Feasibility of fiber optic sensors in sensing high refractive index for the potential application of acquiring solubility and diffusivity of gases and supercritical fluids in polymers

by

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Bachelor of Engineering, University of Victoria, 2014

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

MASTER OF APPLIED SCIENCE

in the Department of Mechanical Engineering

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University of Victoria

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Abstract

Many properties of polymers can be affected by dissolving gases and supercritical fluids at high temperatures and pressures. Solubility and diffusivity are crucial parameters in polymer processing applications that indicates the content of gases and supercritical fluids in a polymer. Hence, different devices for measuring solubility and diffusivity have been researched, but most of the devices used today are very complex, expensive, and requires long experiment time. In this final thesis, the feasibility of fiber optic sensors as measurement devices for solubility and diffusivity of gas/SCF in polymers have been investigated. Many of the polymers used in polymer processing have high refractive index, from 1.40 to 1.60. However, most of the refractive index sensors based on fiber optics only operate in refractive index ranges of 1 to 1.44 because once the surrounding refractive index becomes greater than that of cladding, the total internal reflection is lost and only small portion of the light propagation occurs. This final thesis first reviews the current methods to measure solubility and diffusivity of gases and supercritical fluids in polymers. In addition, different types of fiber optics sensors used for sensing the refractive index are reviewed. Then, the thesis presents cost efficient, but effective fiber optic refractive index sensors, which are the silver nanoparticle coated LPG sensor, uncoated PCF MZI sensor, silver nanoparticle PCF MZI sensor, and the transmission intensity based gap sensor, to sense the surrounding refractive index in the region greater
than the cladding, for the future application of solubility and diffusivity measurement. Moreover, future works that would help in sensing solubility and diffusivity of gas in polymers are also proposed.
# Table of Contents

Supervisory Committee ................................................................. ii  
Abstract ......................................................................................... iii  
Table of Contents .............................................................................. v  
List of Tables .................................................................................. vii  
List of Figures ............................................................................... viii  
Acknowledgments .......................................................................... xii  

Chapter 1.  Introduction ................................................................. 1  
1.1.  Solubility and diffusivity of gas/SCF in a polymer .................. 3  
1.2.  Fiber optics sensors ............................................................... 4  
1.3.  Significance of the problem .................................................. 6  
1.4.  Fiber optic refractive index sensors in refractive index greater than the silica .. 8  
1.5.  Research scope and outline ..................................................... 9  

Chapter 2.  Theoretical background and literature review .............. 11  
2.1.  Methods for measuring solubility of gas/SCF in polymers ......... 11  
2.1.1.  Gravimetric methods ......................................................... 11  
2.1.2.  Piezoelectric methods ....................................................... 19  
2.1.3.  Mamomietric methods ...................................................... 21  
2.2.  Methods for measuring diffusivity of gas/SCF in polymers..... 25  
2.2.1.  Gravimetric method ........................................................... 25  
2.2.2.  Pressure decay method ....................................................... 26  
2.3.  In fiber gratings based refractive index sensors ................. 27  
2.3.1.  Fiber Bragg grating based refractive index sensors .......... 27  
2.3.2.  Long period grating based refractive index sensors .......... 36  
2.4.  Mach-Zehnder Interferometer as refractive index sensor ....... 42  
2.5.  Intensity based fiber optic sensor as refractive index sensor .. 46  
2.6.  High refractive index sensors .............................................. 51  
2.7.  Conclusion .......................................................................... 53  

Chapter 3.  Long period grating sensor for measuring high surrounding refractive index solution 55  
3.1.  Introduction to long period grating sensor .......................... 55  
3.2.  Long period grating sensor in high refractive index .......... 58  
3.3.  Coated Long period grating sensor ...................................... 60  
3.4.  Fabrication of long period grating sensor using the femtosecond laser .......... 61  
3.5.  Silver nano particle deposition using the atomization based coating system ...... 63  
3.6.  Experiment setup ................................................................. 64  
3.7.  Behavior of silver nanoparticle coated long period grating sensor at high refractive index...... 65  
3.8.  Conclusion ........................................................................... 68  

Chapter 4.  Coated Mach-Zehnder Interferometer for measuring high surrounding refractive index solution .......................... 70  
4.1  Mach-Zehnder interferometer sensor .................................... 70  
4.2.  PCF MZI sensor in high refractive index ............................. 76  
4.3.  Effect of coating on PCF MZI ............................................... 80  
4.4.  Fabrication of MZI refractive index sensors ...................... 81  
4.5.  Experiment setup ................................................................. 82
4.6. 40 um MZI at high index ................................................................. 83
4.7. PCF MZI at high surrounding refractive index ................................ 85
4.8. Effect of PCF length in leaky configuration ........................................ 88
4.9. Coated PCF MZI at high surrounding refractive index ...................... 91
4.10. Conclusion ....................................................................................... 97

Chapter 5. Transmission intensity based gap sensor for measuring high surrounding refractive index solution ................................................................. 98
5.1. Introduction ....................................................................................... 98
5.2. Fabrication of the gap sensor ............................................................. 101
5.3. Gap sensor at high surrounding refractive index ............................... 102
5.4. Gap distance parameterization ........................................................... 105
5.5. Polymer testing .................................................................................. 107
5.6. Carbon dioxide desorption in a polycarbonate film ............................ 112
5.7. Conclusion ....................................................................................... 115

Chapter 6. Conclusion and future work ...................................................... 116
6.1. Conclusion ....................................................................................... 116
6.2. Future work ..................................................................................... 118

Bibliography ............................................................................................ 121
Appendix A – Theoretical approaches for solubility study ......................... 135
A.1. Henry’s law ..................................................................................... 136
A.2. Flory-Huggins Theory (Cell Model) ................................................. 136
A.3. Sanchez-Lacombe Theory (Lattice Fluid Model) ................................ 137
A.4. Simha-Somcynsky Theory (Hole Model) .......................................... 138
A.5. Statistical Association Fluid Theory (SAFT) ..................................... 139
Appendix B – Theoretical approaches for diffusivity study ...................... 141
B.1. Steady Diffusion .............................................................................. 141
B.2. Unsteady Diffusion ......................................................................... 142
List of Tables

Table 1-1 Refractive indices of polymers prevalently used in polymer processing........ 7
Table 5-1. Properties of the polymer films ................................................................. 111
List of Figures

Figure 1-1. Schematic of pressure-temperature phase diagram where the triple and critical points are shown. The phase regions are labeled in the diagram................................................. 4
Figure 1-2. Simulated data of coupling wavelength shift against surrounding refractive index on a wide range [27]........................................................................................................ 7
Figure 2-1. Schematic of a McBain Balance in superposed thermostats: (A) heating coil, (B) quartz spring, (C) polymer specimen [42]................................................................. 14
Figure 2-2. High pressure sorption apparatus with a Cahn model 2000 electronic microbalance by Kamiya [56]: (A) pressure chamber containing an electronic microbalance, (B) thermostatically controlled air bath, (C) temperature controller, (D) thermometer, (E) pressure gauge, (F) safety valve, (G) flow meter, (H) columns of active charcoal and silica gel, (I) pressure regulator, and (J) a gas cylinder [56].................. 16
Figure 2-3. Sorption apparatus with a Sartorius S3D-P electro microbalance by Kamiya [58]: (A) Sartorius microbalance, (B) constant temperature water bath, (C) auxiliary furnaces, (D) constant temperature air-bath, (E) pressure chambers, (F) thermocouple, (G) water inlet, (H) gas inlet, (I) polymer sample plus buoyancy corrective (gold wire), (J) counter balance (aluminum rod) [58].................................................. 16
Figure 2-4. The schematic of a solubility measurement apparatus with a MSB by Sato [69].......................................................................................................................... 18
Figure 2-5. Schematic of a sorption apparatus with a QCM used by Miura et al. [81]... 21
Figure 2-6. Schematic of a sorption apparatus with dual sorption pressure decay used by Stern [105]: (A) Auxiliary glass system for volume calibrations, (A1) reference volume, (A2) thermocouple gauge, (A3) mercurial manometer, (B, C, D, E) calibrated volumes of apparatus, (D) gas reservoir, (E) gas absorption cell with polymer sample, (F) differential pressure null indicator, (G) null indicator control, (H) dead-weight gauge, (1-7) needle valves [105]........................................................................................................ 24
Figure 2-7. (a) CO$_2$ Sorption profiles in LDPE and PS samples obtained by the step change in pressure from 11 to 12 MPa at 200 °C. The solid fitting lines are calculated by Fick’s second law [71]; (b) a typical diagram of sorption/desorption experimental measurements [121]............................................................ 26
Figure 2-8. Schematic of common pressure-decay apparatus [122].......................... 26
Figure 2-9. The schematic of a typical Fiber Bragg grating sensor [126].................... 28
Figure 2-10. The spectrum of a Fiber Bragg grating sensor in different refractive index environment [126].............................................................................................. 29
Figure 2-11. The principle of fiber bragg described by Hill et al. [135]. The Bragg resonance for reflection occurs at the wavelength which the grating pitch is one-half of the modal wavelength of the core. The transmitted wavelength would be resultant of wavelength subtracted by the Bragg reflection.......................................................... 30
Figure 2-12. Transmission spectrum using TFBG fabricated by Miao et al. [137]...... 31
Figure 2-13. The comparison in relative shifts between the thinned FBG sensor and unperturbed FBG sensor [138]................................................................................. 32
Figure 2-14. A novel in-fiber structured intensity based tilted fiber Bragg grating sensor with a lateral offset [140]............................................................................................ 33
Figure 2-15. An image of (a) six holes and (b) two-ring triangular photosensitive microstructured optical fibers [141]................................. 34
Figure 2-16. The resonance shift versus refractive indices for (a) six holes and (b) two-ring triangular microstructured optical fibers [141]................................. 34
Figure 2-17. A schematic of a liquid core waveguide based refractometer. The slot for liquid filling can be seen across the fiber Bragg gratings [142]. ......................... 35
Figure 2-18. Reflection spectrum diagrams of the proposed liquid core waveguide based refractometer when the refractive index of the oil is (a) below and (b) above the refractive index of the optical fiber. The arrows in (b) corresponds to the fundamental mode of the liquid core waveguide and the inset focuses on these peaks. (c) Wavelength shift and change of the intensity of the main Bragg peak with respect to RI of the oil. (d) Experimental and simulated wavelength shift of the liquid core FBGs. (e) Sensitivity of the device over the low RI and high RI regime [142]................................. 36
Figure 2-19. The transmission profile of LPG in response to (a) sugar solution (b) ethylene glycol solution (c) salt solution and (d) all three plotted together [143]........ 37
Figure 2-20. The wavelength shift in response to the surrounding refractive index [144]............................. 38
Figure 2-21. Wavelength shift plotted against the changing of cladding radius by etching [144]................................. 39
Figure 2-22. The transmission spectrum of the large mode area LPG PCF in response to high refractive index [153]......................................................... 41
Figure 2-23. The transmission spectrum of (a) LPG PCF, (b) single mode LPG, (c) depicting wavelength shift with respect to surrounding refractive index and (d) depicting intensity change with respect to the surrounding refractive index. .................. 41
Figure 2-24. Transmission responses of LPG sensor in refractive range of 1<n<1.7 from two different literature [19, 31]......................................................... 42
Figure 2-25. The transmission response of LPG-MZI and tapered LPG-MZI to the variation of surrounding refractive index and (b) transmission response of taper-MZI to the variation of surrounding refractive index [154, 155]................................. 43
Figure 2-26. A phase shift based high sensitive refractive index in fiber MZI by Q. Rong and the research group [156]. (a) A schematic of the diameter mismatch MZI and (b) wavelength shift in response to varying surrounding refractive index. .................. 45
Figure 2-27. A refractive index sensor using the core offset MZI configuration by Q. Yao [157].......................................................... 45
Figure 2-28. Change in transmission spectra of PCF MZI with respect to varying refractive index [158].......................................................... 46
Figure 2-29. Intensity based transmission spectrum in MSM configuration [163]........ 47
Figure 2-30. Change in wavelength shift with respect to surrounding refractive index [164, 165].......................................................... 48
Figure 2-31. A schematic of a single mode-thin core-multimode-single mode fiber configuration [166].......................................................... 49
Figure 2-32. Transmission spectra of the STMS configuration with increasing surrounding refractive index [166].......................................................... 49
Figure 2-33. The schematic of multimode-coreless-multimode refractive index sensor [167].......................................................... 50
Figure 2-34. The transmission spectra of the MSM configuration in region I and region II with varying surrounding refractive index [167]...... 51
Figure 2-35. A schematic of TiO$_2$ coated LPG refractive index sensor developed by L. Coelho et al. [172].................................................................................................................. 53
Figure 2-36. The effect of TiO$_2$ coating on the measurement of refractive index. The plots on the right shows the effect of varying thickness [172]................................................................. 53
Figure 3-1. The schematic of a long period grating sensor [17]............................................... 56
Figure 3-2. A spectrum of the long period grating sensor ......................................................... 56
Figure 3-3. Simulated data of coupling wavelength shift against the surrounding refractive index greater than refractive index of the cladding [27]......................................................... 60
Figure 3-4. (a) The femtosecond laser system and (b) the computer controlled 4-axis stage used for aligning fiber [127]........................................................................................................ 62
Figure 3-5. A schematic of the setting used for fabricating LPG [127]........................................ 62
Figure 3-6. Atomization based coating system. ........................................................................ 64
Figure 3-7 A schematic of the refractive index experiment using LPG sensor ................. 65
Figure 3-8. A transmission spectrum for uncoated LPG in the leaky configuration. No noticeable shift in wavelength is seen................................................................. 66
Figure 3-9. A transmission spectrum for 7 passes of silver nanoparticle coating deposited LPG in the leaky configuration. No noticeable shift in wavelength is seen. ................. 67
Figure 3-10. About 0.7 nm wavelength shift occurred in refractive index ranges of 1.4823 to 1.4944, 0.3 nm from 1.4944 to 1.5094, and 0.3 nm from 1.5094 to 1.5365.................. 68
Figure 4-1. A schematic of the Mach-Zehnder interferometer. (a) a light source (b,e) couplers (c) a reference arm (d) a sensing arm (f) an optical spectrum analyzer .......... 72
Figure 4-2. A Mach-Zehnder interferometer created by a pair of long period gratings .. 73
Figure 4-3. A schematic of (a) core mismatch and (b) small single mode fiber Mach-Zehnder interferometer ............................................................................................................. 74
Figure 4-4. A sample spectrum of 40 um Mach-Zehnder interferometer sensor................. 74
Figure 4-5. A schematic of tapered single mode fiber Mach-Zehnder interferometer .... 75
Figure 4-6. A cross section view of a photonic crystal fiber [198].................................. 75
Figure 4-7. a schematic of photonic crystal fiber Mach-Zehnder (a,e) single mode fiber (b,d) collapsed region (c) photonic crystal fiber .................................................. 76
Figure 4-8. Fujikura FSM 40PM fusion splicer used for all splicing operations. .......... 81
Figure 4-9. A schematic of a single mode fiber spliced to the photonic crystal fiber using Fujikura FSM 40PM fusion splicer. ................................................................. 81
Figure 4-10. Endlessly single mode LMA PCF used for MZI configuration................. 82
Figure 4-11. A schematic of refractive index experiment using MZI sensors................. 83
Figure 4-12. Transmission spectrum of 40 um microfiber MZI in response to air and water.......................................................... 84
Figure 4-13. The transmission spectrum of 40 um microfiber MZI. Note that the transmission dip is very small................................................................. 84
Figure 4-14. The transmission spectrum of 40 um microfiber MZI. No noticeable shift has been observed................................................................. 85
Figure 4-15. A transmission spectrum for uncoated PCF MZI in varying high refractive index................................................................................................................. 86
Figure 4-16. A transmission spectrum for uncoated PCF MZI zoomed in at wavelength 1530 nm to 1550 nm. Clearly, the wavelength shifts to the right while the transmission intensity decreases increasing surrounding refractive index. ........................................ 87
Figure 4-17. About 1.6 nm wavelength increased by increasing refractive index from 1.4823 to 1.5365. ................................................................. 87
Figure 4-18. Transmission spectrum of (a) 11 mm PCF MZI (b) 20 mm PCF MZI and (c) 25 mm PCF MZI ........................................................................................................ 91
Figure 4-19. The comparison between a single mode fiber-core only fiber-single mode fiber sensor with a PCF MZI sensor. ........................................................................... 91
Figure 4-20. The transmission spectrum in leaky mode configuration of (a) uncoated 11 mm PCF MZI (b) coated 11 mm PCF MZI (c) uncoated 20 mm PCF MZI (d) coated 20 mm PCF MZI (e) uncoated 25 mm PCF MZI (f) coated 25 mm PCF MZI ............... 95
Figure 4-21. Change of transmission intensity in (a) uncoated and (b) coated 11 mm PCF MZI .................................................................................................................. 96

Figure 5-1. A schematic of D-shaped optical fiber refractive index sensor. Refractive index solution would make direct contact with the core [204] .................................................. 99
Figure 5-2. Light guiding of the femtosecond laser machined microhole refractive index sensor [205] ................................................................. 100
Figure 5-3. A schematic of the gap sensor ................................................................. 102
Figure 5-4. Normalized transmission value with respect to refractive index in the primary experiment setup, but misaligned. ................................................................. 104
Figure 5-5. Normalized transmission spectrum of the gap sensor with increasing refractive index. Note that the intensity increases up to about 1.45 and starts to decrease then. The general trend of the sensor is similar to the primary experiment results. ...... 105
Figure 5-6. The schematic of the gap distance parameterization .................................. 107
Figure 5-7. Change in the transmission intensity with increasing gap distance .......... 107
Figure 5-8. The change in transmission intensity with response to temperature. ........ 108
Figure 5-9. Difference in transparency between solid and molten PCL ....................... 109
Figure 5-10. The transmission intensity of PCL film with varying temperature .......... 110
Figure 5-11. The transmission spectrum of polymer films compared with the RI solutions. ......................................................................................................................... 111
Figure 5-12. 1 mm by 1 mm by 189 μm thick PET film machined using femtosecond laser. .......................................................................................................................... 113
Figure 5-13. A customized pressure chamber used to pressurize PET film. (A) a CO₂ tank (B) a high pressure regulator (C) a valve (D) a sample chamber (E) an exit valve. ...... 113
Figure 5-14. The change in transmission intensity with time as the CO₂ desorbs out of PET film .................................................................................................................. 114
Figure 5-15. The CO₂ sorption film in PET using weight change method in literature. 114
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Chapter 1. Introduction

Polymers are widely used in various applications after becoming high-value products by many processing techniques such as composite manufacturing, blending, and foaming. Depending on how polymers are processed, the properties of finished products can be significantly different. Polymer processes may go through several chemical reactions and/or mechanical processes to shape raw polymer materials from pellets, granules, flakes, or powders, into the desired finished products. The finished product may have different morphologies and properties from the raw materials [1]. Polymer processes involving supercritical fluids (SCFs) and gases are one of the most prevalently used processes and offers wide variety of applications researched in the field. Polymer processes using gas/SCF can be categorized into two major groups. The first group is a process where a polymer is dissolved into a gas/SCF solvent. The second group includes the processes where a gas/SCF is dissolved into a polymer. In both categories, correct understandings about the interactions between a gas/SCF and a polymer are critical for the optimum fabrication process. Many properties of polymers can be affected by the dissolved SCFs and gases at high temperatures and pressures. Among these effects, solubility and diffusivity are crucial parameters in polymer processing applications. Hence, different devices to measure solubility and diffusivity have been researched, but most of the devices used today are very complex, expensive, and requires long experiment time for operation. In this final thesis, the feasibility of the usage of fiber optic sensors as measurement devices for solubility and diffusivity of gas/SCF in polymers have been investigated. The fiber optic sensors have already been investigated
in gas sensing applications, such as in CO$_2$ detection [2, 3]. Therefore, by configuring the experimental setup for a molten polymer, gas diffusing into polymers could be measuring using the fiber optic sensors. As gases dissolve into polymers, it would alter the refractive index (RI) of the polymer resin, which could be detected using fiber optic refractive index sensors. Many of the polymers used in polymer processing have high refractive index, around from 1.4 to 1.68. However, fiber optic sensors offer very low sensitivity when the surrounding refractive index becomes greater than that of the cladding, which is approximately 1.44. This final thesis investigates the feasibility of fiber optic sensors in sensing high refractive index for the future application of solubility and diffusivity measurement.

