0.1 M.^{210,211} The synthetic seawater solution was prepared to mimic the concentrations of typical salt ion species present in standard seawater, i.e., Na⁺ of 0.469 mol/L ²¹⁰, K⁺ of 0.01 mol/L ²¹⁰, Mg²⁺ of 0.0528 mol/L ²¹⁰, and Ca²⁺ of 0.01 mol/L ²¹⁰. Each measurement was repeated for three times to improve testing consistency, with the averaged values presented in the corresponding plots.

AFM Characterisation: Topography images and height profiles of the vertically-aligned 2D MoS₂ layers integrated on PES substrates were acquired with Anasys nanoIR AFM (Anasys Instruments, Santa Barbara, CA) operated in tapping mode with the scan rate of 1 Hz. The AFM probes (Model: PR-EX-T125-10, Anasys Instruments) had a nominal spring constant of 30 N/m.

5.3. Results and Discussions

5.3.1 Vertical 2D MoS2 Layers Preparations and Characterizations

Vertically-aligned (VA) 2D MoS₂ layers of centimetre-scale were grown based on a modified chemical vapour deposition (CVD) process previously developed in our research group. 195-197 A thin film of Mo (~6 nm thickness) was deposited onto a SiO₂/Si substrate via electron beam evaporation, followed by a thermal sulphurisation process inside a CVD chamber, producing VA 2D MoS₂ layers. Conditions for VA 2D MoS₂ layers growth are detailed in *Methods* section. As-prepared VA 2D MoS₂-SiO₂/Si sample following the CVD process was then facilely delaminated via a water-assisted process, with the afloat VA 2D MoS₂ layers gently transferred onto an O₂ plasma treated porous poly(ether sulfone) (PES) membrane substrate (~0.03 µm pore size and ~100 µm thickness, Sterlitech, WA), which has been extensively employed as substrate for assessing desalination performance of 2D materials-based membranes. ¹⁹⁹ The 2D VA-MoS₂/PES surface was then covered with a selectively deposited thin layer of nonpermeable polydimethylsiloxane (PDMS), exposing a pre-defined open region of 2D vertically-aligned MoS₂ layers as the ionic sieving membrane upon solutions contact (Figure 32(a)). The growth, delamination, transfer and integration of 2D layers onto PES substrates followed similar procedures as illustrated on our previous report.²³⁸ With the PES substrate possessing pores greatly exceeding the sizes of salt ion species, the ionic sieving effects can only be attributed to the 2D VA-MoS₂ layers (with the possible mechanism shown schematically in Figure 32(b) and discussed in detail in latter sections of this Chapter).

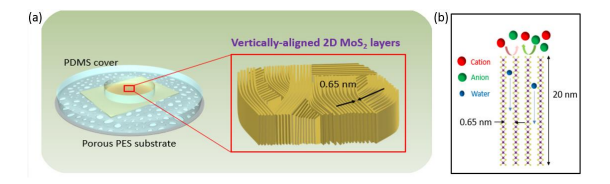


Figure 32 (a). Schematic depiction of 2D VA-MoS₂ layers integrated onto porous PES substrate. A non-permeable PDMS layer is selectively deposited on the MoS₂/PES, leaving an opened area allowing water/solution penetration through the exposed 2D VA-MoS₂ layers. The enlarged view of the red rectangle area shows the vertical alignment of the 2D VA-MoS₂ layers, note the vdW gaps of ~0.65 nm as indicated by the arrows. (b). Schematic illustration of proposed mechanism for the water permeation and salt rejection of 2D VA-MoS₂ layers. The inherent vdW gaps of the vertically-aligned MoS₂ layers pose restrictions towards inbound salt ions mainly due to size exclusions. Whilst the hydrophobic interior regions between the MoS₂ layers facilitate smooth transport of water molecules.

Prior to the integration of VA 2D MoS₂ layers onto PES substrates, the material quality of asgrown 2D MoS₂ layers was first assessed. The structural and chemical integrity of the CVD-grown VA 2D MoS₂ layers were characterised via transmission electron microscopy (TEM) and Raman spectroscopy. A plane-view of the as-grown 2D MoS₂ layers via HR-TEM clearly demonstrates the vertical alignment of 2D MoS₂ layers (Figure 33(a)), with the inset showing a typical interlayer spacing of ~0.65 nm between the adjacent MoS₂ layers. In addition, a cross-sectional view of the 2D MoS₂ layers via TEM (Figure 33(b)) reveals the presence of continuous 2D MoS₂ layers that are of uniform thickness. Shown as an enlarged view of the blue box in Figure 33(b), the crystalline structure of 2D MoS₂ layers is further revealed in

