



Titre: Control and Optimization of Sealing Layer in Films
Title:

Auteur: Zahra Najarzadeh
Author:

Date: 2014

Type: Mémoire ou thèse / Dissertation or Thesis

Référence: Najarzadeh, Z. (2014). Control and Optimization of Sealing Layer in Films [Thèse de doctorat, École Polytechnique de Montréal]. PolyPublie.
Citation: <https://publications.polymtl.ca/1406/>

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URL de PolyPublie: <https://publications.polymtl.ca/1406/>
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Directeurs de recherche: Abdellah Ajji
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Programme: Génie chimique
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UNIVERSITÉ DE MONTRÉAL

CONTROL AND OPTIMIZATION OF SEALING LAYER IN FILMS

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THÈSE PRÉSENTÉE EN VUE DE L'OBTENTION
DU DIPLÔME DE PHILOSOPHIAE DOCTOR
(GÉNIE CHIMIQUE)

AVRIL 2014

UNIVERSITÉ DE MONTRÉAL

ÉCOLE POLYTECHNIQUE DE MONTRÉAL

Cette thèse intitulée:

CONTROL AND OPTIMIZATION OF SEALING LAYER IN FILMS

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en vue de l'obtention du diplôme de : Philosophiae Doctor

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DEDICATION

“To my beloved family”

ACKNOWLEDGEMENTS

First of all, I would like to express my deep and sincere gratitude to my supervisor, Prof. Abdellah Ajji for supporting and challenging me. His wide knowledge and hard work has always been a great motivation for me. I appreciate his patience, encouragements, advice, and guidance as well as all the productive discussions we had.

I am grateful to Prof. Seyyed Hesamoddin Tabatabaei for his support, advices, sharing his industrial and scientific experience, and his comments on the experimental work and papers.

This work would have not been possible without Prolamina and Saputo companies, who provided the financial support throughout the project. I appreciate sharing their up to date industrial experience and knowledge. Especial thanks to Prolamina facility in Terrebonne for allowing me to use some of their equipment during my Ph.D. study.

A special word of thanks is owed to our brilliant and helpful research associate in 3S Pack chair group Ms. Claire Cercl  . I would also like to thank her for translating a part of this thesis to French.

My gratitude to the staff and technicians of Ecole Polytechnique Montreal, chemical engineering department especially Ms. M  lina Hamdine, Ms. Weawkamol Leelapornpisit, and Mr. Guillaume Lessard.

One of the most valuable experiences that I had was the opportunity to work as a part of the great 3SPack research team. The cooperative atmosphere and sharing the results and ideas improved this work. I am especially grateful to my fellow student, Ramin Yousefzadeh, Richard Silverwood.

I would like to thank Jean-Baptiste Bruchet for conducting some experiments for the third paper of this work during summer and autumn of 2013.

My sincere thanks to my friends in   cole Polytechnique of Montreal who their help, discussions, and encouragements were very valuable.

I would also like to thank my family and my husband, Mohammad, who believed in me more than I believed in myself, whose constant support and encouragement gave me the strength to get through many difficulties.

RÉSUMÉ

Le scellage à chaud est un procédé qui permet de sceller hermétiquement un emballage en appliquant de la chaleur et de la pression sur deux films de polymère pendant un temps spécifique (temps de scellage). Le scellage à chaud est utilisé dans le domaine de l'emballage pour une vaste gamme d'applications, et en particulier dans le domaine de l'emballage alimentaire. Depuis les vingt dernières années, les équipements de scellage à chaud fonctionnent sur les lignes de production, à très hautes vitesses, ce qui requiert des conditions de procédé et d'optimisation des matériaux. Les paramètres du scellage à chaud, tels que la température, le temps de scellage, la pression ainsi que les caractéristiques du matériau du film tels que le poids moléculaire, la distribution en poids moléculaire et la présence de petites et de longues ramifications de chaînes sont des facteurs clés pour contrôler la qualité finale de la zone scellée.