The recent advances in fiber optic sensors has extended measurement technologies and telecommunications a step further by providing solutions to current limitations with its unique characteristics. Different types of fiber optic sensors, such as in fiber interferometers or in fiber gratings, are prevalently used for temperature, refractive index, pressure, and strain sensing because fiber optic sensors offer the unique properties such as remote sensing, low propagation loss, high sensitivity, high accuracy, low cost, and immunity to electromagnetic interference. The quality of fiber optic sensors depends on the reliability, robustness, and ease of a fabrication process. The feasibility of fiber optic sensors for detecting the solubility and diffusivity of gas/SCF in a polymer is discussed in this final thesis. In this chapter, the fundamental knowledge in the solubility and diffusivity of gas/SCF in a polymer, fiber optics sensors, limitations encountered by the sensors, and the research scope of this final thesis would be presented.
1.1. Solubility and diffusivity of gas/SCF in a polymer

According to the International Union of Pure and Applied Chemistry (IUPAC) definition [4], solubility is the proportional ratio of a solute in a solvent. Diffusion can be described as the process where a material is transported by the thermal motion of the molecules in a fluid or a matrix. Solubility and diffusivity act as crucial parameters in many polymer processing applications. For example, in the fabrication of nano-/micro cellular foamed plastics, solubility information provides the maximum allowable amount of a gas/a SCF dissolved into a polymer resin at a specific temperature and pressure. Solubility significantly affects the cell nucleation and growth processes and thus, the final foam product properties [5]. Moreover, solubility and diffusivity characteristics are crucial in polymer impregnation processes as they would control additive contents in polymers. In polymer impregnation, both solubility of a solute in a SCF and solubility of a SCF in a polymer are significant parameters [6]. It is also known that SCF solubility in polymers has also a considerable effect in the polymer blend structures in extrusion because a SCF reduces the viscosities and interfacial tensions of the polymers and changes the final blend morphologies [7]. Since solubility and diffusivity have significant effects in the polymer processes as mentioned earlier, further knowledge in solubility and diffusivity measurement methods is essential to allow more flexibility in these polymer processes and the relevant applications.

SCFs are frequently involved in polymer processes because their unique properties provide numerous advantages as an alternative to other organic solvents. A SCF is a substance above its critical temperature and pressure as shown in Figure 1-1. SCFs behave similarly to gas in terms of viscosity and diffusivity while they behave like liquid
for density and solvating properties. The properties such as density, diffusivity, or viscosity can be tuned simply by adjusting the operating pressure and temperature. Moreover, SCFs are also widely used due to their economic advantages. SCFs can be recovered and reused without any purification processes. In addition, the usage of SCFs can benefit many industrial processes from low energy consumption and minimum toxicity when compared to the conventional organic solvents. SCFs are non-carcinogenic, non-flammable, and thermodynamically stable, which are the health and safety advantages of SCFs [8]. As mentioned above, some polymer processes, such as the micro-particle generation use SCFs as solvents to dissolve polymers [7, 9-14]. In this final thesis, the potential usage of fiber optic sensor for measuring solubility and diffusivity of gas/SCF in polymers have been investigated.

![Figure 1-1](image)

**Figure 1-1.** Schematic of pressure-temperature phase diagram where the triple and critical points are shown. The phase regions are labeled in the diagram.

1.2. **Fiber optics sensors**

A fiber optic sensor is a sensor that uses an optical fiber as a basis of the sensing element. Fiber optic sensors are configured in a way so the light from the source
propagates through the optical fiber sensor to the detector, usually in the reflection or transmission mode. Depending on whether the optical fiber sensor actually modulates the light or not, the fiber optic sensors can be categorized as an extrinsic optical fiber sensor or an intrinsic optical fiber sensor. In the extrinsic optical fiber sensor, the transducer, which is not an optical fiber, modulates light, and optical fiber is used to propagate modulated light to a converting device. Extrinsic sensors are used in applications requiring low resolution and low cost because extrinsic sensors generally do not provide high resolution measurements [15]. An example of extrinsic optical fiber sensor is an extrinsic Fabry-Perot type sensor used by S. Chen et al. [16]. In case of the intrinsic optical fiber sensor, the light modulation is caused by the optical fiber, and grants control over the light propagating in the fiber. Although the fabrication is more complex compared to extrinsic sensors, intrinsic sensors offer better performances, and therefore, intrinsic sensors are often used in applications as spectral filters [15, 17]. An intrinsic Fabry-Perot sensor, long period gratings sensor, fiber bragg gratings sensor, and Mazh–Zehnder Interferometer sensors all fall under the category of an intrinsic optical fiber sensor [18-20]. Furthermore, the fiber optic sensor is also categorized into an intensity based sensor or shift based sensor depending on its operating principle. The intensity based sensing is the simplest approach where the intensity modulation of light from optical spectrum analyzer is used for sensing [21]. The disadvantages in the intensity based sensors are that random losses exist in the sensing environment. For example, there could be intensity losses from splice joints, fiber connections, bending. Changing a fiber position could lead to a change in intensity. A shift based sensor, on the other hand, uses the shifts of a phase or wavelength in response to the environment for sensing. The shift
based sensor provides high sensitivity to temperature, strain, and refractive index, and therefore is used widely [22-25].

1.3. **Significance of the problem**

Most of the solubility and diffusivity measurement devices currently used in practices, such as the magnetic suspension balance or the pressure decay system, are expensive, complex, and requires long experimental time. In order to improve upon the mentioned aspects in measuring the solubility, the fiber optic refractive index sensor could potentially be implemented. As gas sorb into a polymer, its refractive index would change, which is already hypothesized using Gladstone-Dale relation and have been observed by R. St-Gelais et al. [26]. By observing this change in refractive index, the solubility and diffusivity of gas in a polymer can be obtained. Fiber optic sensors are ideal for this measurement because they offer accurate measurement of refractive index, even in the harsh environment such as in high temperature, pressure, and electromagnetic fields. However, a significant problem would be encountered when fiber optic sensors are attempted to be used in measuring solubility and diffusivity of gases in a polymer. Fiber optic refractive index sensors offer very low sensitivity when the surrounding refractive index (SRI) becomes greater than that of silica, which is approximately 1.45. For example, as illustrated in Figure 1-2, R. Hou et al. have simulated the sensitivity of long period gratings sensor with varying refractive index and showed that once SRI becomes greater than 1.45, the sensitivity with refractive index becomes almost negligible. This is a significant problem because majority of the polymers used for processing have refractive index greater than 1.45. Some of the main polymers used for polymer
processing are tabulated in Table 1-1. Polymers widely known, such as PMMA, PVA, Nylon 6, PS, PVC, and PE, all have refractive index above 1.5. Therefore, a solution to sense refractive index in regions greater than the cladding is necessary to allow solubility measurement using the fiber optic sensors.

Table 1-1 Refractive indices of polymers prevalently used in polymer processing.

<table>
<thead>
<tr>
<th>Polymers</th>
<th>Refractive Index (n)</th>
<th>Polymers</th>
<th>Refractive Index (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polytetrafluoroethylene</td>
<td>1.35-1.38</td>
<td>Polybenzyl methacrylate</td>
<td>1.57</td>
</tr>
<tr>
<td>Poly-4-methylpentene-1</td>
<td>1.47</td>
<td>Styrene-acrylonitrile copolymer</td>
<td>1.57</td>
</tr>
<tr>
<td>Polymethyl methacrylate</td>
<td>1.5</td>
<td>Polyphenylene methacrylate</td>
<td>1.57</td>
</tr>
<tr>
<td>Polyvinyl alcohol</td>
<td>1.49-1.53</td>
<td>Polydiallyl phthalate</td>
<td>1.57</td>
</tr>
<tr>
<td>Diethylene glycol bis allyl carbonate</td>
<td>1.5</td>
<td>Polyethylene terephthalate</td>
<td>1.58</td>
</tr>
<tr>
<td>Polycyclohexyl methacrylate</td>
<td>1.51</td>
<td>Polystyrene</td>
<td>1.59</td>
</tr>
<tr>
<td>Polyethylene</td>
<td>1.51</td>
<td>Polyvinyl chloride</td>
<td>1.63</td>
</tr>
<tr>
<td>Polyacrylonitrile</td>
<td>1.52</td>
<td>Polyvinyl naphthalene</td>
<td>1.68</td>
</tr>
<tr>
<td>Nylon 6</td>
<td>1.53</td>
<td>Polyvinyl carbazole</td>
<td>1.68</td>
</tr>
</tbody>
</table>

Figure 1-2. Simulated data of coupling wavelength shift against surrounding refractive index on a wide range [27]
1.4. Fiber optic refractive index sensors in refractive index greater than the silica

In order to utilize fiber optic sensors for measuring the solubility and diffusivity of gas/SCF in polymers, the fiber optic sensors must be capable of measuring surrounding refractive index that is higher than the cladding. Most of the fiber optic refractive index sensors set a limitation in its range so the surrounding refractive index is smaller than the refractive index of the cladding. This is because once the surrounding refractive index (SRI) becomes greater than the refractive index of the cladding ($n_{\text{cladding}}$), the total internal reflection is lost and becomes a leaky configuration. As the SRI approaches to the refractive index of the cladding, the light propagation at outer cladding modes become leaky as total internal reflection is lost at the boundary of the outer cladding and the environment. In the early times, the case of a SRI being greater than that of silica (i.e. leaky configuration) did not capture interests because no guided propagation happened into the cladding in the leaky configuration [28]. However, Lee et al.’s work on couplings in a leaky configuration has found a potential in refractive index sensors operating on a wide range of refractive index values [19, 29]. In the leaky configuration, it is conventionally known that light would leak from losing the total internal reflection [30]. For the leaky wave guiding situation, the dispersion relation of effective refractive index in cladding mode no longer has purely real solutions [31]. However, it is important to know that even with the light leaking out in transversely, the light do propagate in a longitudinal direction with attenuated amplitude [32]. This small portion of propagation is from Fresnel reflection and this could enable reasonably low-loss propagation for some applications [31]. Only limited number of studies have reported in Mach-Zehnder Interferometer (MZI) based refractive index sensors in the leaky configuration as well
because the response of MZI to varying surrounding refractive index ceases to be linear in this domain [33]. In leaky configurations, certain optical sensors do react with either a shift based or intensity based response. However, the sensitivity of these responses are very low and insufficient to be used as a refractive index sensor in leaky configuration. Moreover, some researchers endeavored and succeeded in measuring high refractive index but their sensitivity was still low and required usage of high cost fibers and fabrication methods. They also neglected temperature dependence on the refractive index which limits the usage in the practical applications [19, 27, 32-35]. Some of these literature reviews will be discussed in Chapter 2. To apply fiber optic refractive index sensors for the measurement of solubility and diffusivity of gas/SCF in a polymer, the problem of low sensitivity in the leaky configuration must be overcome.

1.5. Research scope and outline

The research scope of this final thesis is to investigate the feasibility of in fiber gratings, in fiber interferometers, and transmission intensity based fiber optic sensors for acquiring solubility and diffusivity of gas/SCF in a polymer. To use fiber optic sensors for solubility and diffusivity measurement, the sensors must be capable of sensing refractive index higher than silica, which has been little of interest. This final thesis suggests a fiber optic based sensor that is capable of sensing in high refractive indices, and potentially be used to measure solubility and diffusivity measurement in polymers. Chapter 1 introduces the solubility and diffusivity of gas/SCF in polymers, fiber optic sensors/interferometers and the problems encountered in leaky configurations in fiber optic refractive index sensors. Chapter 2 reviews and discusses current technologies
developed in literature in solubility measurement, diffusivity measurement, fiber optics as refractive index sensors. Fiber brag gratings sensors, long period gratings sensors, and all the different configurations of Mach-Zehnder interferometers as refractive index sensors would be presented. Moreover, any refractive index studies conducted in leaky mode would also be within the scope of the literature review. Chapter 3 of this thesis presents long period grating sensor for measuring high surrounding refractive index solution. The fabrication of long period gratings sensor using the femtosecond laser, the process of coating LPG, and the behavior of both coated and uncoated LPG in high refractive index solutions are discussed. In Chapter 4, the behavior of smaller core based Mach-Zehnder interferometer, PCF Mach-Zehnder interferometer, and the coated PCF Mach-Zehnder interferometer to the high refractive index solutions are presented. The fabrication process of the discussed MZIs would be introduced as well. In Chapter 5, the novel intensity based gap sensor for measuring high refractive index solutions is described. The sensing principle, experiment setup, and parametric studies are present along with the high refractive index sensing data. Preliminary experiments using polymer films have been conducted to test the feasibility of the gap sensor. Finally in Chapter 6 provide the overall conclusion of the thesis and suggest a potential future works to improve fiber optic sensors in high refractive index environment.
Chapter 2. Theoretical background and literature review

In this chapter, first, the current status of measurement methods for measuring solubility and diffusivity of gas/SCF in polymers are reviewed. Then, the relevant research for the measurement of refractive index using fiber optics sensors is reviewed. The usage of in fiber grating sensors, in line interferometers, and in fiber intensity based sensors used as refractive index sensors are reviewed. Moreover, since measurement of high refractive index leads to loss of perfect internal reflection of the propagating guided mode at the cladding to surrounding refractive index interface, the refractive measurement in the leaky configuration are investigated.

2.1. Methods for measuring solubility of gas/SCF in polymers

Solubility data are crucial in many of polymer processes, such as polymer impregnation, blending, and microcellular foaming. Hence, different techniques for measuring gas/SCF solubility in a polymer have been investigated by numerous researchers. In this section, various theoretical methods used to obtain solubility data are discussed in detail. In addition, the limitations of each method are described as well. Commonly, theories, such as equations of states and empirical equations, are used to improve the accuracy of the measured solubility and diffusivity data. For further information in regards to theories, refer to Appendix A.

2.1.1. Gravimetric methods
The gravimetric method measures solubility of a gas/SCF dissolved in a polymer by acquiring the changes of the sample weight by a gas/SCF. When a polymer sample is exposed to a gas/SCF, the sample swells and the volume change occurs which affects the calculation of solubility data. Therefore, in gravimetric methods, it is critical to compensate for a volume swelling of the sample since the volume swelling correction can affect up to 40% of the measured weight difference depending on gas/SCF and polymer types, and pressure ranges [36]. There are experimental techniques such as a photometric method or a thermomechanical analyzer (TMA) method to measure the sample swelling and buoyancy behaviors [37-40]. Also, the EOS theories such as Henry’s Law, SL-EOS, Flory-Huggins theory, SS EOS, or SAFT EOS are used to account for swelling. Y. Zhang and co-investigators studied the swelling and sorption behaviors of various polymers in the presence of CO₂ at 35 °C and 10.3 MPa [41].

A quartz spring balance is one of the early gas/SCF solubility measuring techniques based on the gravimetric method. This apparatus determines solubility by monitoring the extension of a spring from an increased sample’s weight. Using the Hooke’s Law, the mass of the sample is determined as a function of a spring elongation. The spring used in this technique is fabricated from quartz or similar materials such as tungsten. J.W. McBain is one of the early researchers who used a quartz spring apparatus to study the gas sorption behaviors in rubbers and naturally occurring polymers [42]. Thus, this apparatus is commonly referred as a McBain Balance. The schematic of a quartz spring balance used by McBain can be seen in Figure 2-1. A McBain balance is suitable for the sorption study of organic vapors or gases that are highly soluble in polymers. Only a small portion in a McBain balance is required to be heated and a pressure is held constant.
for solubility measurements. Precautions are necessary when operating a McBain balance as this device is sensitive to a vibration. The preliminary volume calculations required for solubility calculations in other types of gravimetric measurement techniques are not required in a McBain balance. Instead, the sensitivity of a spring and the extension per unit weight calculations must be done for solubility calculations [36]. The sensitivity of a McBain balance is low and therefore it is hard to measure solubility of light gases in a polymer [36]. However, this limitation can be overcome by using more modern and sophisticated types of quartz spring for a higher sensitivity. These modifications on a McBain balance allow the users to operate up to 1.01 MPa [43]. Furthermore, precise temperature and pressure controls can be implemented to improve the experimental accuracy [44, 45]. A McBain balance is compatible with different gases and polymers in the moderate range of temperatures (35 – 178 °C) with modifications. G.E. Serad and co-researchers used a McBain balance to study the sorption and diffusion behaviors of n-Butane in PET [46]. S. Lequin and co-investigators used a McBain balance to study water vapor sorption on cork [47], and Y. Iwai and Y. Arai studied solubility of m-xylene and nonane vapors in PS, and ethylbenzene and nonane vapors in polybutadiene (PBD) in the temperature ranges from 130 to 175 °C and 80 to 130 °C, respectively [48]. A McBain balance is also capable of measuring gas/SCF solubility of a thin film sample. This can be done by suspending the thin film sample in the cell. However, this technique is impossible if the sample is in a semi-liquid state or in a molten state (i.e., high temperature experiments). This limitation can be overcome by utilizing suspended pans or caps but this can cause other experimental challenges such as sample adhesion to the bottom of the pan. As mentioned above, gravimetric methods must consider for the
buoyancy effect and, in case of a McBain balance, the calibration of the unstrained spring needs to be calculated as well before the solubility calculation [44, 49, 50]. The weight, temperature, and pressure sensitivities of a modified McBain balance are ± 0.05 mg, ± 0.5 °C up to 400 °C [51], and ±1 Pa [47], respectively. The sample of 5 to 20 mg can be loaded on the spring [52].

**Figure 2-1.** Schematic of a McBain Balance in superposed thermostats: (A) heating coil, (B) quartz spring, (C) polymer specimen [42]

Usage of electronic microbalances are also available for gas/SCF solubility measurements in polymers. Electronic microbalances can take a smaller sample sized specimen because of a higher sensitivity. In addition, these microbalances have a shorter measurement time compared to McBain type balances [40]. Three electronic
microbalances are frequently mentioned in literatures: ones manufactured by Cahn Instruments, Sartorius Stedim Biotech GmbH, and Mettler Toledo. In a Cahn type microbalance, the sample is positioned in a high pressure chamber and is balanced with a weight suspended on the other end of a beam. Electronic signals with a variable gain amplifier and a digital to analog converter are used to observe any imbalances of the device. A Cahn microbalance was used by Y. Kamiya and the research group to study sorption and dilation of a polymer with various gases [53-56]. Moreover, B. Wong and investigators used a Cahn microbalance to determine solubility and diffusivity of CO$_2$ and HFC134a in PS, filled poly(vinylchloride) (FPVC), and unplasticized poly(vinylchloride) (UPVC) [57]. The schematic of the sorption apparatus using a Cahn microbalance is shown in Figure 2-2. In a Sartorius type microbalance, an electromagnet coil is connected at the center of a beam within a permanent magnet field. The current in the coil changes relatively to the change in the sample’s weight. This fluctuation of current is monitored with a microprocessor to determine the weight of the sample. Y. Kamiya and the group studied CO$_2$ sorption and dilation of Poly(methyl methacrylate) (PMMA) using a Sartorius microbalance [58]. Similarly, L. Phan Thuy and J. Springer studied CO$_2$ sorption in poly(butylene terephthalate) (PBT) [59]. In addition, N. Von Solms and co-authors made a direct measurement of solubility and diffusion of methane and CO$_2$ in high density polyethylene (HDPE) [60]. The schematic of a sorption apparatus using a Sartorius electronic microbalance is shown in Figure 2-3. Several sorption studies in literature used a Mettler AE163 electronic microbalance, which is sensitive up to ± 0.0001g, to study sorption of CO$_2$ in poly(chlorotrifluoroethylene)
(PCTFE), PDMS, PMMA, and PS [61-64]. More recently, C. Gutierrez et al. studied the sorption of CO$_2$ in PS using the same technique [65].