Figure 33(c) with the corresponding atomic arrangements of the 2D MoS₂ layers also illustrated as in the red boxes, which further clarifies the vertical orientation of the 2D MoS₂ layers, and again confirms an average interlayer spacing of ~0.65 nm. Furthermore, the typical Raman spectrum of the as-grown MoS₂ layers (red spectrum line in Figure 33(d)) confirms the presence of 2D MoS₂ layers, with the two distinctive peaks corresponding to the in-plane (E_{2g}^1) and out-of-plane (A_{1g}) vibration modes of 2D MoS₂ layers, respectively. These comprehensive characterisation results conclude that our CVD-grown 2D MoS₂ possess continuous and vertically-aligned layers, with multitudes of intrinsic sub-nanometre vertical channels (e.g., ~0.65 nm vdW gaps). Such microstructural attributes impart VA 2D MoS₂ layers with unique suitability as desalination membranes.

Subsequent morphological characterisations of the as-integrated VA 2D MoS₂ layers on PES substrates were performed via atomic force microscopy (AFM) and Raman spectroscopy. The AFM height profile (Figure 33(e)) corresponding to the yellow line across the VA 2D MoS₂ / PES interface (Figure 33(e) inset) shows that the integrated 2D VA-MoS₂ layers are ~20 nm thick (which is within the reasonable range as produced from a 6 nm Mo seed layer). The Raman profile of the as-integrated VA 2D MoS₂ layers / PES (blue spectrum line in Figure 33(d)) reveals the resembling characteristic peaks of E_{2g}^1 and A_{1g} vibration modes, which are consistent with the observation prior to 2D layers separation and integration (i.e., near-overlapping red spectrum line in Figure 33(d)). The results confirm that after the VA 2D MoS₂ layers integration onto PES substrates, they preserved a maintained structural and chemical integrity.

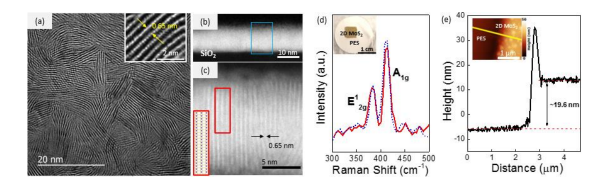


Figure 33 (a). Plane view of the HR-TEM imaging of the smooth 2D VA-MoS₂ layers, clearly showing the vertical alignment of the 2D MoS₂ layers. The inset shows a zoomed-in view of the vertical layers with well-defined vdW gaps of ~0.65 nm as indicated by the yellow arrows. (b). A cross-sectional view of the 2D VA-MoS₂ layers, showing continuous 2D VA-MoS₂ layers with uniform thickness. (c). An enlarged view of the 2D VA-MoS₂ layers as shown in the blue rectangle area in Figure 33(b), showing the vertically-aligned nano-channels formed in between the parallel 2D MoS₂ layers, again with a well-defined vdW gaps distance of ~0.65 nm (indicated by the black arrows). Atomic schematic representation of the vertical MoS₂ nano-channels is also depicted (see the two corresponding areas in the red rectangles). (d). Typical Raman spectra of the 2D VA-MoS₂ layers before (red line) and after (blue line) the PES integration, showing the two distinctive peaks corresponding to the E_{2g}^1 and A_{1g} vibration modes. The inset picture shows a representative sample after the 2D VA-MoS₂ layers (dark gold area) have been integrated onto the PES substrate (white area). (e). Representative AFM height profile of 2D VA-MoS₂ layers integrated onto the PES substrate, here showing a thickness of approximately 20 nm. The height profile was taken along the gold-coloured line on the inset AFM imaging, across the boundary between the PES substrate and the 2D VA-MoS₂ layers.

5.3.2 Water Permeation Assessments

Figure 34(a) illustrates a schematic set-up for water permeation measurements. The 2D VA-MoS₂ layers-integrated PES substrate was covered with another bare PES substrate. After the 2D layers integration, the sample was subsequently sandwiched between two non-permeable PDMS sealing films both exposing pre-defined open areas overlapping with the 2D VA-MoS₂ layers membrane region. Note that this sandwich-like assembly has been extensively employed in configuring nanomaterials-based membranes including 2D materials of large thickness (> 1 μm). ¹⁵⁷ This layered membrane was then loaded inside a syringe filter holder in a typical deadend filtration setup. The water permeation rate was then measured via common external pressure method by weighing the volume of water passing through the membrane area (i.e., the central hole filtration region composed of 2D VA-MoS₂/PES only) as a function of time under a constant pressure. Details are presented in the *Methods* section.

Due to the vertical nature of the 2D MoS₂ layers configuration in our case, the pathways that allow passage for water molecules are treated as parallel continuous nano-channels, which should account for the vast majority of water permeation, with some types of defects (e.g., edge sites and dislocations) may only have very insignificant contributions.