Dans la première partie de cette étude, les relations existantes entre les paramètres du procédé de scellage à chaud et la résistance de l'interface ainsi que les mécanismes ont été étudiés. Au cours du procédé de scellage, les effets de la température, de la pression et du temps de scellage sur la force de scellage du LLDPE sont très importants. Ils ont été illustrés par une carte 3D 'temps-température' et 'pression-température', une méthode qui est facilement applicable à d'autres systèmes. Les mécanismes d'adhésion des polymères semi-cristallins étant mal connus, nous avons donc étudié le comportement en adhésion du LLDPE, un polymère semi-cristallin. Les résultats montrent que la dépendance 'temps-température' de la force d'adhésion est une conséquence de la disponibilité d'une certaine fraction de la phase amorphe du polymère à l'interface du film. La force d'adhésion suit une corrélation linéaire avec la racine carrée du temps de scellage ce qui est également confirmé par les modèles de soudure. La pente de cette corrélation linéaire augmente avec la température.

Par la suite, ce travail met en concurrence les deux principales approches d'analyse du comportement des chaînes du polymère : le point de vue microscopique basé sur la théorie de reptation, et le point de vue macroscopique basé sur une analyse des propriétés du polymère dans son volume. Les mécanismes de rupture et les deux surfaces pelées ont également été analysés après pelage. De plus, la topographie et la morphologie des surfaces pelées, qui ont été scellées auparavant avec différentes conditions de scellage ont été étudiées à l'aide des techniques d'AFM et SEM. Dans le cas d'une rupture des liens, la propagation la rupture génère de nombreuses

fibrilles très orientées dans le sens de la rupture sur la surface du pelage. Les fibrilles étaient visibles en AFM et au SEM sur les surfaces pelées qui avaient été scellées à basses températures. Pour les spécimens scellés à plus hautes températures, la morphologie fibrillaire se développe en fibrilles plus épaisses et plus larges. Ce comportement est une conséquence d'un meilleur mouillage et interdiffusion des chaînes à hautes températures.

Les paramètres de rugosité ont été comparés et agissent comme représentants du degré d'interdiffusion des molécules et d'aire de contact de la surface du film. Augmenter la température de scellage résulte en une surface plus rugueuse ce qui témoigne d'une meilleure diffusion moléculaire.

Dans la deuxième partie de ce travail, le rôle de l'architecture moléculaire du polymère sur l'auto adhésion interfaciale à des températures supérieures à la température de fusion (appelé hot tack) a été étudié. Le poids moléculaire (M_w), la distribution en poids moléculaire (MWD), le nombre et la distribution de longues ramifications (LCB) et de courtes ramifications (SCB) le long des chaînes de polyéthylène sont autant de paramètres moléculaires étudiés. Cette analyse révèle une corrélation linéaire positive entre la force d'adhésion et le poids moléculaire pour des polyéthylènes métallocènes. De plus, la même corrélation est observée pour la viscosité élongationnelle transitoire ce qui met l'emphasis sur le rôle de la force en fondue dans l'obtention d'une bonne force d'adhésion. Cependant l'augmentation de la présence de longues chaînes de ramification ainsi que l'augmentation de leur nombre réduisent la force d'adhésion. Les copolymères métallocènes catalysés éthylène α -oléfine ont généré une force d'adhésion supérieure à celle des polyéthylènes conventionnels et ceci est expliqué par une composition des chaînes plus homogènes pour les premiers. Enfin, il a été déterminé que la force interfaciale d'auto-adhésion était le premier facteur influençant l'interdiffusion des chaînes à travers la surface et, ensuite, la force en fondue du polymère.

Finalement, dans la dernière partie de ce travail, les mélanges de polyéthylènes avec des architectures de chaînes différentes ont été étudiés. La force d'adhésion et la dépendance à la température pour des mélanges binaires de polyéthylènes métallocènes catalysés éthylène α -oléfine et de polyéthylène conventionnels, de différentes compositions, ont été analysées. Il a été observé que le niveau d'amélioration de la force d'adhésion dépend fortement de l'architecture moléculaire et des longues ramifications de chaînes. Les polymères étudiés exhibent un plateau

aux hautes forces d'adhésion pour une certaine plage de température. Ce plateau est appelé 'fenêtre de température au plateau' (plateau temperature window). Notre étude a montré que la largeur de ce plateau est corrélée avec l'aire sous la courbe 'contrainte-élongation' obtenue grâce aux expériences de rhéologie. Il est donc suggéré qu'une plus grande ténacité du matériau en fondue résulte en un plateau de force d'adhésion plus large.