**Figure 2-2.** High pressure sorption apparatus with a Cahn model 2000 electronic microbalance by Kamiya [56]: (A) pressure chamber containing an electronic microbalance, (B) thermostatically controlled air bath, (C) temperature controller, (D) thermometer, (E) pressure gauge, (F) safety valve, (G) flow meter, (H) columns of active charcoal and silica gel, (I) pressure regulator, and (J) a gas cylinder [56]

**Figure 2-3.** Sorption apparatus with a Sartorius S3D-P electro microbalance by Kamiya [58]: (A) Sartorius microbalance, (B) constant temperature water bath, (C) auxiliary furnaces, (D) constant temperature air-bath, (E) pressure chambers, (F) thermocouple, (G) water inlet, (H) gas inlet, (I) polymer sample plus buoyancy corrective (gold wire), (J) counter balance (aluminum rod) [58]
A magnetic suspension balance (MSB) is another famous instrument allowing gravimetric measurements of gas/SCF solubility in polymers. A MSB is differentiated from other gravimetric techniques described earlier due to contactless weight measurements. A microbalance in a MSB is isolated from a high pressure and temperature sample chamber and is positioned in an ambient environment. A MSB is capable of measuring gas/SCF solubility in extreme pressures and temperatures unlike other gravimetric techniques, such as a McBain balance, and shares similar advantages such as a high sensitivity with other electronic microbalances mentioned earlier. A polymer sample is positioned in a sample holder and it is isolated from the balance as shown in Figure 2-4. The sample holder is hung on to a suspension magnet (i.e., permanent magnet), which is kept free in the sorption chamber to allow the transmission of a magnetic attraction force to the external electronic balance. Therefore, the weight measured in the sorption chamber can be precisely transmitted to the isolated microbalance through this suspension magnet. A MSB is used by numerous researchers to measure gas/SCF solubility in polymers. For example, Li, G., Gunkel, F., Wang, J., Park, C.B., and Altstädt, V. measured solubility of N$_2$ and CO$_2$ in PP and Ethylene Octene copolymers from 180°C to 220°C and 160°C to 200°C, respectively, up to pressures of 27.6 MPa [66]. Y. Sato and co-workers found solubility of CO$_2$ in Polyphenylene Oxide (PPO) and PPO/PS blends using a MSB [67]. A MSB can detect mass changes of 3×10$^{-5}$g. A sample mass for measurements is from 0.01 to 30 g with a pressure control of ±10 Pa between 0.1 ×10$^5$ and 1.33 ×10$^5$ Pa, and a temperature control of ±0.05 °C from 20 and 150 °C [52]. Many literatures mention a MSB manufactured by Rubotherm GmbH (Rubotherm) to study solubility and diffusivity of a gas/SCF in
various polymers. B.I. Chaudhary and A.I. Johns studied the sorption of N\textsubscript{2}, Isobutane, and CO\textsubscript{2} in PE in the temperature range of 110 to 200 °C up to 20 MPa [68]. Von Schnitzler et al. measured the sorption of CO\textsubscript{2} in PBT with a MSB in the temperature range of 40 to 120 °C and the pressures up to 30 MPa. Y. Sato and the group studied CO\textsubscript{2} solubility in various polymers and polymer blends using a MSB [67, 69, 70]. Furthermore, S. Areerat and the research group also studied solubility of CO\textsubscript{2} in molten polymers such as HDPE, Low Density Polyethylene (LDPE), PP, Ethylene-Ethylacrylate copolymer (EEA) and PS [71, 72]. In more recent work, Z. Lei and co-authors studied CO\textsubscript{2} sorption in PP. Similarly, E. Aionicesei et al. studied CO\textsubscript{2} solubility in PEG, poly(l-lactide) (PLLA) and poly(d,l-lactide-co-glycolide) (PLGA). In 2015, N. Trupej and co-investigators measured the solubility and diffusion coefficient of a Ar gas in PEG using a MSB [73-76].

![Figure 2-4. The schematic of a solubility measurement apparatus with a MSB by Sato [69].](image-url)
2.1.2. Piezoelectric methods

Piezoelectric crystals can be used to investigate gas sorption behaviors in polymers using the principle of piezoelectric crystal frequencies. The vibration frequency of a piezoelectric crystal changes with respect to its mass, and this frequency change can be measured using a control system [36]. The oscillation frequency of a polymer sample coated atop a piezoelectric crystal is used for the solubility measurement. Prior to the experiment, it is necessary to know the frequencies of uncoated and coated crystals at an ambient environment. Then, the weight of the sample is determined as a function of gas pressures, and through the mass balance, the weight of a gas dissolved in the sample can be calculated. D.C. Bonner and Y.L. Cheng used a piezoelectric crystal sorption device to study N\textsubscript{2} sorption in LDPE [77]. Moreover, this method is well elaborated in the studies of B.J. Brisco and the research group where they measured gas solubility and gas induced dilation in poly(urethane) elastomers using a vibrating beam technique, which involves the measurement of the first harmonic resonant frequency of the beam that can vary with the mass of gas absorbed in a polymer [78-80]. More recently, the work by K. Miura and co-authors with a quartz crystal microbalance (QCM) attracted people’s attention, where quartz crystals are used as mass ‘sensors’ to obtain CO\textsubscript{2} solubility and adsorption data at high pressures [81]. The schematic of a solubility measurement setup using a QCM are shown in Figure 2-5. K. Miura et al. used a platinum resistance thermometer to measure a temperature inside the cell and a strain gauge to measure the pressure of the cell. The solubility was obtained by calculating the frequency difference between the initial frequency and the measured frequency at a certain pressure and temperature. G. Wibawa and co-investigators used a QCM to measure solubility of seven nonpolar organic
solvents such as octane, cyclopentane, cyclohexane, benzene, toluene, ethylbenzene and p-xylene in cis-1,4-polyisoprene, poly (isobutylene) PIB, poly(n-butyl methacrylate), and poly(vinyl acetate) (PVA) [82]. The advantages of a piezoelectric method are; first, this method does not require a volume change correction as other measurement methods due to its vibration frequency based measurement; second, a QCM has a short experiment time; third, a very small change in weights can be measured; and lastly, it is relatively easy to add an automated measurement system [81, 82]. However, the hysteresis error from the adhesion of polymer films to the crystals should be taken into account [36, 83]. Moreover, since a QCM is a fast measurement device, it cannot detect the slow changes during sorption caused by the polymer relaxation which is independent of the film thickness [52, 84]. Preparing a thin polymer film on the crystal is a fundamental challenge, limiting the types of polymers used with a QCM. CO$_2$ solubility in poly 4-methyld-1-pentene (TPX), PMMA, polyimide, PC, PS, chlorinated polyethylene, poly(xylene), poly(2-chloroxylylene), poly(2,3-dichloroxylylene), and teflon were investigated by James Aubert using a QCM up to 40 °C and 9.65 MPa [85]. D. Boudouris and co-researchers studied the sorption behaviors of HCFC-22 and HFC-152a in PMMA, PS, and PC, and M. Pantoula and C. Panayiotou studied CO$_2$ sorption in PMMA and PS [86]. In their work, M. Pantoula and C. Panayiotou compared the obtained solubility data with a QCM with other literature data to elucidate the accuracy of their results [87]. N.S. Oliveira et al. found solubility of N$_2$, O$_2$, CO$_2$ and water in PLA with an overall average absolute deviation of 6% when compared with the theoretical values obtained with the Flory-Huggins theory [88]. More recently, F. Herran and the research group studied the sorption of H$_2$O vapor in PC using a QCM [89]. A.L. Smith and H.M. Shirazi have
obtained the temperature control of 0.0001 °C up to 45 °C and 0.005 °C up to 110 °C in their QCM experimental setup [90].

Figure 2-5. Schematic of a sorption apparatus with a QCM used by Miura et al. [81].

2.1.3. Manometric methods

A manometric method determines the solubility of gases/SCFs in polymers by measuring the pressures or volumes of the gases sorbed in or desorbed from the sample in equilibrium [36]. Early studies were performed in sub-atmospheric pressures (0.101 MPa) [91, 92]. C.S. Venable and T. Fuwa acquired solubility data of different gases in rubber using a direct experimental procedure, which is to calculate solubility data from volume differences between before and after a gas exposure at 100°C in a sub-atmospheric pressure [93]. By monitoring the pressure or volume changes of a gas in a chamber, the solubility of gases in polymers are determined. The chamber volume calibration must be done prior to an experiment in this configuration. In these early direct experimental techniques, one of the challenges is the process of removing all ambient gas phases as soon as possible once the equilibrium is reached, while preventing the
dissolved gas from being pumped out of the sample. The usage of a high power pump for evacuating the chamber removes a gas phase effectively [94, 95].

A phase separation technique is one type of manometric methods used to obtain solubility data. In a phase separation method, the molten polymer sample is exposed to a gas under a certain pressure in an autoclave. Once equilibrium is attained, the solubility can be determined by calculating the amount of gas present in the polymer-rich phase of a sample. This method is mainly applicable to low viscosity polymers as mixings between a low viscosity polymer and a gas/SCF are easier [96]. M. Daneshvar, S. Kim, and E. Gulari used the phase separation method to obtain sorption of CO$_2$ in PEG up to 100 °C and 35 MPa, while Garg et al. obtained CO$_2$ and 1,1-difluoroethane sorption in PDMS and PS in the range of 50 to 160 °C and up to 26 MPa [96-98]. The solubility of propane, N$_2$, and CO$_2$ in PEG was also investigated by Wiesmet and the research group using a phase separation method [99, 100].

A pressure decay method is a prevalently used manometric based solubility measurement technique. This technique was developed by Newitt and Weale in 1948 to measure solubility of H$_2$ and N$_2$ in PS [101]. In a pressure decay method, a polymer sample is placed in a closed pressure vessel with a known volume and a temperature. As a preheated gas is introduced to the chamber, pressures are measured as a function of time until gas sorption to the sample completes. The solubility is determined by calculating the pressure difference between an initial gas amount and a gas amount at equilibrium. It is important to consider for the polymer sample swelling during gas sorption because this changes the total sample volume. Theoretical EOSs can be used to compensate for the swelling volume of a sample due to gas sorption. The solubility
measurement at a high temperature is particularly difficult with a pressure decay method due to the lack of suitable pressure sensors [40, 70]. Moreover, at a high pressure experiment, small gas leakage from the chamber can lead in underestimating the solubility data. Therefore, solubility measurements at high pressures require accurate EOSs, and should take account of Pressure-Volume-Temperature (PVT) behaviors of the polymers at high pressure and temperature conditions. Furthermore, polymer degradation problems could also occur at high temperature experiments [40]. For a pressure decay method, a large amount of samples is required. Hence, a long measurement time is required to obtain solubility data because reaching equilibrium takes a long period of time. The measurement errors can arise when accounting for an initial gas pressure introduced to the cell because a stabilization period is required for a pressure measuring device. During this stabilization period, a substantial amount of gas sorption can occur and this can cause solubility measurement errors. The extrapolation of a pressure decay curve to a zero time was conducted in attempt to correct for this error [102-105]. Moreover, a reference reservoir was used to correct for this type of error [106]. An attempt to use dual sorption cells was made by W.J. Koros, A.H. Chan, and D.R. Paul and they were able to measure solubility of multiple gases in PC without a stabilization related error [107]. Multi-sorption cells were also used to eliminate the errors as in the experiments by Y. Sato’s research group and V.I. Bondar’s research group [108-112]. Using these multi-sorption cells allowed the researchers to perform solubility measurements at high pressures. The schematic of a dual cell pressure decay sorption apparatus is depicted in Figure 2-6. Not only gas sorption behaviors, but also the diffusivity and permeability of different gases in PEO at 35 °C and pressures up to 4 MPa
were obtained using a dual sorption pressure decay method by H. Lin and B.D. Freeman [113]. Moreover, J.E. Groiler and the research group used a vibrating wire sensor with a pressure decay method to obtain solubility of CO$_2$ in medium density polyethylene (MDPE) and poly (vinylidene fluoride) (PVDF) [114, 115]. Recently, D. Li and investigators studied CO$_2$ sorption in solid-state isotactic polypropylene (iPP) in the temperature and pressure ranges of 100 to 150 °C and up to 15 MPa, respectively. S. Kim and co-authors studied the gas sorption of H$_2$, N$_2$, O$_2$, CH$_4$, and CO$_2$ in different thermally rearranged polymers, and C.G.F. Rezende et al. studied the sorption of propylene and propane in polyurethane containing silver nanoparticles [116-118].

**Figure 2-6.** Schematic of a sorption apparatus with dual sorption pressure decay used by Stern [105]: (A) Auxiliary glass system for volume calibrations, (A1) reference volume, (A2) thermocouple gauge, (A3) mercurial manometer, (B, C, D, E) calibrated volumes of apparatus, (D) gas reservoir, (E) gas absorption cell with polymer sample, (F) differential pressure null indicator, (G) null indicator control, (H) dead-weight gauge, (1-7) needle valves [105]
2.2. Methods for measuring diffusivity of gas/SCF in polymers

In this section, two main experimental methods for measuring gas/SCF diffusivity in a polymer under unsteady states are introduced. The advantages and disadvantages of each method are also discussed. To improve the accuracy of the measurement, theories are often used to correct the diffusivity data. For further information in regards to theories, refer to Appendix B.

2.2.1. Gravimetric method

A Magnetic Suspension Balance (MSB) [119, 120] is one of the most popular and precise devices based on the gravimetric method for diffusivity measurements. This technique gathers sample mass change data with respect to measurement time and then fits the data with a diffusivity equation to get a gas diffusivity coefficient.

An electronically controlled magnetic suspension coupling is used to transmit the measured force from the sample enclosed in a pressure vessel to a microbalance located at an ambient temperature and pressure as discussed in an earlier section 2.1.1. When a gas dissolves in a polymeric sample, the weight of the sample grows due to the dissolved gas weight. A mutual diffusion coefficient is then determined by measuring the weight change of the sample with respect to time as shown in Figure 2-7.
Figure 2-7. (a) CO$_2$ Sorption profiles in LDPE and PS samples obtained by the step change in pressure from 11 to 12 MPa at 200 °C. The solid fitting lines are calculated by Fick’s second law [71]; (b) a typical diagram of sorption/desorption experimental measurements [121]

Many diffusion experiments generate permeate flux data with respect to time to fit modified Fick’s second law functions to estimate diffusion coefficients. For example, Figure 2-7a) shows the Fick’s second law fit on the experimental data.

2.2.2. Pressure decay method

![Pressure decay apparatus](image)

Figure 2-8. Schematic of common pressure-decay apparatus [122]

Among all experimental methods for measuring gas diffusivity in a polymer, a pressure-decay method, as shown in Figure 2-8, is attractive because of its convenience, simplicity, and accuracy [123]. In 1963, J. L. Lundberg and his team members used this method to measure diffusivity and solubility of methane in PS at 33 MPa in the temperature range between 100 and 188 °C [103]. In 1969, his group repeated similar experiments using PIB [124]. In a pressure decay method, a molten polymer is molded to form a sheet where the sides of the samples are sealed to satisfy a single-sided diffusion
process. The sample is placed in the center of a high-pressure cell and the both ends of the pressure cell are sealed to ensure a gas can only diffuse in a vertical direction. Prior to a measurement, a vacuum is applied to desorb the sample of any impurities. Then, a testing gas is injected and absorbed in the polymer sample. Gas diffusivity is indirectly calculated by measuring the rate of gas volume or gas pressure changes after gas injection [125]. There are many different kinds of sensors such as laser and nuclear magnetic resonance sensors [122, 125] used in this method to get the graphs of gas volume vs. pressure.

**2.3. In fiber gratings based refractive index sensors**

To utilize fiber optic sensors for the measurement of solubility and diffusivity, in depth comprehension of different fiber optic refractive index sensors is necessary. In this section, in fiber gratings, namely FBG and LPG, based refractive index sensors are reviewed.

**2.3.1. Fiber Bragg grating based refractive index sensors**

A fiber Bragg grating (FBG) is a type of fiber grating that is used widely in sensing applications. FBG is fabricated by inscribing periodic refractive index in the fiber core, which reflects a particular wavelength of light to the direction of the light source due to its phase matching property. Usually, a grating has periods in the order of hundreds of nanometers, and thus is also called short period grating. A schematic of FBG is as shown in Figure 2-9 [126]. FBG behaves like a band stop filter using the principle of Bragg reflection phenomenon [127]. The formation of a periodic index modulation in the optical
fiber core allows particular wavelengths of light to be reflected while transmitting the rest of the light [128]. FBG fiber optics have been investigated for the telecommunication and sensing applications [129]. Furthermore, FBG sensors resulted in high performance sensing for individual or simultaneous measurement of temperature, strain, and refractive index [130-133]. The sample spectrum of FBG refractometer in response to different refractive index environment is shown in Figure 2-10 [126]. The strengths of FBG sensors in single mode fibers are that they offer real time response, accurate measurement, and allow multi parameter measurement. Fiber Bragg gratings can also be inscribed in in fiber interferometers to allow a configuration of Mach-Zehnder interferometer and improve in the performance [18].

![Figure 2-9. The schematic of a typical Fiber Bragg grating sensor][126]
Figure 2-10. The spectrum of a Fiber Bragg grating sensor in different refractive index environment [126]

When the core is exposed to an intensive optical interference pattern, a periodic perturbation of the refractive index along the fiber is formed, called fiber Bragg grating and the FBG is first developed onto optical fiber in 1978 at the Canadian Communication Research Center [134, 135]. The periodic perturbation behaves as a stop-band filter, where a narrow band of optical wavelength is reflected by the scattering from index variations. In FBG, the mode coupling occurs at the Bragg wavelength, $\lambda_B$ which can be obtained by:

$$\lambda_B = 2n_{eff}\Lambda$$ (1)
where \( A \) represents the period of the grating and \( n_{\text{eff}} \) represents the modal index. Each reflection is in phase with the next one as shown in Figure 2-11. Any change in fiber properties such as refractive index, strain, or temperature would change the modal index or grating pitch which would lead to change in Bragg wavelength [135].

\[
\lambda_B \quad \text{refl.} \quad \lambda - \lambda_B \quad \text{trans.}
\]

\[
\lambda \quad \text{inc.} \quad \text{spacing} = \frac{\lambda_B}{2n_{\text{eff}}}
\]

Figure 2-11. The principle of fiber bragg described by Hill et al. [135]. The Bragg resonance for reflection occurs at the wavelength which the grating pitch is one-half of the modal wavelength of the core. The transmitted wavelength would be resultant of wavelength subtracted by the Bragg reflection.

Single mode optical fiber with FBG is used widely used as a refractometer. The effective short period FBG sensor based on the resonance shift is successfully fabricated by C. Chan and researchers and they allowed a refractive index measurement in range between 1.25 and 1.44 [136]. Similarly, tilted FBG (TFBG) sensor based on intensity is investigated by Y. Miao, B. Liu, and Q. Zhao [137]. In case of a single mode fiber, the TFBG enhances the coupling of the light from forward propagating core to backward propagating cladding mode. By analyzing the transmission spectrum of the fabricated TFBG sensor, they observed that with higher refractive indices, the transmission power
decreased linearly as shown in Figure 2-12. Near the leaky configuration (i.e. $n_{SRI} \sim 1.4532$), the transmission spectrum was nearly diminished.

Figure 2-12. Transmission spectrum using TFBG fabricated by Miao et al. [137]

Numerous researchers endeavored to improve the quality of FBG refractive index sensors by applying more advanced techniques. For instance, modelling and characterizing of fiber Bragg gratings are investigated by Ugale et al in order to achieve the maximum reflectivity [128]. Modifications on FBG sensors also improved its performance. For example, the usage of thinned FBGs is researched for refractive index sensing because it has been found that reducing cladding diameter led to a non-linear dependence on surrounding refractive index and effective refractive index, $n_{eff}$ [138, 139]. The Figure 2-13 elucidates the difference in sensitivity to the surrounding refractive
index for the thinned FBG and unperturbed FBG sensors. As the surrounding refractive index increases, the thinned FBG offered more spectral shifts [138]. Another interesting modification on FBG is conducted by T.Gui, H. Tam, P.A. Krug, and J. Albert, in which they developed a novel in-fiber structured intensity based tilted FBG with a lateral offset as shown in [140]. Similar to the core mismatch MZI, lateral offset on FBG allowed cladding modes while still having the Bragg reflected core mode. Therefore, the proposed refractive index sensor uses both the core-mode reflection from the tilted FBG and the intensity based cladding mode from the lateral offset. However, the authors investigated that at high refractive index, the sensitivity of the novel proposed sensor decreases significantly.