Before proceeding with the experimental assessments of the water permeation capability of 2D VA-MoS₂ layers, it is worth exploring the theoretical performance of water permeability of a single nano-channel of vertical 2D MoS₂ layers (and this *per nano-channel* estimation will be further extended to the *per unit area* value).

Since the "openings" for water molecules to penetrate upon contact on the surfaces of 2D VA-MoS₂ layers do not present themselves as geographical "holes", (instead, there are parallel "slits" composed of adjacent nano-channels). Therefore, for the sake of reasonable and

simplified estimations, the preferable "opening" for water molecules permeation is considered as the area defined by the following two dimensional factors: (1). the width of each of the parallel "slits" d=0.333 nm 244 , and (2). the Sulphur-to-Sulphur distance within a single 2D MoS₂ layer d_S =0.316 nm 245 .

The theoretical water permeation capability of 2D VA-MoS₂ layers is based on the Hagen-Poiseuille (H-P) permeability, which describes the water flux under ideal and non-slip flow conditions through cylindrical pores, and is widely adopted as a base for theoretical estimations.^{239,246,247}

The theoretical H-P permeability for vertical 2D MoS₂ layers maybe calculated via the following equation:^{239,247}

$$Q = \frac{\pi (\frac{d}{2})^4}{8\mu} \cdot \frac{\Delta P}{L} \tag{2}$$

Where

Q =The water flux = the volume of water passage per unit time, per unit pressure, per nanoopening

d =Size of nano-opening

 μ = Viscosity of de-ionised water (at 20 °C) ²⁴⁸

 ΔP = Trans-membrane pressure

L =Nano-opening length

Therefore, from Equation. 2, the theoretical H-P permeability is calculated as:

$$Q = 5.54 \times 10^{-18} \frac{L}{h \cdot bar \cdot opening}$$

Furthermore, based on the dimensional factors (d and d_S) mentioned above, the number of nano-openings per 1 cm² for 2D VA-MoS₂ layers can be estimated as $4.88 \times 10^{14} \frac{opening}{cm^2}$, which is the areal density. Note this estimation is at least 50 times more compared to that of the horizontally-aligned CVD-grown 2D MoS₂ layers.⁴¹

Therefore, the theoretical estimation of the Hagen–Poiseuille permeability of a typical 2D VA-MoS₂ layers sample (under ideal, non-slip conditions) can be deduced by combining Equation. 2 and the estimated areal density, which gives:

$$Q_{H-P} = 27 \; \frac{L}{h \cdot bar \cdot m^2}$$

For comparisons, the reported theoretical Hagen–Poiseuille water permeability of vertical CNTs varies in a wide range: from $Q = 0.0053 \frac{L}{h \cdot bar \cdot m^2}$ to $Q = 3.35 \frac{L}{h \cdot bar \cdot m^2}$ (due to varied sample parameters like pore densities and sizes).²⁴⁷

The experimentally assessed water permeation rate, based on the common dead-end filtration method, for 2D VA-MoS₂ layers is $Q = 1656.8 \ (\pm 129.7) \ \frac{L}{h \cdot har \cdot m^2}$.

The prominent difference of the water permeations rates between the theoretical Hagen–Poiseuille permeability and the experimentally assessed values indicate that the vertical 2D MoS₂ layers possess highly slippery nano-channels for water molecules. Comparable and sometimes even more dramatic enhancement effects on water permeability have been reported on vertical CNTs membranes.^{247,249-251}

We believe the hydrophobicity of the vertical 2D MoS₂ layers would most likely account for such contrasting differences. For comparison, the underlying mechanism behind the significant enhancement effects for vertical CNTs has been similarly attributed to hydrophobic—hydrophobic interaction, stemming from the hydrophobicity of the inner wall of CNTs.^{239,252} Furthermore, theoretical MD simulations study also supports the presence of an induced superflow regime inside MoS₂ nanotubes, leading to high water mobility.²⁴³

Comparing to the their horizontally-aligned counterparts, vertical 2D MoS₂ layers clearly show a higher water permeability (see the comparison plot in Figure 34(b)).²³⁸ Such contrast is ascribed to the different pathways that water molecules take. In the case of horizontal-MoS₂ 2D layers, water translocates through the nanopores as well as transports within the interlayer vdW gaps.²³⁸ Whilst in 2D VA-MoS₂ layers, three main factors are at play contributing to the enhanced water permeability: 1). the available passages for water molecules become more abundant, due to the presence of the numerous "slits" like openings (i.e., the vdW gaps between layers of vertical MoS₂); 2). the vertically-aligned layers present much more straightforward "straight-to-the-end" passages, compared to the "zig-zag" routes that water molecules take when migrate through the horizontal MoS₂ layers; and 3). the hydrophobicity of the interior regions between the vertical MoS₂ layers further leads to rather "slippery" pathways for water molecules to move more smoothly.