ABSTRACT

Heat sealing is the process to achieve the hermetic closure of a package by applying heat and pressure on two polymer films for a specified dwell time. The heat sealing process has been used in packaging of a wide spectrum of products, in particular for food packaging. During the past two decades, heat sealing machinery has been developed for very high speed line production which requires very precise process condition and material optimization. Heat seal process conditions, including temperature, dwell time, and pressure as well as film characteristics such as molecular weight, molecular weight distribution, long and short chain branching are the key factors that control the final quality of the joint.

In the first part of this study, we have investigated the interrelated influence of heat seal process parameters on the strengthening of the interface as well as the micro mechanisms involved. The effects of heat seal temperature, pressure, and dwell time on seal strength of LLDPE were illustrated in “time-temperature” and “pressure-temperature” 3D maps, a methodology that is easily applicable to other systems. The interfacial adhesion of semicrystalline polymers was not well understood. We thus studied the adhesion behaviour of LLDPE as a semicrystalline polymer. The results indicated that the time and temperature dependence of seal strength is a consequence of the fraction of amorphous phase available at the interface of films. The seal strength had a linear correlation with the square root of sealing time which was also supported by the welding models. The slope of this linear correlation increased with temperature.

This work was followed concurrently by two main approaches: analysis of polymer chain behaviour from microscopic point of view based on reptation theory and conventional macroscopic analysis based on bulk properties of the polymer. Failure mechanisms and fractured surfaces were also analyzed after peeling. Moreover, the topography and morphology of peeled surfaces, which were sealed at different heat seal conditions, were investigated by AFM and SEM. In the debonding failure mode, the crack propagation caused numerous highly oriented fractured craze fibrils on the peeled surfaces. The fractured fibrils were visible in AFM and SEM peeled surfaces of samples sealed at low temperatures. In specimens sealed at higher temperature the fibrillar morphology developed into thicker and larger fibrils. This behaviour was a result of better wetting and interdiffusion by increasing temperature.

We also utilized the comparative roughness parameters as representatives of the level of film surfaces' contact area and molecular interdiffusion. Increasing the seal temperature resulted in rougher surfaces which indicated the improved diffusion.

In the second part of this work, we investigated the role of molecular architecture of the seal polymer in interfacial self-adhesion at temperatures higher than melting point i.e. hot tack. The analyzed molecular structures include molecular weight (M_w), molecular weight distribution (MWD), amount and distribution of long chain branch (LCB) and short chain branch (SCB) distribution among and along polyethylene chains. Our analysis revealed a positive linear correlation between seal strength and molecular weight for linear metallocene polyethylene. Furthermore, the same relation was observed for elongational transient viscosity emphasizing the role of melt strength in adhesion strength. However, the presence of long chain branching and increasing the long chain branch content reduced the adhesion strength. We explained the superior adhesion strength of metallocene catalyzed ethylene α -olefin copolymers compared to conventional polyethylene resins based on the homogenous composition distribution of their chains. Finally, we argued that the interfacial self-adhesion strength was primarily the outcome of chain interdiffusion across the interface and then the melt strength of the polymer.

In the last part of this work, we focused on the blends of polyethylene resins with different chain architectures. The dependency of adhesion strength on temperature for binary blends of metallocene catalyzed ethylene α -olefins and conventional polyethylene resins were analyzed for a range of different compositions. We found that the level of improvement in adhesion strength depended on the molecular architecture and long chain branching. The polymer shows a plateau of its highest adhesion strength for a range of temperature, called plateau temperature window. Our findings indicated that the plateau broadness was correlated with the area under stress-strain curve of extensional rheological measurements. We thus suggested that the higher melt toughness can result in a broader plateau of adhesion strength.