Figure 2-13. The comparison in relative shifts between the thinned FBG sensor and unperturbed FBG sensor [138]
FBG sensor on different type of optical fibers are also used as a refractometer. The effect of microstructures in the FBG optical fibers on the measurement of refractive index has been studied by M.C. Phan Huy et al. [141]. Two types of microstructures optical fibers (MOFs, or also known as PCF), which are six hole and two-ring triangular MOF, are used for the study as shown in Figure 2-15. As seen from Figure 2-16, for the case of the six hole fiber, the experimental data did not fit with the model but this is because the core shape of the six hole fiber was not exactly circular when the model assumed that the core was circular. In case of two-ring triangular fiber, both multiple FBG mode and single FBG mode agreed well with the theoretical model. Moreover, ear the leaky configuration (i.e. $n \sim 1.45$) both MOF sensors resulted in poor resolution.
Figure 2-15. An image of (a) six holes and (b) two-ring triangular photosensitive microstructured optical fibers [141]

Figure 2-16. The resonance shift versus refractive indices for (a) six holes and (b) two-ring triangular microstructured optical fibers [141].

A unique refractometer based on a micro-slot in a FBG has been studied in 2007. K. Zhou and the research group proposed a liquid core waveguide based refractometer fabricated with a micro-slot in a FBG [142]. As illustrated in Figure 2-17, a micro-slot is fabricated across the FBG in order to allow liquid core principle. In this device, a liquid is used as a core of a waveguide rather than cladding. Therefore, at high liquid refractive index, the light exists mainly in the liquid, and the $n_{\text{eff}}$ is determined by the liquid. In other words, the refractive index higher than cladding could be measured with this device. K. Zhou and the research group first studied the reflection spectra of Figure 2-18(a) below the refractive index of silica fiber and Figure 2-18(b) above that silica. At lower ranges of refractive index, only the main Bragg peak is seen. However, as the
refractive index becomes higher than 1.45, new reflection peak can be seen as shown in the arrows. Moreover, the response of Bragg peak to the variation of the refractive index in terms of the wavelength is as illustrated in Figure 2-18(c). At the low refractive index regime, the sensitivity is same as the literature. However, as the refractive index becomes higher, the trend changes significantly. At the high refractive index regime, the slot initiates to support the propagation modes, which is dependent to the refractive index of the oil (since more peaks appear in Figure 2-18(b)) and peak is increased. As the refractive index continues to increase, the peak decreased due to decreasing coupling from un-etched fiber section. More importantly, a linear relationship is achieved when the wavelength shift is plotted with the varying refractive index above 1.46 as shown in Figure 2-18(d). The sensitivity of the device is found to be higher in the high refractive index region as shown in Figure 2-18(e).

**Figure 2-17.** A schematic of a liquid core waveguide based refractometer. The slot for liquid filling can be seen across the fiber Bragg gratings [142].
Figure 2-18. Reflection spectrum diagrams of the proposed liquid core waveguide based refractometer when the refractive index of the oil is (a) below and (b) above the refractive index of the optical fiber. The arrows in (b) corresponds to the fundamental mode of the liquid core waveguide and the inset focuses on these peaks. (c) Wavelength shift and change of the intensity of the main Bragg peak with respect to RI of the oil. (d) Experimental and simulated wavelength shift of the liquid core FBGs. (e) Sensitivity of the device over the low RI and high RI regime [142].

2.3.2. Long period grating based refractive index sensors

Since the fiber Bragg gratings offer higher sensitivity to strain and temperature and require expensive interferometric techniques, which are often a complex process, for determining wavelength shifts caused by the surrounding refractive index, long period
gratings became a fascinating topic for the researchers in measuring the surrounding refractive index. One of the feature of LPGs is to sense the refractive index of the surrounding material. This comes from the dependence of the attenuation band wavelengths on the effective index of refraction of the cladding modes, which is also dependant to the surrounding refractive index. J.H. Chong et al. have demonstrated a refractive index sensor in different refractive indices of sugar, ethylene glycol, and salt solutions, up to refractive index of 1.44, using the long period gratings [143]. As illustrated on Figure 2-19, the increase in refractive index on all sugar, ethylene glycol, and salt solutions have led the spectrum to shift to the left (i.e. decreased in wavelength).

![Figure 2-19](image_url)

**Figure 2-19.** The transmission profile of LPG in response to (a) sugar solution (b) ethylene glycol solution (c) salt solution and (d) all three plotted together [143]

The behavior of LPG to the surrounding refractive index after etching has also been studied. K.S. Chiang and co-investigators have described the shift in the resonance wavelength of LPG in response to etching of cladding and the change of surrounding
refractive index for the first time in 2000 [144]. Although Vasiliev et al. have already demonstrated that etching the LPG with HF acid led the resonance wavelength of LPG to shift to longer wavelengths, they did not study the effect of etched LPG to the variations of surrounding refractive indices [145]. Therefore, K.S. Chiang and the co-investigators developed analytical formulas for the wavelength shift of etched LPG in response to the surrounding refractive and verified the accuracy of the data with the numerical simulations and experimental results as shown in Figure 2-20. K.S. also discovered that the sensitivity of the variation of the surrounding refractive index increased as the cladding radius decreased as shown in Figure 2-21.

![Figure 2-20](image)

**Figure 2-20.** The wavelength shift in response to the surrounding refractive index [144]
The long period gratings on photonic crystal fiber (PCF) is also used as a sensor to measure the surrounding refractive index. The core of the PCF is in the form of a missing air channel in the center of the fiber while the cladding has microstructured array of air channel running along the fiber axis. Since PCFs allow wide spectrum of design possibilities form the variation of air channel geometry or lattice structure, the optical properties not known in typical single mode optical fiber could be investigated. LPG on PCF has been studied by numerous researchers. J.S. Petrovic et al have theoretically analyzed that PCF LPG is sensitive to strain and surrounding refractive index [146]. L. Rindorf et al. used PCF LPG as a refractive index transducer in measuring thickness of DNA [147]. Different methods are available to fabricate LPG into PCFs, such as UV imprinting, periodic deformation through CO₂ laser, or using the arc discharge [148-152]. In the work of Y. Zhu, Z. He, and H. Du, the authors compared the difference in the
transmission spectrum of LPG written in CO₂ laser in endlessly single mode (ESM) PCF and in single mode fiber [152]. The authors observed that the transmission spectrum of LPG PCF shifted to the right with increasing surrounding refractive index while the transmission spectrum of single mode LPG shifted to the left with increasing surrounding refractive index as shown in Figure 2-23. According to the research group, the equation (3) from Chapter 3, can also be expressed as [152]:

\[
\frac{\partial \lambda_i}{\partial n_{\text{sur}}} = \left[ \frac{\partial n_{\text{core eff}}}{\partial n_{\text{sur}}} - \frac{\partial n_{\text{cl eff}}}{\partial n_{\text{sur}}} \right] \Lambda
\]  

(2)

According to the equation (2), the positive wavelength dependence of the PCF LPG suggests that the dispersion of core and cladding modes (i.e. \( \frac{\partial n_{\text{core eff}}}{\partial n_{\text{sur}}} > \frac{\partial n_{\text{cl eff}}}{\partial n_{\text{sur}}} \)) dominates the phase matching condition [152]. Moreover, in the leaky configuration (\( n > 1.45 \)), the wavelength of single mode LPG shifts towards lower wavelengths and the intensity of the peak attenuation is reduced while the wavelength of LPG PCF is still positive and there are no noticeable change in the intensity of the peak attenuation. Similarly, H. Dobb, K. Kalli, and D.J. Web reported experimental results of sensing properties of both endlessly single mode (ESM) LPG PCF and large mode area (LMA) LPG PCF [153]. Similar to Y. Zhu’s experiment, the spectral shift of the LPG PCF is to the right with increasing surrounding refractive index. Moreover, at the leaky configuration as illustrated in Figure 2-22, the spectral shift do exist but experiences extremely low sensitivity, which would limit the usage in high refractive index applications. The intensity for LMA LPG PCF, however, continues to decrease, from the Fresnel reflections occurring at the cladding-SRI interface [153].
Figure 2-22. The transmission spectrum of the large mode area LPG PCF in response to high refractive index [153].

Figure 2-23. The transmission spectrum of (a) LPG PCF, (b) single mode LPG, (c) depicting wavelength shift with respect to surrounding refractive index and (d) depicting intensity change with respect to the surrounding refractive index.

The transmission spectrum of LPG in response to the leaky configuration has been investigated. H. Patrick et al. have experimentally obtained wavelength shift and
transmission of 275 um period LPG for refractive index of 1 < \( n_{\text{SRI}} \) < 1.72 [19]. Furthermore, Stegall et al. also have experimented wavelength shift and transmission of LPG with period 290 um for refractive index of 1 < \( n_{\text{SRI}} \) < 1.734 [31]. From Figure 2-24, the wavelength shift of LPG in leaky configuration do exist, but the sensitivity is not sufficient for use in high refractive index applications. Typical LPG in single mode optical fibers without any further processing, such as thin film layer coating atop optical fiber, high refractive index sensing is difficult.

![Figure 2-24. Transmission responses of LPG sensor in refractive range of 1<\( n_{\text{SRI}} \)<1.7 from two different literature [19, 31]](image)

2.4. Mach-Zehnder Interferometer as refractive index sensor

Mach-Zehnder interferometers (MZI) in many different configurations are widely used as the refractive index sensor. In all configurations of fiber optic interferometers, the processes of beam splitting and beam combining are required because the fiber optic interferometer operates using the interference between the two beams which have propagated through different optical paths. As a refractive index sensor, it is critical to have one of the optical paths to be positioned in a way so that the signal can be affected by the surrounding refractive index. Both intensity based and phase based fiber optic
interferometer sensors has been already researched in sensing applications. A phase shift
based refractometer is obtained using a pair of LPG [154]. The first LPG couples the
incident light guided to the core of a single mode fiber to the cladding mode of the fiber.
The coupled cladding mode is again recoupled to the core mode of the fiber by the
second LPG. As shown in Figure 2-25, the transmission wavelength of LPG MZI shifts to
the left (i.e. towards the lower wavelength) as the refractive index increases, similar to
the typical LPG refractive index sensor. However, the sensitivity of LPG MZI is low
compared to the typical LPG. The refractive index sensitivity of MZI is improved by
introducing tapers to MZI. The fiber tapering causes the change in the diameter of the
core mode, and therefore, portions of the core mode couples with the cladding mode. As
shown in Figure 2-25(b), the wavelength of tapered MZI shifts to the left, and having
three tapers improved the refractive index sensitivity. By introducing a taper into LPG-
MZI by J.F. Ding et al., the refractive index sensitivity of the LPG-MZI have been
substantially enhanced.

Figure 2-25. The transmission response of LPG-MZI and tapered LPG-MZI to the variation of
surrounding refractive index and (b) transmission response of taper-MZI to the variation of
surrounding refractive index [154, 155]
Having a mismatch in diameter is another configuration of MZI that has the capability to sense the surrounding refractive index. By splicing a smaller or larger core single mode fiber to a typical single mode fiber, a core diameter mismatch MZI is configured, and in this case, the light in the core mode would couple to the cladding mode of the smaller fiber at the splice point and would recouple with the uncoupled core mode at the other end of the splice point as shown in Figure 4-3(b). In 2012, Q. Rong and the research group developed a phase shift based high sensitive refractive index in fiber MZI by sandwiching a single mode fiber in between two thin-core fibers (TCFs) as illustrated in Figure 2-26(a) [156]. The wavelength of TCF-SMF-TCF MZI shifted to the right (i.e. towards higher wavelengths) as it is emerged to the higher surrounding refractive index, which could be seen from Figure 2-26(b). Also, the shorter length of SMF sandwiched in between the TCF led to higher sensitivity in comparison to the longer ones. A refractive index sensor based on core offset has been investigated by Q. Yao et al. [157]. In their core offset configuration of MZI, FBG is implemented to control the number of cladding modes in the MZI. As illustrated in Figure 2-27, the amplitude of the measured transmission intensity decreases with increased refractive index. The phase, on the other hand, shifts to the right (i.e. increases in wavelength) by 0.4 nm as the surrounding refractive index increased by 0.025 [157].
Figure 2-26. A phase shift based high sensitive refractive index in fiber MZI by Q. Rong and the research group [156]. (a) A schematic of the diameter mismatch MZI and (b) wavelength shift in response to varying surrounding refractive index.

Figure 2-27. A refractive index sensor using the core offset MZI configuration by Q. Yao [157]. The refractive index sensing using photonic crystal fibers are prevalently used in the MZI configuration. PCF is not only used because its microstructure allows a variety of different configurations, but also is used because PCF sensors survive in the harsh environments, such as in high temperature and pressure. Therefore, although PCF offers higher losses compared to single mode fibers, the development of PCF as sensors is continued. In order to achieve a PCF MZI, PCF must be fusion spliced onto the single mode fiber. During the fusion splicing, a small segment of PCF collapses, diffracting some of the core mode light to couple to the cladding mode. A simple and compact PCF MZI refractive index sensor is developed by R. Jha and co-investigators [158].
investigators have successfully spliced a LMA PCF into a single mode fibers with collapsed regions, and used two devices, one with length of 17 mm and the other with 32 mm, to observe the behavior of transmission spectra with respect to varying refractive index. As shown in Figure 2-28, the wavelength shifts to the right and the amplitude of the transmission intensity increases with increased surrounding refractive index, and the 32 mm length device experience larger values of the wavelength shift [158, 159]. The PCF MZI offered higher sensitivity near the refractive index of the fiber, and above that value, according to R. Jha et al., the cladding mode became effectively a radiation mode and the interference pattern disappeared. In low regions, ranging from 1.33 to 1.35, the shift still existed and the sensitivity value was similar to the work by M. Deng et al. [160]. Moreover, according to H.Y. Choi et al, the problem with high loss in PCF, which is a material property, is reduced by introducing LPG into PCF MZI [161, 162].

![Figure 2-28](image-url)  
**Figure 2-28.** Change in transmission spectra of PCF MZI with respect to varying refractive index [158].

2.5. **Intensity based fiber optic sensor as refractive index sensor**

A cost effective, simple, and compact in size optical fiber refractive index sensor can also be fabricated by splicing different types of optical fibers. An intensity based
refractive index sensor is developed by Joel Villatoro and David Monzon-Hernandez, using a diameter mismatch [163]. In this case, J. Villatoro and D. Monzon-Hernandez used a multimode-single mode-multimode (MSM) structure to fabricate the index sensor. The principle of the structure is similar to MZI, where the propagating core more is coupling to the cladding mode due to diameter mismatch. However, the SMS or MSM is differentiated to different section from MZI because there is a new field of study based on the intensity based and phase shift based SMS or MSM configuration index sensor. In the MSM index sensor developed by J. Villatoro, the core diameter mismatch is obtained very economically and simple because all it required is splicing a single mode fiber in between the two multimode fibers. As illustrated in Figure 2-29, J. Villatoro observed increase of the amplitude of transmission intensity with the increased surrounding refractive index. In the leaky configuration (n>1.44), the change in transmission is still observed.

![Figure 2-29. Intensity based transmission spectrum in MSM configuration [163]](image)

A refractive index sensor based on single mode-multimode-single mode (SMS) configuration has also been investigated. Two research groups, one led by P. Wang and the other led by Q. Wu, developed a wavelength shift based refractometer by splicing a
multimode fiber in between the two single mode fibers [164, 165]. From both studies, the increase in surrounding refractive index caused the wavelength to shift to the right, as can be seen from Figure 2-30. Moreover, Q. Wu did analysis on the effect of the multimode fiber diameter on the sensitivity and discovered that smaller multimode fiber diameter increased the sensitivity. In both studies, the refractive index sensitivity is higher near the refractive index of the cladding (n~1.44).

![Figure 2-30. Change in wavelength shift with respect to surrounding refractive index [164, 165]](image)

In 2014, a similar type of refractive index sensor to SMS is developed by M. Shao et al. [166]. The difference of this refractive index sensor to the SMS configuration is that first, single mode fiber is spliced to the thin cored fiber, which is spliced to the multimode fiber core at the other end and then spliced back to the single mode fiber resulting a STMS configuration, as illustrated in Figure 2-31. With a STMS configuration, M. Shao et al. observed a wavelength shift to the right in the increasing surrounding refractive index and observed a decrease in intensity with rising surrounding refractive index. Furthermore, the researchers conducted a parametric study on the length of the thin cored fiber, length of the multimode fiber core, and the diameter of the multimode fiber core with the varying surrounding refractive index and discovered that 8 mm L_{TCF}, 15 mm
L_{MMFC}, and 90 μm D_{MMFC} resulted in highest refractive index sensitivity as shown in Figure 2-32 [166].

![Figure 2-31. A schematic of a single mode-thin core-multimode-single mode fiber configuration](image1)

![Figure 2-32. Transmission spectra of the STMS configuration with increasing surrounding refractive index](image2)

A wide range of surrounding refractive index has been studied using a multimode-coreless silica fiber (CSF)-multimode configuration. Y. Jung et al. have succeeded in fabricating a refractive index sensor by splicing multimode-CSF-multimode fibers together as seen in Figure 2-33. A unique aspect of this configuration is the ease of coupling to a source or another fiber due to large core diameter. In this configuration, the core modes from MMF1 are coupled to the cladding modes of the CSF and at the second splice point, the cladding modes are recoupled to the core modes of MMF2 [167]. Y. Jung et al. analyzed spectrum of the MCM configuration from wavelength of 400 nm to
1600 nm and categorized into two regions, where region I is from 700 nm to 1100 nm and region II from 1200 nm to 1450 nm as shown in Figure 2-34. In the region I, Y. Jung et al. did observed change of intensity with varying refractive index. The researchers were able to sense change of refractive index using the intensity based MCM configuration sensor. Interestingly, the sensitivity of the region I for MCM configuration increased in the leaky mode. In the region II, the wavelength shift behavior with varying surrounding refractive index was observed. As seen in Figure 2-34, the wavelength shifts to the right with respect to the surrounding refractive index in the range of $1.30 < n_{SR1} < 1.44$. However, in the region II, as soon as the leaky configuration is achieved ($n_{SR1}$~1.45), the wavelength shift did not occur and there were no noticeable change in the region II.