5.3.3 Salt Rejection Evaluations

In addition to the superior water permeability capability discussed above, the salt rejection performances of 2D VA-MoS₂ layers were also assessed.

The proficiency of 2D VA-MoS₂ layers on PES supports for salt ions rejection was determined via the bespoke measurement set-up (depicted in Figure 34(c)) as described in detail in our previous report, following similar procedures and testing conditions.²³⁸ In brief, a leak-free solution passage via only the sandwiched 2D VA-MoS₂ layers connects the permeate and feed side containing de-ionised water and different types of testing salt solutions respectively. The levels of the migration of salt ions were quantified via the conductivity measurement method well established in the field;¹⁵⁷ i.e., the salt rejection rate was calculated by $\left(1 - \frac{C_P}{C_F}\right) \times 100\%$, where C_P and C_F are the conductivity of the permeate and feed side, respectively.²⁰⁸ Measurement procedures, conditions as well as the different types of salt solutions underwent testing are detailed in the *Methods* section. Figure 34(d) presents the plots of seawater ion rejection rates obtained at identical interval times during the course of three hours towards various types of respective salt ion species, whilst the plots for multi-ionic mixture solutions are given in Figure 34(e).

The results (as given in Figure 34(d) and Figure 34(e)) show a high level of capability for 2D VA-MoS₂ layers acting as reliable hindrance to the passage of common seawater salt ion species at various concentrations, for both single-ionic and multi-ion mixture solutions (with an overall salt rejection rate of > 99.7%).

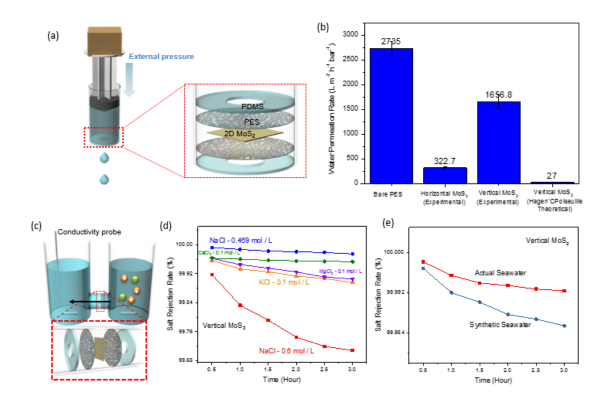


Figure 34 (a). Schematic depiction of the water permeation measurement set-up. The enlarged view shows the sandwiched membrane composed of 2D VA-MoS₂ layers integrated on PES substrate, with another bare PES covered on top of them. The selectively-deposited PDMS regions are also shown with the circular areas left open for water/solution penetration. (b). The comparisons of water permeation rates for: 1). bare PES substrates, 2). horizontal 2D MoS₂ layers, ²³⁸ 3). vertically-aligned 2D MoS₂ layers, and 4). the theoretical Hagen-Poiseuille permeability of 2D VA-MoS₂ layers. (c). Schematic representation of the salt rejection rates measurement set-up. The sandwiched membrane composed of 2D VA-MoS₂ layers / PES, with selective PDMS sealing is shown in the enlarged view. (d). Salt rejection rates of 2D VA-MoS₂ layers towards various types of seawater ions over a continuous 3-hr time period. (e). Salt rejection rates of 2D VA-MoS₂ layers towards synthetic and actual seawater samples over a continuous 3-hr time period.

Due to the similarity of assessment methodology and testing conditions, the following discussions focus on the salt ions rejection performance of 2D VA-MoS₂ layers as compared to their horizontal counterparts, in addition to comparisons with vertical-CNTs.

It is found that across all the different types of ionic solutions, the vertically-aligned 2D MoS₂ layers show a slight yet consistently higher performance in terms of salt rejection rates under similar testing conditions compared to the horizontally-aligned 2D MoS₂ layers²³⁸ (representative optical imaging of 2D H-MoS₂/PES and TEM imaging of the horizontal layers are shown in Figure 35(a)). The respective salt rejection performance comparison plots are given in Figure 35(b)-(f) for single-ionic solutions, and in Figure 35 (g)-(h) for multi-ionic mixture solutions. This improvement of performance may due to two possible reasons (also illustrated in Figure 32(b)): 1). Compared to H-MoS₂ layers, which possess nano-pores of varying degrees of sizes, the VA-MoS₂ layers instead contain filtration networks of nearconsistent dimensions (i.e., tiny "slits" of approximately 0.65 nm in width. See Figure 33(a) and Figure 33(c)). Such narrow-sized vdW gaps "slits" are well-suited for allowing the smooth and even maybe accelerated passage for water molecules, which have a diagonal length of ~0.275 nm.²⁰³ And at the same time, the hydrated salt ions face both dimensional exclusions upon encountering the narrow "slits" due to their larger sizes (also note the actual Sulphur-to-Sulphur distance of ~0.3 nm.²⁴⁴), and the energetic impedance of having to lose their solvation shells if they are to squeeze through. 225,226 Additionally 2). The thickness of 2D VA-MoS₂ layers are inherently higher than that of the H-MoS2 layers, since the growth of vertical 2D MoS₂ layers prefers thicker (>~3 nm) Mo-seed layers. ¹⁸⁵ Thus possessing lengthened barriers preventing the salt ions from passing through.