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LIST OF SYMBOLS AND ABBREVIATIONS

AFM	Atomic force microscopy
BOPP	Biaxially oriented polypropylene
CCD	Comonomer composition distribution
De	Deborah number
DSC	Differential scanning calorimetry
D_{self}	Self diffusion coefficient
EVA	Ethylene vinyl acetate copolymer
f_a	Amorphous fraction
FDA	Food and Drug Administration
GPC	Gel permeation chromatography
H	Microscopic property at interface
h	Film thickness (μ)
HDPE	High density polyethylene
HFFS	Horizontal form fill seal
LCB	Long chain branch
LDPE	Low density polyethylene
LD	Low density polyethylene
LLDPE	Linear low density polyethylene
LL	Linear low density polyethylene
LVE	Linear viscoelastic behaviour
m-PE	Metallocene catalyzed polyethylene
M_c	Critical entanglement molecular weight
MFI	Melt flow index (g/10min)

M_w	Weight average molecular weight (kg/mol)
MWD	Molecular weight distribution
N	Number of points within the AFM image
NMR	Nuclear magnetic resonance
PBS	Polybutylene succinate
PCL	Polycaprolactone
PDI	Polydispersity index
PE	Polyethylene
PE-g-MA	Polyethylene grafted maleic anhydride
PET	Polyethylene terephthalate
PHA	Polyhydroxyalkanoate
PLA	Polylactic acid
PVA	Polyvinylalcohol
PVOH	Polyvinylalcohol
R_a	Arithmetic average of the absolute values of the surface height deviations (nm)
R_g	Radius of gyration (nm)
RMS	Root mean square average of height deviations (nm)
S	Interface strength
SBR	Styrene-butadiene rubber
SCB	Short chain branch
SER	Sentmanat extensional rheometer
SEM	Scanning electron microscopy
SSp	Plateau seal strength (g/25.4mm)
t	Time (s)

T_c	Crystallization temperature ($^{\circ}\text{C}$)
TESP	Tapping etched silicon probe
T_g	Glass transition temperature ($^{\circ}\text{C}$)
T_{mf}	Final melting point ($^{\circ}\text{C}$)
T_{pi}	Plateau initiation temperature ($^{\circ}\text{C}$)
T_{pf}	final plateau temperature ($^{\circ}\text{C}$)
T_{si}	Seal initiation temperature ($^{\circ}\text{C}$)
wt	Weight fraction
v	Peel rate (mm.s^{-1})
VFFS	Vertical form fill seal
X_c	Degree of crystallinity
Z	Height from mean plane of the AFM image (nm)
Zn	Ziegler-Natta catalyst

Greek letters

$\dot{\epsilon}$	Strain rate (s^{-1})
τ	Saturation time (s), reptation time (s)
η_e^+	Transient elongational viscosity (Pa.s)
η^*	Complex viscosity (Pa.s)
ΔH_S	Heat of fusion (J.g^{-1})
ΔH_U	Heat of fusion for a theoretically 100% crystalline polyethylene (J.g^{-1})
ΔH_T	cumulative heat of fusion (J.g^{-1})
μ	Micron
$\dot{\gamma}$	Strain rate (s^{-1})
ω	angular frequency (rad/s)

CHAPTER 1

INTRODUCTION

The flexible packaging industry grows every year to replace other kinds of containers in food industry because of cost benefits and environmental advantages such as landfill occupation space. The majority of pouches are made of multilayer polymer films and are sealed mostly by a heating method. The final package integrity ultimately depends on the seal quality. The seal protects the product from environmental influences and provides a barrier against oxygen to prevent spoilage of product and thus increases shelf life.

The seal layer is the last layer of a multilayer film which is in contact with the product inside the package. Generally, the seal layer is attached to the barrier layer by an adhesive or tie layer. The most common method of sealing is heat sealing. Heat sealing is one of the major processes of flexible packaging. Heat sealing is bonding two polymer surfaces by applying heat and pressure for a certain period of time. In heat sealing, the surfaces are forced into intimate contact while they are in at least a partially molten state. Many factors determine the quality of a heat seal. They can be classified as heat seal process parameters or film properties.