![Figure 2-33. The schematic of multimode-coreless-multimode refractive index sensor [167]](image-url)
Figure 2.34. The transmission spectra of the MSM configuration in region I and region II with varying surrounding refractive index [167]

2.6. High refractive index sensors

As been discussed a number of times, fiber optic refractive index sensor encounters a decrease in sensitivity when measuring surrounding refractive index that is higher than that of the cladding (i.e. leaky configuration). In this section, a post process, mainly coating, on fiber optic sensor to improve sensitivity in leaky configuration has been reviewed. Since coating of the optical fiber is very complex and is a distinct field of research, only the basic principles would be discussed in this section. It has been investigated previously both experimentally and theoretically that LPG’s central
wavelengths of the core-cladding mode coupling bands depend with the optical thickness of high RI coatings. At a certain thickness, one of the low order cladding modes is phase match to a mode of the waveguide formed by the coating [168-171]. This is called mode transition region, where the cladding modes are reorganized with each mode taking the properties of its adjacent lower order cladding mode [168]. Therefore, by adjusting the optical thickness of the coating so the LPG operates in this mode transition region improves the sensitivity of the cladding mode effective index and thus the resonance wavelengths to the surrounding refractive index [169]. L. Coelho et al. have developed a fiber optic refractive index sensor based on a LPG that has been coated with a titanium dioxide (TiO2) thin film as illustrated in Figure 2-35. The thermal evaporation of pure titanium in a controlled oxygen atmosphere is used to deposit thin film of TiO2 onto the LPG. As seen from Figure 2-36, there are no noticeable wavelength shift in uncoated LPG in refractive index range of 1.46 to 1.6. However, by depositing 40 nm layer of TiO2 onto the LPG sensor, wavelength shifts occurred from refractive index of 1.46 to 1.6 (i.e. leaky configuration). Moreover, the changes in sensitivity with varying coating thickness has been investigated by the same research group as shown in Figure 2-36, and showed that different coating thickness could be used depending on the desired range of surrounding refractive index [172]. However, this coating on LPG require very uniform coating throughout the coated region, and the thickness of the coating must be controlled in a nanometer scale, which is a complex and expansive procedure.
2.7. Conclusion

The measurement devices used for obtaining solubility and diffusivity of gas/SCF in polymers are expansive, complex, and take a long experimental time. A new potential solubility and diffusivity measurement sensor has been suggested using in fiber gratings
and fiber optic interferometers. In this chapter, most widely used solubility measurement
devices and diffusivity measurement devices reviewed. Then, the refractive index sensors
fabricated by in fiber gratings and in line interferometers are reviewed as well to validate
the feasibility of using fiber optic sensor for the measurement of solubility and diffusivity
of gas/SCF in polymers. Since refractive index of most polymers are higher than the
cladding, the fiber optic refractive index sensors must be capable of operating in leaky
configuration. Therefore, literature reviews on high refractive index sensors are also
included in this Chapter.
Chapter 3. Long period grating sensor for measuring high surrounding refractive index solution

The LPG sensors are one of the most frequently used fiber optic sensor in measuring refractive indices. As mentioned in Chapter 2, bare LPG sensors cannot be used when the surrounding refractive index becomes higher than that of the cladding (i.e. the leaky configuration). Hence, a high refractive index thin layer coating atop the LPG sensor enables LPG sensors to operate even in the leaky mode. To improve the cost efficiency and the sensitivity in the leaky mode, especially in the range of 1.48 to 1.55, for the potential application of measuring solubility and diffusivity in polymers, a new type of coated LPG sensor has been suggested. The LPG has been grated using the femtosecond laser, and it is coated with silver nanoparticles using the atomization based coating system. In this Chapter, the feasibility of silver nanoparticle coated LPG for measuring solubility of gas/SCF in polymer is tested by experimenting the measurement of refractive index in the leaky configuration.

3.1. Introduction to long period grating sensor

While FBGs have periods in the order of hundreds of nanometers, long period fiber gratings (LPG or LPFG) offer periods in the order of 100 micrometers extending to a millimeter. A typical long period grating schematic is illustrated in Figure 3-1 [17]. Due to the gratings, the guided core mode couples to the forward propagating cladding modes, where it is affected by the effect of absorption and scattering. The discovery of LPG, has captured attentions from numerous researchers and have impacted the development of the fiber sensing technology. Because coupled light from core mode into forward
propagating cladding mode is dependent to the surrounding environmental conditions, the change in temperature, pressure, or refractive index cause change in the light propagation properties of the long period gratings. Hence, LPG sensors are widely used by researchers for different applications. A sample spectrum through LPG sensor can be seen from Figure 3-2. The peak in the plot changes with respect to the environmental conditions and therefore makes a very reliable sensor.

![Schematic of a long period grating sensor](image)

**Figure 3-1.** The schematic of a long period grating sensor [17]

![Spectrum of the long period grating sensor](image)

**Figure 3-2.** A spectrum of the long period grating sensor
Different fiber optic sensors exist and they are used in numerous applications for diverse purposes. As discussed above, in fiber gratings sensor and in line optical fiber interferometers offer different configurations, and each have ability to sense different measurand, such as temperature, strain, pressure, and refractive index. This final thesis mainly focuses on the feasibility of fiber grating sensors and optical fiber interferometer sensors as the high refractive index sensors because most of the refractive indices of the polymers are higher than the refractive index of cladding, which is approximately 1.44.

In the LPG refractive index sensing, the main condition for phase matching between the forward propagating core mode and cladding mode, which is known as phase matching condition, can be described by the following equation [173]:

\[
\lambda_M = \left[ n_{\text{eff}}^{\text{core}}(\lambda) - n_{\text{eff}}^{\text{cl}(m)}(\lambda) \right] A_L
\]  

(3)

where \( n_{\text{eff}}^{\text{core}} \) represents the effective index of the core mode, \( n_{\text{eff}}^{\text{cl}(m)} \) represent the effective index of the \( m^{th} \) order cladding mode, and \( A_L \) represent the period of the grating. Because the cladding interacts with the environment, the cladding modes can move from the fiber into the environment due to absorption and scatterings. Therefore, LPGs are used extensively as a tool for refractive index sensing. The coupling of wavelengths for the fundamental core mode and forward propagating cladding modes, is the refractive index sensing principle of the LPG sensor. From Equation (3), the coupling resonant wavelength is a function of the effective indices of the guided core and cladding modes and the grating period. The effective indices are dependent with the fiber core and cladding indices and radii. Moreover, the effective indices of the cladding modes are strongly dependent to the surrounding refractive index \((n_3)\). The change in surrounding
refractive index leads to a change in the effective index \( n_{e\text{ff}}^{cl} \) of the cladding modes. The higher order modes results in greater index modifications. The coupling wavelength corresponding to a particular cladding mode depends on \( n_{e\text{ff}}^{cl} \), which is dependent to the surrounding refractive index \((n_3)\). Therefore, a change in the surrounding refractive index \((n_3)\) will cause a shift in the value of \( \lambda_{PM} \). Typical LPG refractometer restricts the refractive index of a SRI to a value less than the effective index of the cladding mode (i.e. \( n_3 < n_{e\text{ff}}^{cl(m)} \)). In a single mode fiber, the fundamental mode is well guided into the fiber core without any changes from the surrounding refractive index. Moreover, if temperature, pressure, and strain is kept constant, the grating period should remain constant. Thus, the shift of resonance cladding mode with respect to the effect of the surrounding refractive index \((n_3)\) \((d\lambda_M/dn_3= 0)\) is given by [174]:

\[
\frac{d\lambda_M}{dn_3} = \frac{d\lambda_M}{dn_{e\text{ff}}^{cl}} \frac{dn_{e\text{ff}}^{cl}}{dn_3}
\]  

(4)

For the typical operational range, an increase in effective index of the cladding mode results in a negative shift of the coupling wavelength \((d\lambda/dn_{cl}) < 0\). As a result, a typical response of LPG to the surrounding refractive index shows a negative shift of a transmission dip as it encounters with higher surrounding refractive index than air (i.e. \( n_{\text{SRI}} > n_{\text{air}} \)).

3.2. Long period grating sensor in high refractive index

In fiber gratings, the wavelength coupling from the core mode to the cladding modes is dependent on the difference between the effective core index and the effective cladding
index, the radii of core and the cladding, and the grating period. Changing any of these parameters cause shift in the transmission spectrum. According to R. Hou and the co-investigators, the cladding mode profiles are also sensitive to changes in the surrounding refractive index at leaky configuration. The published experimental data by H.J. Patrick, A.D. Kersey, and F. Bucholtz show that LPG both transmission spectrum shifts and intensity changes due to the effect of Fresnel reflection even in the range of 1.40 to 1.72 despite the fact that sensitivity is extremely low [19]. Although the aforementioned literature data have not reported any analysis on spectral shifts, Hou and the researchers have discussed the three cases of $n_{SRI} < n_{cl}$, $n_{SRI} \approx n_{cl}$, and $n_{SRI} > n_{cl}$, and analyzed the spectral shift in case of $n_{SRI} > n_{cl}$ using their mathematical model. The simulated data by Hou and the researchers are shown in Figure 3-3. Cladding modes are most accurate when calculated using the three-layer model [175] but this is feasible only for the case of $n_{SRI} < n_{eff,cl} < n_{cl}$. Therefore, the two-layer model is used by Hou and the researchers to calculate the cladding modes in their simulations, which provided reasonably accurate values. The two-layer model treats cladding and core as one multimode fiber and the environment as the new cladding only in the LPG region [27]. In the Figure 3-3, the wavelength shift in the nonlinear region of $1.475 < n_{SRI} < 1.54$ is too small to be calculate probably because of the transition from total internal reflection mode to a leaky configuration mode. At $n_{SRI} = 2.666$ the wavelength shifts become +1 nm. Even though these spectral shifts are very small, they should be able to be measuring using a high resolution optical spectrum analyser. However, the spectral shifts in the leaky mode is very small and would still have limitation as a refractive index sensor in the region of the $n_{SRI} > n_{clad}$. 
Figure 3-3. Simulated data of coupling wavelength shift against the surrounding refractive index greater than refractive index of the cladding [27]

3.3. Coated Long period grating sensor

Fiber optic refractive index sensors encounter a decrease in sensitivity when measuring surrounding refractive index that is higher than that of the cladding. It has been known that coating on top of the fiber optic sensor to improve sensitivity. LPG’s central wavelengths of the core-cladding mode coupling bands depend with the optical thickness of high RI coatings. At a certain thickness, one of the low order cladding modes is phase match to a mode of the waveguide formed by the coating [168-171]. This is called mode transition region, where the cladding modes are reorganized with each mode taking the properties of its adjacent lower order cladding mode [168]. Therefore, by adjusting the optical thickness of the coating so the LPG operates in this mode transition region improves the sensitivity of the cladding mode effective index and thus the resonance wavelengths to the surrounding refractive index [169]. As discussed before, Coelho et al.
has successfully measured high refractive index using the nanolayer deposition of TiO$_2$ on LPG sensors [172]. Nanoparticle coating on fiber optic sensors has been recently investigated. The deposition of nanoporous TiO$_2$/Polyion on LPG sensors have been found to improve sensitivity of glucose solutions in comparison with uncoated LPG sensors at a thickness of approximately 230 nm [176]. Similarly, the deposition of SiO$_2$ nanoparticles and Au nanoparticles on LPG sensors are researched by Sergiy et al. and Tang et al., respectively, and observed improvement in the refractive index sensitivity [169, 177]. In this thesis, the effect of silver nanoparticles on LPG sensor is of interest.

3.4. Fabrication of long period grating sensor using the femtosecond laser

The femtosecond laser (Spectra-Physics ultrafast Ti: Sapphire laser) shown in Figure 3-4, was used to create RI change in the fiber core. The kilohertz ultrafast laser has a pulse width of 120 femtosecond with the center wavelength of 800 nm. A computer controlled 4-axis stage was used for aligning and scanning the fiber with laser radiation. The schematic of experimental setup for the writing long period gratings in a single mode fiber is illustrated in Figure 3-5. A set of neutral density (ND) filters was used to primarily tune down the pulse energy. After the ND filters, a computer controlled variable attenuator was used to set laser output power to the desired amount. A computer controlled electronic shutter was used for turning on/off the laser beam selectively to write index in the desired location at the fiber core. To obtain the beam diameter of 1.5 mm, an iris diaphragm was used. Then the laser beam was guided into a microscope and focused by objective lens (Numerical aperture: 0.55) into the core. The fabrication of gratings with femtosecond pulses were observed through a CCD camera mounted above
the dichroic mirror. Fiber jacket was removed using a jacket stripper and was rinsed with acetone in prior to writing LPGs in the bare fiber. The software, GOL3D was used to generate scanning path of the laser head. The period of refractive index modulation is 433 µm. In the index modified area, the refractive index was written for a length of 100 µm using the laser scanning (period: 2 µm) at the speed of 50 µm/sec. 42 periods are fabricated over 18.186 mm with power of mW. The fiber was coupled with a broad band light source (AFC BBS-1550) and an optical spectrum analyzer (PHOTONETICS Walics) to monitor the process [127].

**Figure 3-4.** (a) The femtosecond laser system and (b) the computer controlled 4-axis stage used for aligning fiber [127]

**Figure 3-5.** A schematic of the setting used for fabricating LPG [127]
3.5. Silver nano particle deposition using the atomization based coating system

Spray coating of silver nano particles was performed on femtosecond laser grated LPG sensors. An ultrasonic atomizer was utilized to create a mist of fine droplets. The ultrasonic atomizer is a very cost efficient method of coating because it converts the electrical energy into vibrational energy by means of a piezoelectric crystal that generates standing waves within the liquid layer that is present on its surface. When the amplitude of waves is increased such that a critical height is attained, the waves become unstable and fine droplets are detached from the crests and are carried to the nozzle by forcing air into the enclosed chamber where they are generated. This leads to spray formation, which achieves high velocity by second air stream flowing through nozzle. The nozzle is connected to NC based machine to facilitate the coating process. The schematic of the atomization based coating system is illustrated in Figure 3-6.
In this specific work, spray coating was performed with 42 inches per minute of feed rate at a distance of 13 mm from surface, while the air pressure was 145 kPa. 20 cycles of spray coating were performed on the LPG sensor. The resulted coating was then sintered in the furnace at 250 °C for 150 minutes to prevent the coating from wearing off during the refractive index experiment.

3.6. Experiment setup

Both uncoated and silver nanoparticle coated LPG sensors fabricated from the femtosecond laser has been tested under high refractive index solutions of 1.4823,
The olive oil, with refractive index of 1.458, and the cinnamon leaf oil, with refractive index of 1.5365, are mixed together to prepare a range of high refractive index solutions of 1.4823, 1.4944, 1.5094, and 1.5365. Abbe Refractometer is used for testing the refractive index of the solutions. The schematic of the experiment is as shown in Figure 3-7. The broad band light source (AFC BBS-1550) is connected to a FC/APC single mode fiber connector which is fusion spliced onto a LPG sensor. The other end of the LPG sensor is fusion spliced to another FC/APC single mode fiber connector which is connected to the optical spectrum analyzer (PHOTONETICS Walics). In air, at full configuration, the LPG peaks at 1550 nm with about -22 dBm. The refractive index solution is introduced the region shown in Figure 3-7. At steady state, the spectrum is saved, and after, the LPG sensor is cleaned and rinsed with acetone until the original spectrum is obtained in air.

**Figure 3-7** A schematic of the refractive index experiment using LPG sensor.

### 3.7. Behavior of silver nanoparticle coated long period grating sensor at high refractive index

The change in transmission spectrum of uncoated LPG in refractive index of 1.4944 and 1.5365, which is the refractive index of most polymers, has been observed in the experiment setup discussed above. As expected, with uncoated LPG, there is no noticeable wavelength shift from a transmission spectrum with the leaky configuration.
(surrounding refractive index >1.45). As shown in Figure 3-8, with surrounding refractive index of 1.5365, the transmission spectrum shows a dip of -9 dBm and at surrounding refractive index of 1.4944, the transmission spectrum shows a dip of -7 dBm. Only the amplitude of the transmission power increases by approximately 2 dBm with the refractive index increment of about 0.04.

![Graph](image.jpg)

**Figure 3-8.** A transmission spectrum for uncoated LPG in the leaky configuration. No noticeable shift in wavelength is seen.

In order to investigate the effect of silver nanoparticle coating on high surrounding refractive index, 7 and 20 passes of the silver nanoparticle coating deposited LPG sensor has been tested in refractive index of 1.4823, 1.4944, 1.5094, and 1.5365. The transmission spectrum of the 7 passes of coated LPG of coated LPG resulted as shown in Figure 3-9. For 7 passes of coated LPG, no visible wavelength shifts occurred, similar to the uncoated LPG. Only the transmission intensity changed from -4 dBm, -4.5 dBm, -6.5 dBm, and -7.5 dBm for refractive indices of 1.4823, 1.4944, 1.5094, and 1.5365,
respectively. After deposition of thin layers of silver nanoparticles, the transmission dip decreased by about 3 dBm.

![Figure 3-9](image-url)

**Figure 3-9.** A transmission spectrum for 7 passes of silver nanoparticle coating deposited LPG in the leaky configuration. No noticeable shift in wavelength is seen.

Interestingly, when 20 passes of silver nanoparticle is deposited to LPG sensor, the wavelength shifted to the left with increasing value of surrounding refractive index. As shown in Figure 3-10, about 0.7 nm wavelength shift occurred in refractive index ranges of 1.4823 to 1.4944, 0.3 nm from 1.4944 to 1.5094, and about 0.3 nm shifts has been observed from 1.5094 to 1.5365. There was approximately an overall wavelength shift of 1.3 nm towards to shorter wavelengths (i.e. shift to the left), which corresponds with the literature, from surrounding refractive index range of 1.4823 to 1.5365. This value of wavelength shift is significant considering back to the Figure 3-3, where no significant wavelength shifts has occurred in the simulated transmission spectrum of LPG sensor in leaky configuration. In case of the transmission intensity, not much change has occurred, and the transmission dip plateaued out at 7.5 dBm. If sensitivity could be optimized
further with coating, the silver nanoparticle coated LPG sensors could be feasible for obtaining solubility and diffusivity of gas/SCF in polymers.

Figure 3-10. About 0.7 nm wavelength shift occurred in refractive index ranges of 1.4823 to 1.4944, 0.3 nm from 1.4944 to 1.5094, and 0.3 nm from 1.5094 to 1.5365.

3.8. Conclusion

The behavior of femtosecond laser written LPG sensor in high refractive index has been investigated through a series of experiments. The cost efficient coating method based on the atomization coating system is implemented to deposit thin layers of the silver nanoparticle to the LPG sensor. The first test is done with uncoated LPG, which showed no noticeable wavelength shift. Similarly, 7 passes of silver nanoparticle coated LPG did not have any wavelength shifts. However, with 20 passes of silver nanoparticle coating on the LPG sensor, the transmission spectrum shifted towards shorter wavelength with increasing wavelength. From the refractive index of 1.4823 to 1.5365,
approximately 1.3 nm wavelength shifted to the left. This is significant because the wavelength shift occurred in the leaky configuration, where the total internal reflection is lost in the propagation mode. With further optimization with the coatings, the silver nanoparticle coated LPG could be used as a simple and cost efficient method to measure solubility and diffusivity of gas/SCF in polymers.
Chapter 4. Coated Mach-Zehnder Interferometer for measuring high surrounding refractive index solution

The numerous investigations on Mach-Zehnder interferometers as refractive index sensors in different configurations have been conducted by many researchers, as mentioned in Chapter 2. In MZI, the optical path differences between two modes, one reference arm and the other sensing arm, in the optical fiber eventually recombines and gives different interference patterns with the environment. In this Chapter, attempts to sense the refractive index in the range of 1.48 to 1.55, for the potential application of measuring solubility and diffusivity in polymers, has been made using a smaller core MZI configuration, PCF MZI, and silver nanoparticle coated PCF MZI configuration. The same atomization based coating system used for coating LPG was utilized to deposit thin layers of silver nanoparticles. The validity of the aforementioned MZI for measuring solubility of gas/SCF in polymers is tested by sensing for the measurement of refractive index in the leaky configuration.

4.1 Mach-Zehnder interferometer sensor

The fiber optic interferometric sensors offer distinctive properties such as a miniature sized configuration, immunity to electromagnetic wave, and high sensitivity detection. Therefore, interferometric sensors have become popular over conventional sensors for applications such as environmental monitoring [127]. In-fiber Mach-Zehnder interferometers, in particular, offer high sensitivity not only to temperature, strain and pressure, but also to surrounding refractive index changes. In all types of fiber optic
interferometers, the processes of beam splitting and beam combining are required because the fiber optic interferometer operates using the interference between the two beams which have propagated through different optical paths of a single fiber or two different fibers [161, 178]. It is critical to have one of the optical paths to be positioned in a way so that the signal can be affected by the external perturbations. The interferometers send out sensing information as a signal, which can be analyzed either in phase or intensity, and performs with high accuracy and high sensitivity. Both intensity based and phase based fiber optic interferometer sensors has been already researched in sensing applications [179, 180]. Moreover, the miniaturization is a current trend in fiber optics and, the bulky optical instruments such as beam splitters and combiners can be replaced by having fiber optic interferometers as in-line structures. In line structure, which have two optical paths in a single line, also offers high stability and coupling efficiency.