Despite many MD simulations works^{253,254} as well as extensive experimental investigations^{229,247} on the potential of VA-CNTs membranes for desalination applications, their practical feasibility has been limited by the actual pore sizes which remain larger than 1 nm ^{213,239,247}, greater than the hydrated radius of typical salt ions (e.g., Na⁺ of 0.72 nm). ^{225,239} It is worth noting that extensive molecular dynamics simulations have shown that the inner pore diameter of VA-CNTs should reach ~0.6 nm for optimal salt rejection rates. 240,253,254 Although the present technologies have the capability to reduce the pore diameter of VA-CNTs to this range, a critical pore size of ~0.7 nm would pose as a limiting factor, beyond which the transport of water will be impeded due to the confinement effect. 240,253,255 Therefore, although VA-CNTs have demonstrated exceptionally high experimental water permeability (e.g., up to $10,500 \frac{L}{h \cdot bar \cdot m^2}$) ²⁴⁷, their practical prospect as salt rejection membranes remains uninviting. 239,240,256,257 For example, molecular modelling indicates that even with a pore diameter of 0.75 nm for VA-CNTs membranes, they would only yield a 58% of salt rejection capacity. 253,254 Furthermore, experimental investigations of the salt rejection capacity of VA-CNTs membranes show that with an average pore size of 1.6 nm, only up to 60% ion exclusion efficiency was observed.²⁵⁸ Therefore, alternative to seawater desalination, vertical CNTs have instead found promising application potential as filtration membranes towards larger-sized species like heavy hydrocarbons and microbial contaminants. 256,259

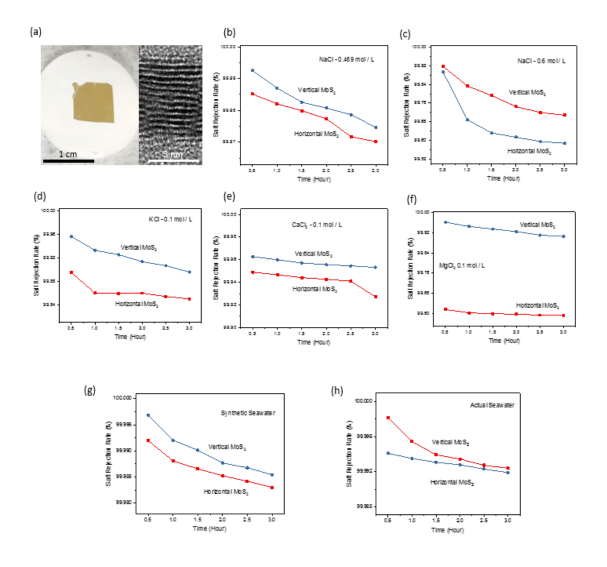


Figure 35 (a). Representative optical imaging of the 2D horizontal-MoS₂ layers integrated onto PES substrate. Along with a top-view HR-TEM imaging showing the horizontally-aligned 2D MoS₂ layers. (b). Comparison plots of the respective salt rejection rates for 2D vertically-aligned MoS₂ layers and 2D horizontally-aligned MoS₂ layers towards 0.469 mol/L NaCl solution over a continuous 3-hr time period under similar measurement conditions. (c). Comparison plots of the respective salt rejection rates for 2D vertically-aligned MoS₂ layers and 2D horizontally-aligned MoS₂ layers towards 0.6 mol/L NaCl solution over a continuous 3-hr time period under similar measurement conditions. (d). Comparison plots of the respective salt rejection rates for 2D vertically-aligned MoS₂ layers and 2D horizontally-

aligned MoS₂ layers towards 0.1 mol/L KCl solution over a continuous 3-hr time period under similar measurement conditions. (e). Comparison plots of the respective salt rejection rates for 2D vertically-aligned MoS₂ layers and 2D horizontally-aligned MoS₂ layers towards 0.1 mol/L CaCl₂ solution over a continuous 3-hr time period under similar measurement conditions. (f). Comparison plots of the respective salt rejection rates for 2D vertically-aligned MoS₂ layers and 2D horizontally-aligned MoS₂ layers towards 0.1 mol/L MgCl₂ solution over a continuous 3-hr time period under similar measurement conditions. (g). Comparison plots of the respective salt rejection rates for 2D vertically-aligned MoS₂ layers and 2D horizontally-aligned MoS₂ layers towards synthetic seawater over a continuous 3-hr time period under similar measurement conditions. (h). Comparison plots of the respective salt rejection rates for 2D vertically-aligned MoS₂ layers and 2D horizontally-aligned MoS₂ layers towards actual seawater over a continuous 3-hr time period under similar measurement conditions.