The process parameters influencing the seal are energy (often temperature), pressure, dwell time, and the machine design. Distinct levels of energy are required for different materials, thicknesses, package types, and processing steps. Dwell time is the time during which heat and pressure is applied. It should be easily adjustable to fractions of a second because of the economic concerns oblige the modern packaging lines to run as fast as possible. Likewise, the pressure between the jaws should be easily adjustable and uniform through the dimensions of seal area. These factors may vary for different materials.

The film properties are thickness, material characteristics, and post treatment of the film (e.g. corona treatment, printing). Material characteristics such as density, crystallinity, molecular weight, molecular orientations, polarity, and additives in the resin change the film properties and affect the sealing conditions and performances.

All of these factors tend to interact in a complex way. For example, the amount of heat available may be limited by the capacity of the heater elements, by the rate of heat transfer from the sealing bar and its coating, by the type of product being packaged, or by the amount of energy

consumption in order to maintain the jaw's temperature. Increasing the dwell time would permit increasing the heat available at seal layer surface, but this is not economically beneficial because fewer packages will be produced per minute and the production line speed will reduce.

At the microscopic scale, a variety of molecular mechanisms are engaged in heat sealing such as melting, wetting, inter-diffusion, and re-crystallization. During heat sealing, adhesion is developed between two surfaces. As a fast self-adhesion process, heat sealing has seldom been studied from a fundamental perspective. One reason is that it has to be performed in a very short dwell time far less than the time required for most welding and adhesion processes. Furthermore, the architecture of polymer chains plays a significant role on all micro-mechanisms involved in sealing. Moreover, there is a vast possibility of choices of materials to be used as seal layer. This makes it difficult to find an optimal combination of two or more polymers in blends to obtain a high quality product.

Research in this area has been inadequate and limited to the companies that produce materials and design seal layer for superior performance. Therefore, lack of fundamental studies in academic research labs led to limited information in open literature.

The main objective of this work is to understand how the process and material parameters affect the final seal quality. To achieve this goal, the first part of this thesis is dedicated to provide a clear understanding about the influence of seal process parameters on final seal quality. Then, several types of polyethylene resins were analyzed by performing systematic experiments to obtain a comprehensive knowledge about the role of molecular architecture of the polymer on heat seal performance. Finally, the role of blending a conventional polyethylene with metallocene catalyzed ethylene α -olefins copolymer was investigated. The correlation of rheological properties and heat seal performance of the resins was also studied.

This dissertation is based on three articles that have been accepted or submitted to scientific journals and consists of the following sections:

- Chapter 2 provides a broad literature review considering the related issues and followed by the originality and main objectives of this dissertation.
- The summary and organization of the articles are described in Chapter 3.

- The main achievements of the thesis are given in the format of three scientific papers in Chapters 4, 5 and 6.
- Chapter 7 presents a general discussion of the main results.
- Finally, Chapter 8 presents the final conclusions of this work and the recommendations for future works.

During this project and in collaboration with other members of 3SPack, we had the opportunity to compare the heat seal properties of blends of biodegradable polymers with the blends of polyolefin based copolymers. The result of this sub project has been published in International Polymer Processing (IPP), available in Appendix A.

CHAPTER 2

LITERATURE REVIEW

2.1 Heat seal process

Heat sealing is bonding two polymer surfaces while applying heat and pressure for a certain period of time. In heat sealing the surfaces are forced into intimate contact while they are in at least a partially molten state (Theller, 1989). There are several methods of heat sealing. The main differences between the common types of seal methods are in how they supply the energy and/or pressure to the sealant in the seal area.

In impulse-sealing, heat is supplied by sending an electrical impulse into a resistant wire or ribbon. In dielectric sealing which is specified for polar sealants, energy is provided by an alternating electrical field which heats up polar sealants. In laser sealing energy is provided by laser radiation and the sealing area absorbs heat from the laser light. In ultrasonic sealing energy is provided by friction of the sealants due to ultrasonic vibrations. Cold sealing is when the seal is achieved without heat and only by adding mechanical pressure. It is the alternative for heat sensitive products (Selke et al., 2004). Conductance sealers, also named as conductive sealer or heated bar sealer, are the most common type of heat sealers in commercial usages. This system typically consist of two metal jaws, one or both of which are electrically heated. (See Figure 2-1) Two films are brought into intimate contact by heated jaws and thus are bonded. The bar sealing is the subject of this study which uses the least expensive equipment compared to others, and it is the most preferred technique.