According to B.H. Lee and the research group, types of fiber optic interferometers could be categorized into four, which are the Fabry-Perot, Mach-Zehnder, Michelson, and Sagnac, and the fabrication methods and the operational principles for each of the interferometers are different for each of them [161, 178, 181-184]. Out of the four categories, this final thesis focuses only on Mach-Zehnder interferometer sensors.

Mach-Zehnder interferometers (MZIs) are diversely used in fiber optic sensors because they provide numerous easy and economical configurations. General schematics of the MZI can be seen from Figure 4-1. In a MZI, an incident light from the light source is split into two independent arms by a fiber coupler. One arm is a reference arm, which is kept isolated from the external perturbation, and the other arm is the sensing arm, which is exposed to the external perturbation. Then, the propagated incident light is recoupled by
the second fiber coupler, and the recoupled light has interferences. The interferences of the recoupled light heavily depends on the optical path distances between the two arms, and the external perturbation on the environment such as the temperature, pressure, or refractive index changes the optical path distances of the MZI, resulting in change of intensity or shift in phase or wavelength.

![Diagram of Mach-Zehnder Interferometer](image)

**Figure 4-1.** A schematic of the Mach-Zehnder interferometer. (a) a light source (b,e) couplers (c) a reference arm (d) a sensing arm (f) an optical spectrum analyzer

As mentioned before, the reason MZI is used prevalently in fiber optic sensing is that it can be easily configured. A miniature and compact in line MZI can be fabricated by having a recoupled light and uncoupled light intersect at the core, causing interference. For instance, an in line structure MZI is obtained by having a pair of LPG as shown in Figure 4-2. The incident light guided to the core of a single mode fiber is coupled to cladding mode of the fiber by the first LPG. The coupled cladding mode is then recoupled to the core mode of the fiber by the second LPG. T. Allsop and co-investigators have demonstrated a high sensitivity refractometer based on the in-line fiber long period grating Mach-Zehnder interferometer in 2002 [185].
MZI can also be fabricated by splicing another single mode fiber to a single mode fiber, if only the core of the two mismatches. As seen in Figure 4-3a, an offset in splicing the two single mode fibers makes MZI. Some of the light in core mode would be coupled to cladding mode and some would stay uncoupled due to the offset. At the other end of the offset splice, the cladding mode would recouple with the uncoupled core mode light. W.C. Wong and the research group have investigated the use of phase shift based core-offset MZI for refractive index sensor and suggested that having a rounded tip caused higher order cladding mode which resulted in higher refractive index sensitivity [186]. Similarly, splicing a smaller single mode fiber to a single mode fiber would also create MZI, as shown in Figure 4-3b. In this case, the light in the core mode would couple to the cladding mode of the smaller fiber and recouple with the uncoupled core mode at the end of smaller fiber. A sample spectrum of 40 um MZI sensor used in refractive index sensing is illustrated in Figure 4-4. Similarly, Q. Rong et al have used thin-core fibers with single mode fibers to demonstrate the core diameter mismatch MZI refractometer [156]. The core diameter mismatch can be used not only with single mode fibers, but also with multimode and hollow fibers [187, 188]. The principle of the MZI configuration is the same with the core diameter mismatch in this case, where the mismatch in the core diameter couples core mode lights into the cladding mode light. Another simple yet cost effective way to configure a MZI is by tapering the optical fiber. As shown in Figure 4-5,
the fiber tapering causes the change in the diameter of the core mode, and therefore, portions of the core mode couples with the cladding mode. However, due to the fiber tapering, the tapered MZI becomes very fragile near the taper. Tapering can be implemented with different types of MZIs and even with in fiber gratings [18, 154, 189-194].

**Figure 4-3.** A schematic of (a) core mismatch and (b) small single mode fiber Mach-Zehnder interferometer

**Figure 4-4.** A sample spectrum of 40 um Mach-Zehnder interferometer sensor
Different types of fibers could be spliced to the single mode fibers and configure MZI. Photonic crystal fibers (PCF) have a period pattern of microscopic voids throughout the fiber, shown in Figure 4-6, and the design of this microstructure grants the unique modal and guidance properties [158]. By fusion splicing a PCF with the single mode fiber, MZI can be configured. During the fusion splicing, a small segment of PCF collapses and this diffracts the core mode light to couple to the cladding mode resulting in MZI configuration, as shown in Figure 4-7. Zheng et al. have successfully fabricated a PCF MZI strain sensor that is insensitive to temperature and refractive index, and they used wavelength shift to sense the strain [195]. The drawback of the PCF MZI is that the losses are high compared to other offset methods. However, in line PCF MZI sensors can endure harsh environments such as high temperature and is used widely in numerous applications [195-198].
As discussed above, MZI is capable of measuring temperature, strain, pressure, and refractive index using the reference and the sensing arm. In case of the SRI sensing, the sensing arm is exposed to the varying refractive index while the reference arm is isolated. Changes in the signal from the sensing arm caused by varying refractive index modifies the optical path difference of the MZI and can be analyzed by observing the change in the interference pattern. Not much investigation has been done on MZIs in leaky configuration. If any, only very limited number of studies reported in MZI based refractive index sensors in the leaky configuration because the response of MZI to varying surrounding refractive index ceases to be linear in this domain [33].

4.2. PCF MZI sensor in high refractive index

In PCF Mach-Zehnder interferometer under guided configuration (i.e. SRI < 1.45), the fundamental single mode fiber mode diffract as it enters the collapsed PCF region, and from this diffraction, the mode broadens with the excitation of core and cladding modes in PCF region. Since the PCF core and cladding modes’ propagation constant differ, the modes obtain different phase difference throughout the propagation. The phase difference

![Figure 4-7. A schematic of photoic crystal fiber Mach-Zehnder (a, e) single mode fiber (b, d) collapsed region (c) photonic crystal fiber](image-url)
depends with wavelength of the guided light, $\lambda$, and the distance of the modes travel, $L$. (i.e. PCF length). Once the modes reach the second collapsed region of PCF, they further diffract and recombine [159]. The transmission of the PCF MZI can be expressed as the two-mode interferometer [159, 199]:

$$I = I_1 + I_2 + 2\sqrt{I_1 I_2} \cos \left( \frac{2\pi \Delta n_{\text{eff}} L}{\lambda} \right)$$

(5)

where $I$ refers to the intensity of the total interference signal, $I_1$ and $I_2$ are the intensities for the core and the cladding modes, respectively, and $\Delta n_{\text{eff}}$ is the difference between the effective refractive indices for core and cladding modes as:

$$\Delta n_{\text{eff}} = (n_{\text{eff}}^{\text{core}} - n_{\text{eff}}^{\text{clad}})$$

(6)

The variation in the surrounding refractive index would change the effecting refractive index of the cladding mode. Thus, according to eqns (5, 6), the transmission of the MZI would be affected with varying surrounding refractive index [200]. The length of the PCF is critical as well since not only it is dependent with the transmission of the PCF MZI, but also it is in relation to the fringe period, $\Lambda$, of MZI [198]:

$$\Lambda = \frac{2\pi \lambda}{(\beta_1 - \beta_2) \cdot L}$$

(7)

where $\lambda$ is the central wavelength of the light source, $\beta_1$ and $\beta_2$ are the propagation constants of the core and cladding modes, and $L$ is the length of the PCF region.

According to J-N Wang and J-L Tang, the longer PCF region would lead to shorter fringe period (i.e. more peaks) and lead to different SRI sensitivity.
Once the MZI sensor becomes in the leaky configuration, the operation of the interferometer changes. According to O. Duhem et al, the surrounding refractive index higher than the cladding allows a part of energy to be reflected at the interface of external medium and cladding and the other part of energy to be lost by Fresnel refraction. Fresnel reflection becomes the main source for the leaky cladding modes and transforms the guided cladding modes to leaky mode. O. Duhem et al. accounted for Fresnel losses by considering that only $\gamma$ of the coupled power is present at the end of the MZI region [34]:

$$\gamma = \exp[-mTL]$$

(8)

where $m$ is the number of Fresnel reflections per unity of length, $T$ is the Fresnel transmission coefficient, and $L$ is the length of the MZI region.

In leaky mode configuration, the effective index of the leaky mode becomes a complex number where real parts represent modal phase information in propagation and imaginary parts represent modal attenuation. When surrounding refractive index changes in leaky mode, the real part of the effective index for the leaky mode almost keeps constant, and the imaginary part of the effective index changes [32]. Therefore, in the leaky mode configuration, or at regions of refractive index greater than cladding, the transmission intensity of the MZI sensors should be investigated. The leaky mode interference in the cladding mode of MZI can be approximated using the quasi-leaky mode mentioned by Yang et. Al [31]. Quasi-leaky mode is an equivalent closed numerical model that can be used to represent radiation modes in open leaky waveguides [31]. Assuming that the single mode fiber and the leaky mode region are perfectly aligned, the modes are sufficient in the field expansion in the following investigation. At $z = 0$, which is the first
interface of single mode fiber and the leaky mode region, the electrical field can be expanded in terms of equivalent leaky modes in the leaky mode region:

\[ E_{in} = \sum_{m} \eta_m E_m \]  

(9)

where \( E_{in} \) is the electrical field of the fundamental mode from input single mode fiber, \( E_m \) is the \( m^{th} \) leaky mode in leaky mode region, and \( \eta_m \) is the excitation coefficient of \( m^{th} \) leaky mode which can also be used as:

\[ \eta_m = \frac{\int E_{in} \cdot E_m ds}{\int E_m \cdot E_m ds} \]  

(10)

As the electrical field in leaky mode region is caused by the modal interference of all the excited leaky modes with different propagation constants, at the certain distance \( z \), it is:

\[ E(z) = \sum_m \eta_m E_m e^{i\beta_m z} e^{-\alpha_m z} \]  

(11)

where \( \beta \) and \( \alpha \) are phase constant and attenuation constant of \( m^{th} \) excited leaky mode in leaky mode region. At \( z = \) Length of leaky mode region (i.e. second interface between leaky mode region and single mode fiber, or at the second splice point), the coupling of leaky mode region and guided single mode fiber is established, and its normalized output intensity can be approximated by:

\[ E T_{out} 10 \log_{10} \left( \frac{|\int E(L) \cdot E_{out} ds|^2}{|\int E(L)|^2 ds \int |E_{out}|^2 ds} \right) (dB) \]  

(12)
where $E_{\text{out}}$ is the electrical field of the fundamental mode in the output single mode fiber, which should equal $E_{\text{in}}$ if the input and output have use same single mode fiber.

4.3. Effect of coating on PCF MZI

As mentioned earlier in the Chapter, in the PCF MZI, effective refractive index of the cladding mode in the PCF is sensitive to the surrounding refractive index in the evanescent field where this cladding mode extends to. Recently, it has already been studied that sensitivity of the PCF MZI refractive index sensors can be significantly enhanced by applying a thin layer of coating with a material that has greater refractive index than silica. The coating atop PCF region would shift the cladding mode’s light distribution to near the cladding-SRI interface, which would result in an increase of the energy density in this region [201, 202]. From the increase of energy density at the PCF cladding-SRI interface, the intensity of the evanescent wave also increases, and thus increasing the refractive index sensitivity. It is known that the coating thickness and refractive index of coating is critically related with the PCF MZI sensitivity. In case of thickness, the thickness of the coating in tens of nanometer, which is very difficult and expansive to achieve, affects the sensitivity.

In this final thesis, the coating of nanoparticles on the fiber optics is an interest. According to Urrutia et al. in their review paper in regards to nanoparticle coating with fiber optic sensors mentioned that usage of nanoparticle coatings on MZI sensors are not very common [203]. Therefore, the effect, if any, of silver nanoparticle coating on PCF MZI would be investigated in this Chapter.
4.4. Fabrication of MZI refractive index sensors

All the MZI refractive index sensors used for the experiments were fabricated using the splicing technique. The Fujikura (FSM 40PM) fusion splicer, shown in Figure 4-8, was used for all splicing operations. For the fabrication of 40 um microfiber MZI, the standard single mode fiber was spliced with a 40 um microfiber with an arc power of 10 bit exposed for 1000 milliseconds. For the fabrication of PCF MZI, the fusion splicer was used to splice standard single mode fiber to a PCF with an arc power of 10 bit exposed for 1500 millisecond. As illustrated in Figure 4-9, the splicing created the collapsed region, which allows the core mode to diffract to the cladding modes, in the PCF portion. The length of PCF for the PCF MZI was 16 mm. The PCF used for MZI fabrication is the endlessly single mode LMA PCF (NKT Photonics A/S) as shown in Figure 4-10.

**Figure 4-8.** Fujikura FSM 40PM fusion splicer used for all splicing operations.

**Figure 4-9.** A schematic of a single mode fiber spliced to the photonic crystal fiber using Fujikura FSM 40PM fusion splicer.
4.5. Experiment setup

Different MZI sensors are used to sense in high refractive index from 1.4823 to 1.5365, similar to the Chapter 3. The 40 um microfiber MZI, uncoated PCF MZI, and silver nanoparticle coated MZI are fabricated and are used for high refractive index sensing as shown in Figure 4-11. The broad band light source (AFC BBS-1550) is connected to a FC/APC single mode fiber connector which is fusion spliced onto a MZI sensor. The other end of the MZI sensor is fusion spliced to another FC/APC single mode fiber connector which is connected to the optical spectrum analyzer (PHOTONETICS Walics). Each MZI sensor has two fusion splices on the joints of PCF or 40 um micro fiber with single mode fibers. Once all configuration has been setup, the transmission spectrum at air is marked as reference point. Then, the high refractive index solution is introduced to the regions shown in the schematic. Once the change in the spectrum has been saved, the refractive index solution is washed away, and the MZI is cleaned and rinsed with acetone until the reference spectrum is achieved in air.
4.6. 40 um MZI at high index

Similar to the LPG refractive index configuration, the transmission spectrum of 40 um microfiber MZI has been observed using an optical spectrum analyzer (OSA) and a light source. When MZI was connected to the optical spectrum analyzer, multiple peaks could be seen in air, with transmission peaks having a value of about 5 dBm, as shown in Figure 4-12. To test how the 40 um microfiber MZI behaves in refractive index below the cladding (i.e. guided mode), the sensor was put in tap water, which has refractive index of about 1.3. In water, the transmission peaks shifted to the left to the lower wavelength. The intensity of the transmission dip decreased by approximately 0.5 dBm. A clear shift to the left verified the fact the 40 um MZI functions properly as a refractive index sensor in index regions below the cladding. However, when the sensor was inserted to a solution with high refractive index (i.e. leaky mode), the intensity of the peaks substantially decreased as shown in Figure 4-13. The intensity of the transmission dip has decreased to about 1 dBm, and no visible wavelength shifts have occurred as shown in Figure 4-14. Moreover, no visible trend in transmission intensity is seen as well. This experiment has
verified that 40 um MZI sensors cannot be used as a refractive index sensor in the leaky configuration.

**Figure 4-12.** Transmission spectrum of 40 um microfiber MZI in response to air and water.

**Figure 4-13.** The transmission spectrum of 40 um microfiber MZI. Note that the transmission dip is very small.
4.7. PCF MZI at high surrounding refractive index

The transmission spectrum of PCF MZI in response to a variation in surrounding refractive index greater than cladding has been observed through OSA and a light source. First, the transmission wavelength shifts has been observed in the tap water, which has refractive index about 1.3. Unlike the 40 um microfiber MZI sensor discussed in 4.4., the transmission peak shifted to the right, towards the higher wavelengths. Then, the PCF MZI is put in the high refractive solutions to observe its performance in the leaky configuration. In comparison to the 40 um microfiber MZI, the PCF MZI sensor did have a transmission wavelength shift trend in response to high refractive indices as shown in Figure 4-15 and Figure 4-16. Although the achieved the transmission intensity in leaky mode was low and the peaks were broad, the peak wavelengths have shifted to the higher wavelengths towards the right. As seen in Figure 4-16, clearly, as the surrounding
refractive index increases, the wavelength of the peak shifted to the right. From surrounding refractive index range of 1.4823 to 1.5365, about 1.6 nm wavelengths shifted to the right, as illustrated in Figure 4-17. The low sensitivity of the wavelength shifts should be further investigated, but as mentioned in the literature, real part of the effective index of the cladding mode, which determines the modal phase information, almost keeps constant with varying surrounding refractive index. In terms of transmission intensity of the PCF MZI, clearly, noticeable decreasing trend in transmission intensity has been shown with increasing refractive index. Although very small, the PCF MZI sensors could be used to sense high refractive index by observing both changes in intensity and wavelength shift.

![Figure 4-15](image.png)

**Figure 4-15.** A transmission spectrum for uncoated PCF MZI in varying high refractive index.
Figure 4-16. A transmission spectrum for uncoated PCF MZI zoomed in at wavelength 1530 nm to 1550 nm. Clearly, the wavelength shifts to the right while the transmission intensity decreases increasing surrounding refractive index.

Figure 4-17. About 1.6 nm wavelength increased by increasing refractive index from 1.4823 to 1.5365.
4.8. Effect of PCF length in leaky configuration

In previous section, although very small, the wavelength has shifted to the right in 16 mm PCF MZI sensor. The literature mentions that the real part of the effective refractive index of the cladding mode almost stays constant, which means that the change in the modal phase information is very small. The transmission intensity of the PCF MZI sensor also decreased with increasing surrounding refractive index. To investigate further in the wavelength shift and transmission intensity, 3 different lengths of PCF region, 11 mm, 20 mm, and 25 mm has been investigated. The fringe space in MZI is known to depend on the length of the interferometer section. The longer interferometer, the more peaks are observed in the PCF MZIs as shown in Figure 4-18. In leaky mode configuration, the cladding mode loses most of its energy due to the Fresnel refraction, and only little amount of cladding modes from Fresnel reflection is obtained at the end. According to Duhem et al., the Fresnel losses depend with the length of MZI section as well [34], and therefore, transmission spectrum of the longer PCF MZIs (20 mm and 25 mm) has much more flat spectrum (smaller peaks). The 11 mm PCF MZI in leaky configuration had more noticeable peaks as shown in Figure 4-18. For all three lengths of PCF MZIs, the wavelength shifts to the right very slightly, but no noticeable trends were observed, especially because the peaks are broad as well. However, it can be seen that they peaks shift to the right. In terms of transmission intensity, the intensity decreases with increasing refractive index for all three PCF MZIs (i.e. amplitude of the peaks get greater). The smaller the PCF region are, the change in transmission with respect to the varying surrounding refractive index increased (i.e. better sensitivity). The transmission intensity change with varying SRI for PCF MZI behaved following the quasi-leaky mode.
approximation mentioned by Yang et al. [31]. At the maximum of the peak, higher RI had highest transmission intensity and lower RI resulted in lowest transmission while at minimum of the peak, visa-versa. Comparing the simulated quasi-leaky mode of SMF-core only fiber-SMF RI sensor with PCF MZI sensor as shown in Figure 4-19, the PCF MZI sensor behaved similar to the quasi-leaky mode simulation. If the PCF MZI sensors are used for high RI sensing, it should be used as intensity based sensors operating in quasi-leaky mode.
Figure 4-18. Transmission spectrum of (a) 11mm PCF MZI (b) 20 mm PCF MZI and (c) 25 mm PCF MZI

Figure 4-19. The comparison between a single mode fiber-core only fiber-single mode fiber sensor with a PCF MZI sensor.