Comparisons of water permeability of 2D VA-MoS₂ layers (the present work) with reported performances of VA-CNTs and commercial ultrafiltration (UF) are given in Table 3 Comparisons of membrane specifications and water permeability., along with the corresponding materials properties of the testing samples. (Note: Amongst the reported results for VA-CNTs, there are vast performance differences because of inherent variations of sample parameters like pore densities and sizes).

Table 3 Comparisons of membrane specifications and water permeability.

Membrane specification s	This	Lee et al. ²⁴⁷	Yu et al. ²⁵⁰	Zhan g et al. ²⁶⁰	Baek et al. ²³⁹	Hind s et al. ²⁴⁹	Holt et al. ²⁴⁶	Du et al. ²⁴¹	UF 247
(CNT) inner diameter,	N/A	4.1	3.0	10	4.8	7.5	1.6	10	5.7
Thickness,	19.6	10 ⁶	7.5×1 0^5	$\begin{array}{c} 1.2 \times 1 \\ 0^5 \end{array}$	2×1 0 ⁵	5×1 0^3	2.5×1 0^3	0^6	2×1 0^4
Pore density, 10^{10}pore $/\text{cm}^2$	4.88×1 0 ⁴	300	290	10	6.8	6	25	2.4	9
Water permeability at 20 ℃, L m ⁻² h ⁻¹ bar ⁻¹	1,657	10,50	2,740	1,913	1,10	606	267	234	32
Calculated water	270,00	677	276	7,348	159	33,4 81	578	53	4,09 6

permeability					
at 20 ℃,					
$\times 10^{-4} \mathrm{L}\mathrm{m}^{-2}$					
h ⁻¹ bar ⁻¹					

5.4 Summary

In summary, we experimentally evaluated the performances of vertically-aligned CVD-grown 2D MoS₂ layers (approximately 20 nm in thickness) as water permeable membranes for desalination and ionic sieving. This ultra-thin membrane presented an excellent combination of high water permeability and high ionic sieving capability for a variety of tested ions, which with improved performances compared to horizontally-aligned 2D MoS₂ layer layers membranes. This experimentally confirmed superiority is believed to be a combined result of small thickness and the hydrophobic interiors regions of the vertical MoS₂ layer sheets. The present study merits further investigations towards the potential exploration of 2D materials for high-efficiency membrane and desalination technologies.

CHAPTER 6: CONCLUSIONS

In summary of the present dissertation, the behavior of the internal potential barrier in a polymer-derived amorphous SiAlCN ceramic was studied by measuring its complex impedance spectra at various dc bias as well as different testing and annealing temperatures. The complex impedance spectra of the polymer-derived a-SiAlCN were measured under various dc bias voltages in a temperature range between 50 and 150 °C, as well as different annealing temperatures (1100-1400 °C). All spectra, regardless of temperature and bias, consist of two semi-circular arcs, corresponding to the free-carbon phase and the interface, respectively. The impedance of the free-carbon phase is independent of the bias, while that of the interface decreased significantly with increasing dc bias. It is shown that the change of the interfacial capacitance with the bias can be explained using the double Schottky barrier model. The charge-carrier concentration and potential barrier height were estimated by comparing the experimental data and the model. The results revealed that increasing testing temperature led to an increased charge-carrier concentration and a reduced barrier height, both following Arrhenius dependence, whereas the increase in annealing temperature resulted in increased charge-carrier concentration and barrier height. The phenomena were explained in terms of the unique bi-phasic microstructures of the material. The research findings reveal valuable information of temperature-dependent properties of polymer derived ceramics, and should contribute towards more precise understanding and control of the electrical as well as dielectric properties of polymer derived ceramics.

Furthermore, the desalination performances and underlying mechanisms of two-dimensional CVD-grown MoS₂ layers membranes have been experimentally assessed. Based on a successful large-area few-layer 2D materials growth, transfer and integration method, the 2D

MoS₂ layers membranes showed preserved chemical and microstructural integrity after integration. We experimentally evaluated the performances of CVD-grown few-layer 2D MoS₂ of ~7 nm thickness as water permeable membranes for desalination and molecular sieving. This near atom-thick membrane presented an excellent combination of high water permeability and high ionic sieving capability for a variety of tested ions, outperforming previously developed 2D MoS₂ layer membranes of much larger thickness. Such performances are attributed to the dimensional and geometrical effect, as well as the electrostatic interaction of the inherently-present CVD-induced atomic vacancies (as verified by STEM characterization and MD simulation) for governing both water permeation and ionic sieving at the solution/2D-layer interfaces. The present study merits further investigations towards the potential exploration of 2D materials for high-efficiency membrane and desalination technologies.