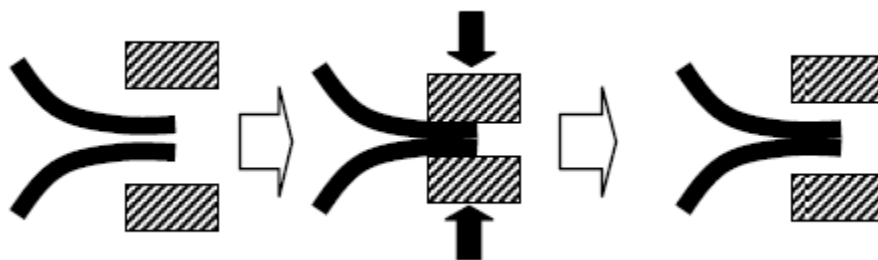


Figure 2-1 Schematics of heated bar sealing of two films

In bar sealing, the design of jaw surface affects the seal integrity and strength. Although the simple and ideal seal jaw might be flat, in practice there is a risk of damage in sealed area when there are folds or tucks in the films. The integrity of seal would thus be endangered. In some applications, the flat type seal bar is frequently used, but other different types are also common. Patterned, serrated, crimp or embossed seal bars give the seals extra strength (Selke et al., 2004; Theller, 1989). In the case of non-uniform film thickness, flat bars cause weaker seal than jaws designed with serrations. Serrated jaws can improve seal appearance. They are generally used to ensure that the two films are adequately stretched to have intimate contact (Theller, 1989). Figure 2-2 illustrates the vertical and horizontal serrations in the jaw surface.

The flat form of jaw was thus used in this study for simplicity and to prevent the complications in seal cross section and seal surface in jaw pattern. In order to prevent sticking, the molten seal material to the jaws, seal bars are normally covered by a layer of poly (tetrafluoroethylene) (Teflon®) or other types of non-stick coatings. It can prevent damage to the seal and jaw and buildup of residual material at the jaws surface (Troughton, 2008).

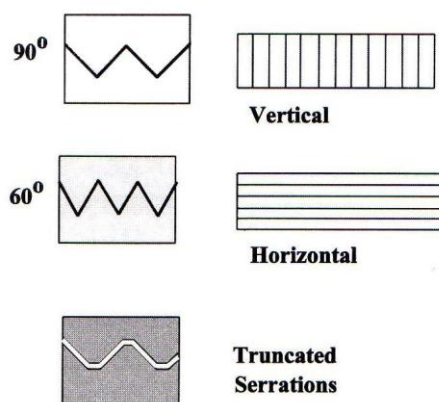


Figure 2-2 Cross section and top view of serrated seal jaws (Selke et al., 2004)

Stehling and Meka (1994) described the molecular mechanisms involved in the heat sealing of a single-layer film of semicrystalline polymers. As illustrated in Figure 2-3, both films are brought in contact and heated through the jaws. The crystals melt, and the pressure increases the molecular contact and the wetting of the two surfaces. If the contact time is sufficient, the polymer chains of the two surfaces diffuse through the interface and create entanglements. The seal interface is strengthened by molecular chain interpenetration across the interface. Only after

the polymer fully melts, enough penetration occur to ensure maximum seal strength (Stehling and Meka, 1994). Finally, cooling and crystallization strengthen the assembly.