4.9. Coated PCF MZI at high surrounding refractive index
The effect of thin layers of silver nanoparticle deposited at outer surface of PCF MZI in high refractive index has also been studied. Since the coating has affected the performance of the LPG sensor, similar phenomena are expected on the PCF MZI sensors. Therefore, the 20 pass of silver nanoparticles were deposited into PCF region of the sensors, for all 11 mm, 20 mm, 25 mm, respectively, and the refractive index experiment was conducted as did in section 3.5. The comparison of the uncoated PCF MZI and coated PCF MZI can be seen from Figure 4-20. Unlike the LPG sensor, the coating did not critically affected in the performance of the PCF MZI sensors. The wavelength shift of the PCF MZI sensors in leaky configuration is still almost negligible. However, for all lengths of the PCF MZI sensors, the transmission intensity did increased very slightly. For instance, in Figure 4-21, the sensitivity of the transmission intensity with respect to increasing surrounding refractive index increased very slightly with the silver nanoparticle coating. The silver nanoparticle coating in PCF MZI sensors are not as effective in leaky mode configuration refractive index sensing as the LPG sensors.
Figure 4-20. The transmission spectrum in leaky mode configuration of (a) uncoated 11 mm PCF MZI (b) coated 11 mm PCF MZI (c) uncoated 20 mm PCF MZI (d) coated 20 mm PCF MZI (e) uncoated 25 mm PCF MZI (f) coated 25 mm PCF MZI
Figure 4-21. Change of transmission intensity in (a) uncoated and (b) coated 11 mm PCF MZI
4.10. Conclusion

The response of two different configurations of MZI sensors, 40 um microfiber MZI and PCF MZI, has been experimented with varying refractive index in the leaky configuration. The 40 um microfiber MZI sensor failed to sense high refractive index solutions and PCF MZI sensors showed decreasing trend in transmission intensity with increasing high refractive index values in leaky mode configurations. The sensitivity of the wavelength shift in PCF MZI were almost negligible and thus could not be used. Moreover, it was found that as the length of PCF region increased, the peaks became smaller from higher Fresnel losses. Then, the effect of thin layer silver nanoparticle coatings on the PCF MZI has been investigated. The deposition of silver nanoparticles atop PCF MZI sensors did not have significant effects. The transmission intensity decreased by very little and therefore the sensitivity based on transmission intensity increased by little. If sensitivity optimizations could be done by adjusting coating thickness and the PCF length, PCF MZI sensors based on transmission intensity could potentially be used to measure the solubility and diffusivity of gas/SCF in polymers.
Chapter 5. Transmission intensity based gap sensor for measuring high surrounding refractive index solution

The transmission intensity based fiber optic sensors are frequently used for measuring the refractive index, especially in the region of high refractive indices. Commonly as discussed in Chapter 2, intensity based fiber optic sensors have single mode-multimode-single mode configurations or similar multimode configurations; but a new type of intensity based sensor in transmission mode is introduced, based on a gap in between to single mode fibers. In this Chapter, a simple and cost effective intensity based fiber optic sensor is introduced and its potential to be used as a measurement technique for solubility and diffusivity of gas/SCF in polymers is discussed. The change in transmission intensity with increasing refractive index from 1 to 1.6 is observed. Moreover the parametric study on the gap distance of the sensor to the transmission intensity has been conducted from the 0 um to 2000 um.

5.1. Introduction

Transmission intensity based fiber optic sensors has been used to measure the refractive index. C. Chen et al. has fabricated a novel class of multi-D- shaped optical fiber refractive index sensor where they constructed several D-Sections in multimode optical fiber using the femtosecond laser pulses[204]. As illustrated in Figure 5-1, regions of the multimode fiber has been machined into a D-shape so the core is exposed to the surrounding environment. The operating principle of the sensing bases on the attenuated total internal reflection along the fiber and the changes of attenuated light intensity of the
multi D-shaped fiber with respect to surrounding refractive index. Similarly, Y. Wang et al has fabricated a refractive index sensor based on a microhole in single-mode fiber created by the use of femtosecond laser micromachining, which is also based on the transmission intensity [205]. They discovered that if the hole diameter is close to the diameter of the core, no light would be directly guided in the fiber core without passing through the microhole. Therefore, the transmission behavior is critically dependent to the refractive index of the hole (i.e. RI of the environment solution). At certain refractive index, the refracted light would be guided to the core. However, at certain refractive index, only the second refracted light, or even the third refracted light is guided in to the core as shown in Figure 5-2. Since first refracted guided light would have less transmission loss than the second and third refracted guided light, the transmission intensity losses would be higher at certain refractive indices.

**Figure 5-1.** A schematic of D-shaped optical fiber refractive index sensor. Refractive index solution would make direct contact with the core [204]
Figure 5-2. Light guiding of the femtosecond laser machined microhole refractive index sensor [205]

Inspired by the transmission intensity based fiber optic sensors fabricated by both C. Chen and Y. Wang, a new transmission intensity based fiber optic sensor has been fabricated. Instead of partially introducing the core with the surrounding environment, introducing a gap in between the transmission configuration has been investigated. At one end of the gap (or at the first core to surrounding environment environment), the light would be coupled to the surrounding refractive index. At the second surrounding environment to core interface, some of the refracted light would be recoupled to the core mode. As discussed by Y. Wang, depending on the refractive index values of the surrounding environment, either first, second, third, and so on, refracted light would be coupled back to the core mode, and this would have an effect in the transmission loss. Therefore, the change in transmission intensity with respect to varying surrounding refractive index solution would be investigated in this Chapter.
5.2. Fabrication of the gap sensor

Since typical LPG and MZI based refractive index sensors have low sensitivity when the surrounding refractive index is greater than the refractive index of the cladding, referred as leaky mode, a new type of sensor has been fabricated to sense changes in high refractive index. As shown in Figure 5-3, the proposed gap sensor have very simple setup and operates with two single mode optical fibers aligned on the fixed stage. The sensor operates in transmission mode where one fiber tip receives the signal from the light source and transmits to another fiber tip which sends the signal to the optical spectrum analyzer. In between two optical fibers, a small gap exists, which creates a direct contact point between the core and the surrounding material. Therefore, the light propagating the core would couple to the surrounding environment, and only some would recouple back at the other end. As the gap sensor does not depend on the total internal reflections, the gap sensor should operate even in the leaky configuration. Theoretically, changes in surrounding refractive index should change the amount of light coupled and recoupled at the gap, and thus result in different transmission spectrum. The transmission losses should vary depending on the surrounding refractive index because at certain refractive index, first, second, and/or third refracted light would be guided back to the core mode. (first refracted light would have the least amount of transmission losses)
5.3. Gap sensor at high surrounding refractive index

The change in light intensity with varying surrounding refractive index in the gap sensor have been investigated. In the gap sensor, the alignment in between two optical
fibers are critical because the light propagates in the core, which is only about 8 micrometers. Two 8 micron cores must be aligned perfectly in all 3 axis to receive a full transmission spectrum from one core to another. Even if the cores are misaligned very slightly, the transmission intensity would be too low to observe any change in the surrounding refractive index. For instance, the primary gap sensor experiment has been conducted by placing two single mode fiber in a 125 um slot, aligning until the maximum transmission spectrum is achieved. Then, in that alignment configuration, different refractive index solutions, from 1.33 to 1.5365 has been tested. As illustrated in Figure 5-4, the transmission spectrum increases as the refractive index solution reaches 1.4823. Once refractive index solution increases further, the transmission spectrum starts to decrease again. However, obtaining the alignment of gap sensor in 125 um slot was very difficult and required a long time. For the application of the solubility and diffusivity measurement, more accurate and reliable method to measure refractive index is required and therefore the pigtail fixture method has been implemented.
In order to obtain perfect alignment, a new experiment setup with the two single mode fiber tips connected using the FC/APC couplers and joined with a ferrule connector are developed. Since the ferrule connector had threads, exact gap distances could be kept constant during the experiment. Hence, the distance between the gap and alignment is kept consistent, leaving only the surrounding refractive index as a parameter of this experiment. Moreover, refractive index solutions that goes up to refractive index of 1.61 has been prepared to see more results than the primary experiments. The result of the experiment is as shown in Figure 5-5. From refractive index of 1 to 1.4823, the transmission intensity increased. However, from refractive index of 1.4823 to 1.61, the transmission intensity decreased, which is a similar trend with the primary experiments. The maximum transmission intensity is obtained when the refractive index was closest to the refractive index of cladding, which is approximately 1.45. It can be estimated that at 1.45, the maximum transmission intensity could have been obtained. Instead of single
mode fibers, which has about 8 μm diameter core, 62.5 μm core diameter multimode fiber has been used to observe the transmission spectrum with varying refractive index. Ideally, with larger core, there would be larger amount of light propagating through the gap and more light receiving. However, a quick experiments using the multimode fiber has shown that the transmission spectrum actually has become worst and, no noticeable trends were obtained. Moreover, the gap sensor alignment was not easy even with the multimode fibers, and this may have been caused by larger loss of the multimode fibers.

![Normalized Transmission Spectrum](NormalisedTransmission.png)

**Figure 5.5.** Normalized transmission spectrum of the gap sensor with increasing refractive index. Note that the intensity increases up to about 1.45 and starts to decrease then. The general trend of the sensor is similar to the primary experiment results.

### 5.4. Gap distance parameterization

To investigate the effect of the gap distance on the transmission intensity, a parameterization experiment has been conducted. In air, which has refractive index
approximately 1, the transmission spectrum of two single mode fiber tips were observed with incrementing the gap distance. The schematic of the experiment is as shown in Figure 5-6. In this experiment, the only the varying parameter is setup to be the gap distance. The two single mode fibers are placed on to the V-groove using the HFV001 (Thorlabs) stage and the fibers are fixed in position by a strong magnet to ensure the consistent alignment. One of the fiber is attached on to the micro stage to pull away the fiber accurately to increase the gap distance by 20 micrometers. As illustrated in Figure 5-7, the result has been observed as expected. The strongest transmission intensity is obtained when the gap distance is at zero, and as soon as the gap is introduced, the transmission intensity decreased in a linear trend. Although the smaller gap of, about 40 um would be ideal for the refractive index sensing, washing away the residual refractive index solutions and placing a new solution becomes harder with smaller gap because of the capillary effect. Therefore, gap distance of about 100 – 200 um would be more ideal for refractive index sensing.
5.5. Polymer testing

The feasibility of transmission intensity based gap sensor with polymer has been investigated. Firstly, the temperature stability of the gap sensor has been studied. The
single mode pigtails are fitted into a fiber sleeve and is put in a temperature controlled furnace. The transmission intensity of the gap sensor at every 10 °C is measured through the OSA and the result is as seen in Figure 5-8. There are no noticeable changes in the transmission intensity observed with varying temperature. Therefore, the proposed gap sensor itself is insensitive to environment temperature.

**Figure 5-8.** The change in transmission intensity with response to temperature.

The change in transmission intensity with varying temperature with a polymer film filling the gap has been investigated to test the feasibility of polymers with the gap sensors. The polycaprolactone (PCL) 45k Mn has been fabricated as a film with 71 μm and are inserted in between the fibers. 45k Mn PCL is white when it is in a solid form but as it becomes in a polymer melt, its transparency changes as shown in Figure 5-9. The
change in transmission intensity as the polymer changes phases have been studied as illustrated Figure 5-10. The melting point of PCL is 60° C and as shown in Figure 5-10, as temperature increases, the transmission intensity increases gradually. When the temperature approaches the melting temperature, significant changes in transmission occur since the transparency changes with phase change.

Figure 5-9. Difference in transparency between solid and molten PCL.
Figure 5-10. The transmission intensity of PCL film with varying temperature

Then, the change in transmission spectrum with three different polymer film has been observed. Polymer films of fluorinated ethylene propylene, polypropylene, and PET with refractive indices of 1.34, 1.49, and 1.64 respectively, have been placed at the gap to measure the transmission as tabulated in Table 5-1. The thickness of the films were all 51 μm. The change in transmission intensity for the three polymer films have been obtained as shown in Figure 5-11. The trend of the change in transmission with polymer films matched with the trend from the RI solution test, where from FEP to PP (i.e. RI from 1.34 to 1.49) the transmission spectrum increased and from PP to PET (i.e. RI from 1.49 to 1.64), the transmission spectrum decreased. Although the spectrums obtained with
polymer films had higher intensities over all, this may be from the fact that the gap distances were shorted with the films.

**Table 5-1. Properties of the polymer films**

<table>
<thead>
<tr>
<th>Polymers</th>
<th>Refractive Index (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FEP (Fluorinated ethylene propylene)</td>
<td>1.34</td>
</tr>
<tr>
<td>PP (Polypropylene)</td>
<td>1.49</td>
</tr>
<tr>
<td>PET (Polyester)</td>
<td>1.64</td>
</tr>
</tbody>
</table>

**Figure 5-11.** The transmission spectrum of polymer films compared with the RI solutions.
5.6. Carbon dioxide desorption in a polycarbonate film

A desorption process where the transport of carbon dioxide (CO\(_2\)) out of a CO\(_2\) saturated polymer has been observed using the transmission intensity gap sensor. The PET film has been fabricated as a film and is put in a customized pressure chamber as shown in Figure 5-13. A PET film with dimension of 1 mm by 1mm by 51 \(\mu\text{m}\) has been machined using a femtosecond laser as shown in Figure 5-12. The machined PET film has been placed in the sample chamber and have been pressurized with pressure of 1250 psi of CO\(_2\) at room temperature for 48 hours. Next, the sample is taken out and the thickness has been measured before it is placed at the gap in a fiber sleeve between the pigtails. The entire setup process has taken 3 minutes as the pressure is released. Once the PC film is firmly placed at the gap, the transmission intensity is measured every 3 minutes. The change in transmission intensity over time has been plotted in Figure 5-14. The thickness of the PET film before and after 90 minutes are measured to be 51 \(\mu\text{m}\). The transmission intensity decreased gradually until about 80 minutes until it reached plateau. The increase in refractive index would cause decrease in transmission intensity according to the experiment conducted in high refractive index. It is assumed that mixture of CO\(_2\) and polymer would have slightly lower RI than that of the pure polymer. Moreover, the desorption plot obtained by the gap sensor had opposite trend from the CO\(_2\) sorption plot obtained with weight change method in literature, as shown in Figure 5-15, which is an interesting result considering that sorption is gas in taking of polymer and desorption is gas leaving the polymer. Further experiments should be conducted to validate this hypothesis.
**Figure 5-12.** 1 mm by 1 mm by 189 μm thick PET film machined using femtosecond laser.

**Figure 5-13.** A customized pressure chamber used to pressurize PET film. (A) a CO₂ tank (B) a high pressure regulator (C) a valve (D) a sample chamber (E) an exit valve.
Figure 5-14. The change in transmission intensity with time as the CO$_2$ desorbs out of PET film.

Figure 5-15. The CO$_2$ sorption film in PET using weight change method in literature.
5.7. Conclusion

The new intensity based fiber optic refractive index sensor has been introduced and the change in transmission intensity with increasing surrounding refractive index has been observed. The gap sensor consisted of very simple setup and operated with two single mode optical fibers aligned on the fixed stage. The core diameter of the single mode fiber was about 8 micrometers, and therefore, the aligning process is found to be difficult. The refractive index experiment with single mode fiber gap sensor showed an increasing trend until 1.428, then which started to decrease. The usage of multimode fibers have led to higher losses and resulted in no noticeable trend in the refractive index experiment. Since the single mode fiber gap sensor operated even in the leaky configuration, the gap sensor could be used to measure the solubility and diffusivity of gas/SCF in polymers, if the aligning process can be configured with polymer resins. Therefore, the transmission intensity based gap sensor has been used to experiment behaviors with polymer films. First, the temperature independence of the gap sensor has been experimented. Then, the change of transmission intensity as the PCL film changes its phase has been observed. As PCL changed its phase from solid to polymer melt, its transparency become clearer and the transmission intensity has increased. The CO\textsubscript{2} desorption test on PET film has conducted as the increase in transmission intensity with time has been observed. Further experiments should be conducted to verify that transparency change with CO\textsubscript{2} injection caused this result.
Chapter 6. Conclusion and future work

The overall conclusion of the final thesis along with the future works are presented in this Chapter.

6.1. Conclusion

The solubility is a proportional ratio of a solute in a solvent, and diffusion can be described as the process where a material is transported by the thermal motion of the molecules in a fluid or a matrix. In many of polymer process applications, solubility and diffusivity act as crucial parameters. Therefore, numerous measurement methods for obtaining solubility and diffusivity of gas/SCF in polymers have been investigated. This final thesis looks into detail of different methods for obtaining solubility and diffusivity. Most of the solubility and diffusivity measurement devices discussed are very expansive, complex to use, and require long experiment time. Hence, the potential availability of fiber optic sensors as solubility and diffusivity of gas/SCF in polymers have been discussed in this final thesis. Fiber optic sensors are widely used in many applications, especially as refractometers because it is accurate, reliable, cost efficient, and simple. The in fiber grating sensors, such as FBG and LPG, and the in line interferometer sensors, mainly Mach-Zehnder interferometer, are introduced and have been researched to verify that the fiber optic sensors could be used with polymers. Most of the refractive index sensors based on fiber optics only operate in refractive index ranges of 1 to 1.44 because once the surrounding refractive index becomes greater than that of cladding (i.e. 1.45), the total internal reflection is lost and only small portion of the light propagation occurs
(i.e. leaky configuration). Since most of the polymers used for processing have refractive index above 1.4, the fiber optic sensor must be overcome sensing limitations in the leaky configuration. This final thesis suggests different fiber optic sensors, a silver nanoparticle coated LPG sensor, a PCF MZI sensor, a silver nanoparticle coated PCF MZI sensor, and a transmission intensity gap sensor, that could be used to sense refractive index in leaky configuration.

The femtosecond laser written LPG sensor coated with silver nanoparticles is tested for sensing refractive indices higher than that of cladding. For coating silver nanoparticles, the cost efficient atomization based system is implemented to deposit thin layers to the LPG sensor. With 20 passes of silver nanoparticle coating on the LPG sensor, the transmission spectrum shifted towards shorter wavelength with increasing wavelength. From the refractive index of 1.4823 to 1.5365, approximately 1.2 nm wavelength shifted to the left.

The response of two different configurations of MZI sensors, 40 um microfiber MZI and PCF MZI, has been experimented with varying refractive index in the leaky configuration. The 40 um microfiber MZI sensor failed to sense high refractive index solutions. PCF MZI sensors showed decreasing trend in transmission intensity with increasing high refractive index values in leaky mode configurations. The sensitivity of the wavelength shift in PCF MZI were almost negligible and thus could not be used. Moreover, it was found that as the length of PCF region increased, the peaks became smaller from higher Fresnel losses. Then, the effect of thin layer silver nanoparticle coatings on the PCF MZI has been investigated. The deposition of silver nanoparticles atop PCF MZI sensors did not have significant effects. The transmission intensity
decreased by very little and therefore the sensitivity based on transmission intensity increased by little.

The new transmission intensity based fiber optic refractive index sensor has been introduced. The gap sensor consisted of very cheap and simple setup, which operated with two single mode optical fibers aligned on the fixed stage. The core diameter of the single mode fiber was about 8 micrometers, and therefore, the aligning process is found to be difficult. The change in transmission spectrum is observed even in the leaky configuration using the gap sensor, and it has also been found that the distance of the gap inversely affected the transmission spectrum. The polymer testing has been experimented by observing transmission intensity changes of PCL, PET, FEP, and PP films. The change in refractive index with polymer film resulted the similar trend with the RI solution test. Moreover, a desorption test of CO$_2$ in PET have opened up more possibility of using fiber optic sensors for acquiring solubility and diffusivity of gas/SCFs in polymers.

The proposed sensors are capable of sensing surrounding refractive index greater than the cladding, even in regions from 1.4 to 1.6, where the most of polymer’s refractive index is. Therefore, the fiber optic sensors potentially could be used to measure the solubility and diffusivity of gas/SCF in polymers if the following future works could be implemented.

**6.2. Future work**

In order to use fiber optic sensors as solubility and diffusivity measurement devices, several more researches in the fiber optic sensors other than sensing in leaky
configuration must be conducted. First of all, the sensitivity of all, in fiber gratings sensors, in line interferometer sensors, and transmission intensity based sensors must be enhanced. The sensitivity of the proposed silver nanoparticle coated LPG sensor could be improved by optimizing the number of passes coated onto the LPG sensor. Similarly, the parametric study of coating passes and length of PCF segment in PCF MZI sensors would contribute in improving the sensitivity of the PCF MZI sensors. Furthermore, it has been known that having tapers in fiber optic sensor enhances the sensitivity. Therefore, adding in fiber tapers along with the proposed sensors could cause an increase in sensitivity. In case of the transmission intensity based gap sensor, varying the fiber tips would have an effect in sensitivity. For example, having FC/PC pigtail instead of FC/APC would allow more sensitivity because FC/APC has slight angle at the fiber tip to reduce Fresnel reflection. Second, the fiber optic sensors must withstand harsh environment. Since the measurement of solubility and diffusivity of gas/SCF in polymers require high temperature and high pressure, only the robust sensors could be utilized for this application. The fixtures that houses the sensor should prevent any gas/SCF leakages while maintaining constant high temperature. Moreover, polymer resins are non-Newtonian fluid, meaning that viscosity of polymer resin should be taken into account. This may be more difficult as the fibers are small in size. However, by searching literatures on harsh environment fiber optic sensors, solutions should be available.