CHAPTER 7: FUTURE WORK

The interfaces of nanomaterials are integral parts for defining the properties and performances of many types of materials. Therefore, the investigations of the roles of interfaces should also be extended to the studies of inherent properties of their bordering phases as well as the influences of the externally applied fields. The following suggestions are proposed for aiding future research efforts.

(a).

The eventual aims of the investigations on the electrical and dielectric properties of polymer derived ceramics is multi-fold: (1) to understand the origin and fundamental mechanism of conductions; (2) to decipher the microstructure-processing-properties relationships (i.e., how the different parameters / conditions / techniques would influence its electrical / dielectric performance); (3) to gain in-depth knowledge of better controlling / targeting its structures and properties towards different and wider applications (e.g., harsh environment sensing, and environmental barrier coatings). There are a number of factors that could influence its eventual performance, for example: (a) the ratio of initial ingredients; (b) the processing temperatures; (c) temperatures under which the PDC materials operate; and (d) contaminations (especially oxygen species) introduced during processing steps. Based on the results and discussions of the present investigations, in order to continue to build a more thorough understanding of controlling the electrical and dielectric properties of PDCs (for scientific research as well as practical applications purposes), the following further recommendations are provided as a suggestive guideline:

(1). Since the PDCs materials are particularly notable for their high-temperature-stable properties as well as promising future applications under harsh environments, it would be empirical to investigate the performance of PDCs (in particular their electronic / dielectric properties) under in-service conditions / environments. Though many research works have demonstrated the structural stability of PDCs, its long-term influence over PDCs microstructures and properties still demand in-depth efforts before the materials render practically useful. In addition, impedance measurements of even wider ranges should be investigated as temperatures could have shifted peaks towards higher ends.

One additional aspect of a future research direction worth pursuing of is to study the synergistic effects of the material's responses to both temperatures and external compressive pressures. It was found / observed that the polymer derived SiAlCN ceramics, at the same time, show prominent responses to varying degrees of temperatures and pressure. This effect could undermine its future applications for harsh environment sensing, since the actual operating environment of such sensors would have both rapidly-changing temperatures and pressures present. Since these two effects arise from different origins, it truly worth more further explorations on the improved control of this effect. It is suggested that one approach to reduce such effect would be to study how the processing / annealing route would affect such synergistic effect. And another more resistive path would be to combine electrical engineering knowledge via clever design of circuits to minimise the effect, which would normally require product shape-control technique during processing.

(2). To study the effect of microstructures on the double Schottky barriers model in controlling PDC dielectric properties. Especially the electronic band / energy structures of PDCs with different annealing / pyrolysis histories. The presence of varying levels of defects in different

PDCs dictates the relative complexity of determining their band structure. However, various experimental microstructural characterisation techniques should be useful for identifying the electronic structures of PDCs. Examples include: optical absorption measurement, X-ray photoelectron spectroscopy (for bond information). They could provide critical information on the microstructural evolutions and assist the evaluations of how various factors may affect the relative permittivity as well as dielectric loss of polymer derived ceramics, which are crucial indicators of the practical applicability of these materials for future applications.

(3) The aims of studying the electronic as well as the dielectric properties of PDCs are to have more precise and dependable controls for practical sensing applications. It is therefore imperative to conduct further investigations on the integration and packaging processes of the sensing materials (i.e., PDCs) into an integral system. The reliability and accuracy of the sensing system first depends on the innate microstructures of the PDC sensing material. From a holistic scientific point of view, the materials processing will have fundamental effects on the performance of PDCs, which requires further investigations as well as future explorations of the applicability of novel processing techniques.

(b).

The filtration / selective separation properties of 2D layers-based membranes are just beginning to gain substantial research attentions, and the practical applications are still in its infancy. Significant endeavors and discoveries await in developing device-level / large-scale applications of them.

(1). As a fundamental aspect of research and development, viable methods for the growth, transfer and integration of large-area 2D membranes are vital for the implementations of 2D

layers-based desalination devices / equipment. Most importantly, the long-term durability as well as defect-free integration and operations still require extensive investigations.

- (2). As a novel type of nanomaterials, the long-term immunological bio-compatibility, toxicology, and bio-persistence of 2D materials shall be established via intensive research efforts to ensure the health aspects are addressed properly.
- (3). Another potential avenue for future investigations would be the evaluation of controlled pore size generations to produce nanopores with desired sizes to meet the needs of selective filtrations. Detailed studies are required to fine-tune the operational procedures and parameters for optimal nanopore size distributions.

APPENDIX: COPYRIGHT PERMISSIONS

CHAPTER 3

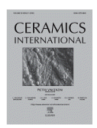












Title: Effect of temperature on internal

potential barriers in polymerderived amorphous ceramics

Hao Li,Baisheng Ma,Linan An

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CHAPTER 4





Author:







Experimental Realization of Few Layer Two-Dimensional MoS2

Membranes of Near Atomic Thickness for High Efficiency Water Desalination

Hao Li, Tae-Jun Ko, Myeongsang

Lee, et al

Publication: Nano Letters

Publisher: American Chemical Society

Aug 1, 2019

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Figure 2 and Figure 4

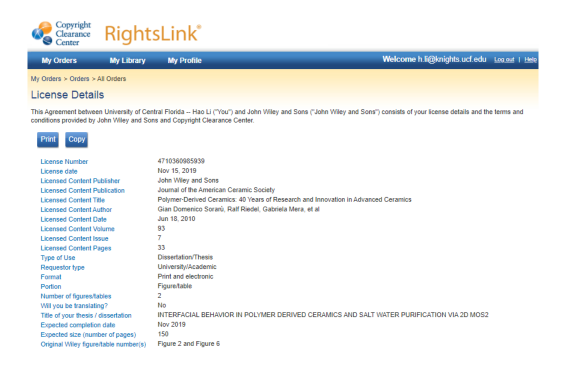


Figure 6, Figure 7, Figure 8, and Figure 9

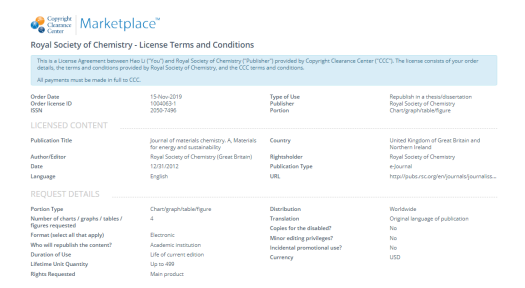


Figure 10 and Figure 11

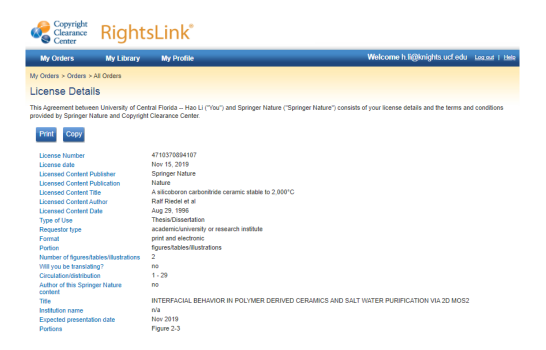


Figure 12

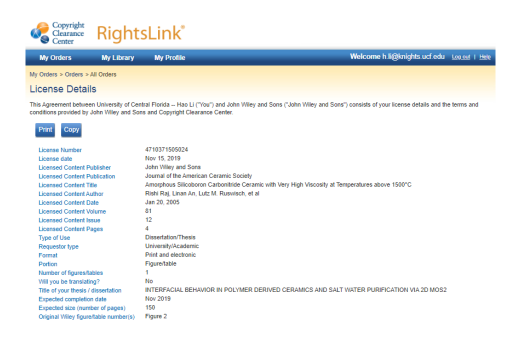


Figure 13 and Figure 14

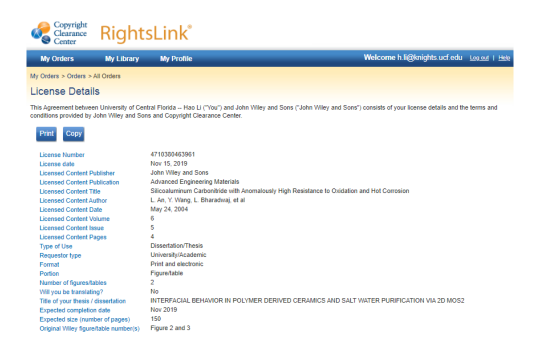
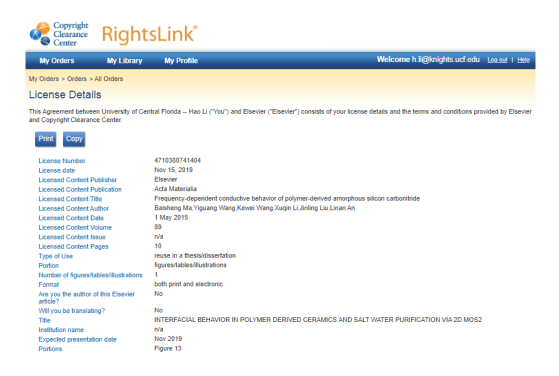


Figure 15



LIST OF REFERENCES

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