Investigations on sapphire fibers would allow more flexibility in using fiber optics as measurement device for measuring solubility because sapphire fibers offer unique characteristics. As sapphire fibers have refractive index of about 1.73, measuring refractive indices of most polymers with sapphire fiber sensors would not lead to leaky
configuration (i.e. would not lose total internal reflection if LPG sensor could be fabricated using the sapphire fiber). Moreover, the melting point of sapphire is over 2000 °C, and therefore, numerous studies have been investigated using sapphire fibers for sensing in harsh environment [206]. Hence, sapphire fibers would be a perfect fit if several limitations could be overcome. Being a hard crystal with high melting temperature, splicing becomes a problem in sapphire fibers [207]. In order to fabricate in fiber gratings sensors or in line interferometers, splicing is inevitable. In addition, sapphire fibers are coreless fibers, which means that no cladding exists. Therefore, cladding should be manually fabricated by modifying index with femtosecond laser, or depositing thin layers of coating. Creating fiber gratings would be challenging as well as not much studies have been done in literature on sapphire fibers due to the challenges and expensive cost. However, the advantages that sapphire fibers offer are very attractive, and have high potential. Thus, fabricating fiber optic sensors with sapphire fiber would be beneficial in solubility and diffusivity measurement applications.
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Appendix A – Theoretical approaches for solubility study

Many thermodynamic models are developed to study polymer melts and multi-component polymer blend systems. These theoretical thermodynamic models have been utilized by researchers to study the phase equilibrium behaviors between a gas/SCF and a polymer. Theoretical thermodynamic models can be useful tools to study many fundamental properties such as solubility and diffusivity of polymer/gas mixture systems. The basic theories and various equations of states (EOSs) are briefly discussed in this section.

An EOS is a mathematical relation among volume, pressure, temperature, and the composition of a closed system. In general, EOSs can be categorized into two types: empirical and theoretical models. The empirical EOSs are used more in the interpolation of obtained data, and thus result in poor predictions when extrapolated to high temperature or pressure conditions much beyond the ranges of experimental data. Furthermore, the empirical EOSs are difficult to use in systems with strong polar molecules or supercritical components. In contrast, the theoretical methods based on statistical thermodynamics offer many advantages over the empirical approaches. These models require less mixture data for the parameter fitting, allowing more reliability in the extrapolation at high temperatures and pressures, and simplifying the calculations of multi-component mixtures or systems with supercritical components. The following sections describe the theoretical approaches based on statistical thermodynamics.
A.1. Henry’s law

Henry’s law is not considered as an EOS model but it is a relation to estimate gas/SCF solubility in a polymer. According to Durill et al. Henry’s law is reliable only at low pressures [96, 208]. These researchers tested up to 30 MPa assuming that a Henry’s law constant is independent of pressure [209, 210]. However, at high pressures, a gas sorption isotherm has deviated significantly from the Henry’s law [96]. This deviation has been improved through the Langmuir type isotherm which accounts for gas solubility in interstitial spaces and micro-voids [211, 212]. Michaels et al. correlated the solubility of He, N$_2$, O$_2$, Ar, CH$_4$, CO$_2$, and ethane in polyethylene terephthalate (PET) and studied the deviations of gas sorption isotherms from the Henry’s law [213]. The Henry’s law represents reliable data in low pressures below 2 MPa and thus is commonly used in many solubility studies [107, 214]. The relation between a Henry’s law constant and a temperature can be described as: $H_1=H_0 \exp(-E_s/RT)$, where $H_0$ and $H_1$ are the Henry’s law constants, $E_s$ is the heat of solution, and $T$ is the temperature [210].

A.2. Flory-Huggins Theory (Cell Model)

The classical Flory-Huggins theory is based on a lattice model for describing polymer/solvent or polymer/polymer mixture systems. In a lattice model, all the cells are occupied with segments (i.e., solvent molecule or segment of a long polymer chain). The volume among different segments (segments of a long polymer chain or small solvent molecules) is assumed to be the same. Moreover, an assumption is made where a polymer segment and a solvent molecule could replace one another in the liquid lattice. The
energy of mixing for a mixture system is from both enthalpy of mixing and entropy of mixing.

The Flory-Huggins theory is a special model of the general cell model where all the cells are occupied by either the polymer chain segments or the small solvent molecules. This cell model demonstrates that the thermodynamic properties of a mixture strongly depend on the thermodynamic properties (volume, compressibility, etc.) of each pure component.

Schematic of a Flory-Huggins thermodynamic model [215]. Black circles and white circles represent polymer molecule chains and small solvent molecules, respectively.

Y. L. Cheng et al. first used the Flory-Huggins theory to study gas solubility in a polymer [216]. The Flory-Huggins theory is useful to investigate solubility and phase equilibrium within limited pressure and dissolved gas concentration ranges, but the concern is that this theory fails to predict isothermal data at high pressure and high dissolved gas concentration ranges as it ignores the volumetric changes of a polymer/gas mixture system by pressure, temperature, and mixing [217].

A.3. Sanchez-Lacombe Theory (Lattice Fluid Model)

To overcome the limitations of the Flory-Huggins theory, Sanchez and Lacombe proposed a dimensionless theoretical model based on a fluid-lattice theory (i.e., the SL EOS). Vacant cells are adopted in Sanchez-Lacombe theory, and these cells are introduced in the lattice to illustrate the extra entropy change in the system as a function
of volume and temperature of the polymer/gas mixture system. The changes in volume only occur when new holes or vacant sites occur on the lattice because the lattice size, or cell volume (i.e., occupied volume), is fixed. To obtain a pure component, SL EOS requires three EOS parameters such as a characteristic temperature $T^*$, a characteristic pressure $p^*$, and a characteristic specific volume $v_{sp}^*$, similar to the Flory-Huggins theory.

The SL EOS is a quantitative theory and it is applicable for both gaseous and liquid states. The chemical potential of a gas component can be calculated the same way as that of a liquid phase. Therefore, the SL EOS is not required to define a standard state for a gas (i.e., gas standard state fugacity) unlike the Flory theory. The SL EOS model is the most widely used EOS model, and C. Panyaiotou extended this SL EOS further by accounting for the strong interactions (i.e., hydrogen bonding) between the polymer and the solvent [218]. The SL EOS model is simple while being capable of extrapolating the available experimental data to high temperatures and pressures and thus is used widely to investigate gas solubility in polymers [37, 41, 108-111].

A.4. Simha-Somcynsky Theory (Hole Model)

To improve the Flory-Huggins theory for the liquid state (i.e., polymer melts), Simha and his co-workers proposed the SS theory based on the lattice-hole theory [219, 220]. The lattice-hole model considers a lattice of cells, each of which can accommodate either a small molecule or a chain segment. Also, similar to the lattice fluid model, the hole model improves the liquid state cell model by introducing the vacancies in the lattice, which can describe the thermal expansion [219]. In obtaining the characteristic
parameters of SS EOS, it uses the “square well” approximation to the cell potential and the nearest neighbor contributions to the lattice energy are not included. The resulting coupled equation of states must be solved simultaneously with an expression minimizing the partition function with respect to the fraction of occupied sites. Unlike SL EOS, in the SS theory, the change in a cell volume, or the occupied volume, is also allowed with temperature and pressure. According to Q. Gao, C.B Park, X. Xu, and J. Wang, SL EOS better fits for the data of gas while SS EOS better fits for data of polymers [221].

A.5. Statistical Association Fluid Theory (SAFT)

Statistical Association Fluid Theory or SAFT, is an EOS based on the lattice theory [222] where it assumes that real fluids exist in a continuous space rather than in a lattice [96]. The SAFT EOS has been proven to successfully forecast the thermodynamic properties of small molecules over a wide density range, and has also been applied effectively to polymeric fluids and their mixtures. Instead of using a hard sphere reference fluid as in many other engineering EOS, the SAFT is developed from an idea of a reference fluid incorporating both the molecular geometry and molecular association effects. In the SAFT model, the fluids (molecules) are described as equi-sized spherical segments interacting according to a square-well potential. Therefore, the component (molecules) is well described with the following three component parameters: \( m \), \( v^0 \), and \( u^0/k \), where \( m \) is the number of segments per molecule, \( v^0 \) is the segment molar volume in a close-packed arrangement in milliliters per mole of segments, and \( u^0/k \) is the segment energy or the well depth, a temperature-independent energy parameter, in kelvin.
The SAFT EOS can be used for various gas/SCF polymer systems at high pressures [100, 223]. S.H. Huang et al. extended the SAFT EOS theory to study the mixtures of real, molecular, and macromolecular fluids over a wide density range to investigate gas solubility in a polymer [223, 224]. This model is more complex than the SL EOS but it accounts for the specific interaction such as hydrogen bonding among the segments of polymer molecules [40].
Appendix B – Theoretical approaches for diffusivity study

Two kinds of diffusion, steady and unsteady diffusion, are discussed here. The diffusivity values of these two types of diffusion are always obtained by applying mathematic models with the corresponding experimental data. Common equations and models are described below.

B.1. Steady Diffusion

For steady diffusion, a flux does not change with time but with space. It is expressed as:

\[ J = -D \nabla C \]  \hspace{1cm} (13)

where \( J \) is the fluxes of atoms in atoms/(m\(^2\) \cdot s), \( D \) is a diffusion coefficient in m\(^2\)/s, and \( \nabla C \) is a concentration gradient in atoms/m\(^4\). It is called a Fick’s first law. Considering a unidirectional case, this equation can be simplified as:

\[ J_x = -D \frac{dC}{dx} \]  \hspace{1cm} (14)

This simplification is valid in a membrane diffusion case when the thickness of a membrane is much smaller than the other dimensions. The mean diffusion coefficient over the entire range of concentrations can be obtained by the “time lag” method developed by R. M. Barrer and E. K. Rideal in 1939 [225]. In a steady state, the mean diffusion coefficient is directly proportional to time:
\[ D = \frac{l^2}{6\theta} \]  

Where \( D \) is the diffusion coefficient, \( l \) is the length of a polymer sample which a gas travels through, and \( \theta \) is the time lag which is the intercept of the time axis with the extrapolated linear steady state portion of the pressure-time curve.

**B.2. Unsteady Diffusion**

This is the case where a gas flux changes with respect to time before a diffusion process reaches equilibrium. Many different models are used to describe this type of diffusion: Crank’s model, Dual mode model, Gas-Polymer Matrix Model, Free Volume Model, Mass Transfer Model, and some Empirical Models. Some of these models have very limited applications. For example, a Dual mode model and a Gas-Polymer Matrix Model are only used for glassy polymers. The detailed descriptions on each of these models are described below.

1) Crank’s model

Crank’s model developed by J. Crank and G. S. Park is most used, since this model is easy to understand, and suitable for various experimental geometries, accommodating different boundary conditions. A sample shape has a great effect on gas transport behaviors. In 1968, Crank and Park derived several equations based on the sample geometries. The following equation was used to describe unsteady gas diffusion through a polymer film[226]:

\[ D = \frac{l^2}{6\theta} \]
where $M_t$ and $M_\infty$ represent the mass uptakes of a penetrant at time $t$ and at a long time, respectively. $L$ is a diffusion length. This model is mostly used in pressure decay experiments. This equation can also be applied to diffusion cases along a cylindrical rod or tube with a length of $L$, where one end of its surface is sealed and the other end is maintained at a constant concentration [226].

Although the simplified model is widely used, the original model \( \frac{M_t}{M_\infty} = 2 \left( \frac{D t}{\pi L^2} \right)^{1/2} \) gives more comprehensive descriptions of solubility and diffusivity calculations [227].

2) Dual-mode Sorption Model

This model divides diffusing molecules into two different populations: (i) the molecules dissolved by an ordinary dissolution process whose concentration is $C_D$, and (ii) the molecules trapped in micro-voids which do not diffuse whose concentration is $C_H$.

The dissolved gas molecules obey The Henry’s law:

\[
C_D = k_D p
\]  

(17)

The concentration of the trapped gas molecules in micro-voids is given by the equation of Langmuir [228]:

\[
C_H = \frac{C_H^{bp}}{1 + bp}
\]  

(18)

where $C'_H$ is a Langmuir saturation constant, which is directly related to the global volume of a specific site (i.e., micro-voids), and $b$ is an affinity constant for these sites, $p$ is the pressure.
These two populations have their own constant diffusion coefficients \((D_D, D_H)\) [229] and \(J\) is the total flux.

\[
J = J_D + J_H = -D_D \frac{\partial C_D}{\partial x} - D_H \frac{\partial C_H}{\partial x}
\]  

\((19)\)

3) Gas-Polymer Matrix Model

Different from the dual mode model described in the previous section, this gas-polymer matrix model is based on the assumption that there is only one population of a penetrant but with an interaction between a solute and a polymer matrix. The basic concept involves an activation energy which represents the necessary energy for the separation of macromolecular chains by cooperative motions of sufficient amplitudes to allow a penetrant to execute its diffusional jump. This model assumes that the presence of a penetrant reduces the intermolecular forces between the polymer chains, resulting in a smaller necessary activation energy for the separation of chains. The increase in a penetrant concentration leads to a higher diffusion coefficient due to an enhanced segmental mobility of the chains. From a mathematical viewpoint, this theory proposes the following relation to express the interaction between a penetrant and a polymer [229]:

\[
D = D_0 (1 + \beta C) \exp(\beta C)
\]  

\((20)\)

where \(D_0\) is a diffusion coefficient in a zero-concentration limit, \(\beta\) is a constant relating the excess activation energy of chain separation to the depression of the glass-transition temperature of the polymer by a gas, and \(C\) is a gas concentration. As for the concentration, it can be expressed as:

\[
C = \sigma_0 p \exp(-\alpha C)
\]  

\((21)\)

where \(\sigma_0\) is a solubility coefficient at infinite dilution, \(p\) is pressure, and \(\alpha\) is a constant describing the effect of gas-polymer interactions on solubility changes. The
smaller \( \alpha \) is, the less effect a penetrant has on solubility [230]. This model gives a better understanding of molecular interactions, gas perturbation, and thermodynamics than the dual-mode sorption model.

4) Free Volume Model

The basic idea of this theory is that a gas molecule can move from an original position to another when a local free volume exceeds a certain free volume fraction limit. This model explains the relationship among a diffusion coefficient and parameters such as a glass transition temperature, a penetrant shape and size, a temperature, and a concentration.

A free volume fraction, \( f \), is one of the most important parameters for solubility and diffusivity definitions. It is defined as:

\[
f = \frac{V_f}{V_{Tot}} = \frac{V_{Tot} - V_{occ}}{V_{Tot}} \tag{22}\]

where \( V_{Tot} \) is a total volume, \( V_{occ} \) is an occupied molecules volume and \( V_f \) is a free volume.

A thermodynamic diffusion coefficient, \( D_T \), is given by H. Fujita and K. Kishimoto [231] as:

\[
D_T = RT A_d \exp \left( -\frac{B_d}{f} \right) = RT m_d \tag{23}\]

where \( R \) is a gas constant, \( T \) is a temperature, \( A_d \) is a parameter related to a penetrant size and shape, \( B_d \) is a characteristic parameter of an available free volume fraction, and \( m_d \) is a mobility of diffusing molecules relative to a polymer. The combined free volume fraction, \( f \), is defined as:
\[ f = \phi_1 f_1 + \phi_2 f_2 \]  

(24)

where \( \phi_i \) and \( f_i \) are corresponding volume fraction and the free volume fraction for a component \( i \). The index 1 refers to the diffusing molecule and the index 2 to the polymer. Defined by J. Comyn [232], \( D_T \) can also be expressed as:

\[ D_T = A \exp\left(\frac{\partial \ln C}{\partial \ln a}\right) \]  

(25)

where \( C \) is a penetrant concentration, and \( a \) is a penetrant activity parameter.

Based on the J. Comyn’s research, A. Peterlin modified the equation (11) and proposed the following equation [233]:

\[ D = D(0) \exp(\alpha_d C) \]  

(26)

where,

\[ \alpha_d = \frac{B_d f_1}{(\phi_2 f_2)^2} \]  

(27)

\[ D(0) = RT A_d \exp\left(\frac{-B_d}{\phi_2 f_2}\right) \]  

(28)

Here, index 1 refers to the diffusing molecule and index 2 to the polymer.

However, this model is considered inappropriate for diffusion of small molecules. Therefore, correction for a free volume fraction was applied to the model. The free volume fraction depends on three thermodynamic variables as shown below [233-235]:

\[ f(T, p, C) = f_{\text{ref}}(T_{\text{ref}}, p_{\text{ref}}, 0) + \alpha(T - T_{\text{ref}}) - \beta(p - p_{\text{ref}}) + \gamma C \]  

(29)

The first term describes a standard fractional free volume in a reference state; the second term shows the increase of a free volume with thermal dilation where \( \alpha \) is a thermal expansion coefficient of the free volume; the third term indicates free volume shrinking during a hydrostatic compression (\( \beta \) is compressibility); and the forth term weighs the influence of a gas concentration on a free volume.
5) Mass Transfer Model

This theory is based on the Fick’s second law. This model can determine a diffusion coefficient by measure a gas concentration in the polymer phase. The basic procedure of this model is to fit experimental gas concentration data with a predicted function.

In a transient state, a penetrant concentration could be expressed by functions of gas location in the sample and time. A diffusion coefficient is an optimization parameter in this function.

The fundamental equation for this model is:

$$\frac{\partial c}{\partial t} = D \frac{\partial^2 c}{\partial z^2}$$  \hspace{1cm} (30)

Where $c$ is a penetrant concentration, $t$ is time, and $z$ is a gas travel direction in the sample, and $D$ is a diffusion coefficient. This equation was modified by many researchers [120, 122, 236]. An initial condition is given as below, describing no gas concentration in a polymer sample at time $t = 0$ [226]:

$$c(z, 0) = 0 \quad 0 < z \leq L$$  \hspace{1cm} (31)

For gravimetric experiments with the samples with one sealed end, the boundary conditions for $z = 0$ are as below with no gas concentration change at the sealed end:

$$c(0, t) = c(t) \quad 0 \leq t \leq T$$  \hspace{1cm} (32)

$$\frac{\partial c}{\partial z} \bigg|_{z=L} = 0 \quad 0 \leq t \leq T \frac{\partial c}{\partial t} = D \frac{\partial^2 c}{\partial z^2}$$  \hspace{1cm} (33)

This model can also be approached with an iterative algorithm with an initial guess value for a diffusivity coefficient. The value for a diffusivity coefficient keeps changing until the difference between calculated and experimental data are within the tolerance limit.
6) Empirical Model

For gas/polymer systems in which gas solubility essentially obeys the Henry’s law (e.g., hydrocarbons in elastomers), the dependence of a diffusion coefficient on the absorbed penetrant concentration has been empirically represented at a given temperature by the equations shown below [237]:

Linear model [238]:

\[
D(C) = D_0(1 + \beta C)
\]  
(34)

Exponent model [234, 239]:

\[
D(C) = D_0 e^{\beta c} c(z, 0) = 0 \quad 0 < z \leq L
\]  
(35)

Concentration and pressure dependent model [240]:

\[
D(C, p) = D(0,0) e^{(\beta p + \alpha C)}
\]  
(36)

Where \(D\) is a diffusion coefficient, \(C\) is a penetrant concentration, \(p\) is a pressure, \(D_0\) is a diffusion coefficient when \(C\) is zero, \(\beta\) is a temperature dependent constant, and \(\alpha\) is a constant related to plasticization.

The equations (21) and (22) obey the Henry’s law. In equation (22), the diffusion coefficient shows a stronger dependence on the concentration. Equation (23) takes hydrostatic constraint and plasticization into account [237].