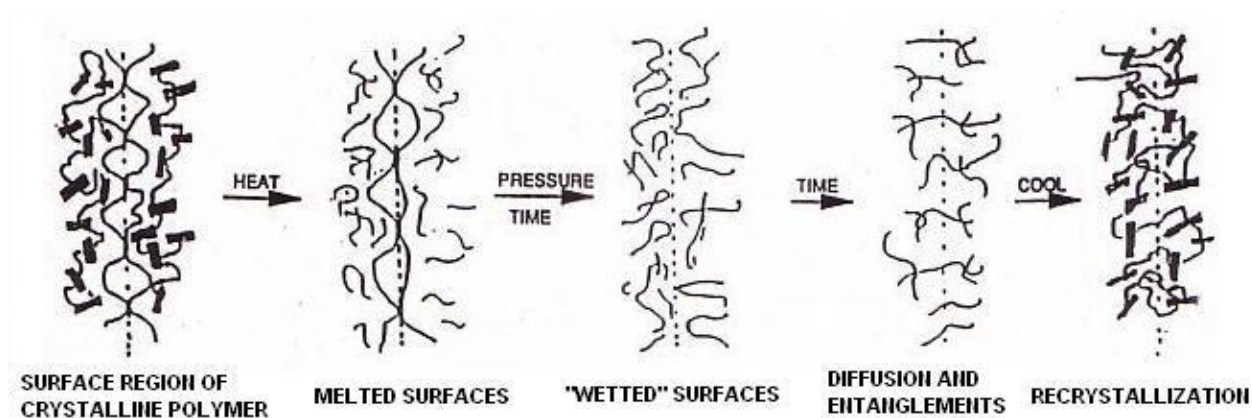


Figure 2-3 Molecular mechanisms involved in the heat sealing of two single-layer films (Stehling and Meka, 1994)

2.1.1 Heat seal properties

There are different properties by which seal performance is examined. The strength of the heat seal is often determined by the required force to pull apart the two films which have been sealed together. The standard procedures have been developed to measure these properties at two different stages: while polymers are still molten and hot, called hot tack, and after polymer reaches ambient temperature, called seal strength. These procedures enable comparisons between the seal performance of different materials and/or different sealing methods.

In this section, the main seal characteristics will be briefly summarized in order to build the theoretical base needed for studying heat sealing of semicrystalline polymers. We first introduce general parameters that are useful to interpret the adhesion test we used in our study.

2.1.1.1 Seal strength

Seal strength is the required force to separate the two sealed adherents. Measurement of seal strength between thin films is generally done using T-peel tests. As illustrated in Figure 2-4, the two legs of a test specimen are pulled at a certain rate and a force/width versus extension curve is obtained. Seal strength is defined as the average force per unit width of seal required to separate progressively two adhered specimens in T-peel test. (F02 Committee, 2000) At a certain

extension, failure of the joint occurs. Depending on the strength of bonded specimens, the seal might show different types of failure. The elongation of the test piece at failure is referred to as seal elongation, and the area under the curve is referred to as seal energy.

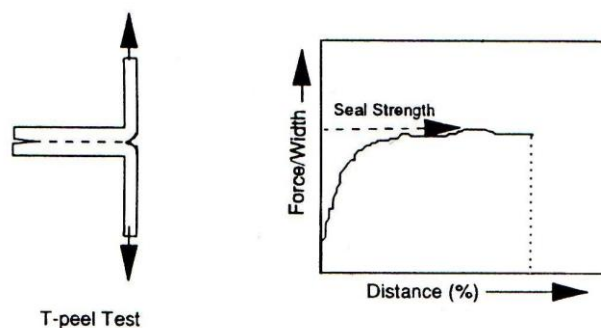


Figure 2-4 Schematic of the T-peel test and obtained force-displacement curve (Meka and Stehling, 1994)

2.1.1.2 Hot tack

Hot tack is the ability of the seal joint to withstand forces while it is still molten and soft. Operationally, it is defined as the strength of a hot seal measured immediately or at a specified delay time after completion of the sealing cycle, and before the temperature of the seal reaches ambient. (F02 Committee, 1998) The hot tack strength is very low compared to the heat seal strength because the latter is measured after the material is completely solidified and cooled to ambient conditions.

Hot tack is a critical property in vertical form-fill-seal (VFFS) modern machinery because the product weight exerts a force on the bottom seal while it is not solidified yet. The product is generally dropped into the package without any delay. Figure 2-5 illustrates a schematic of VFFS machine as well as schematics of different hot tack performance in response to the force applied by bearing the weight of the product. In Figure 2-5a, a deformed seal area with possibility of seal distortion and leakage is shown.

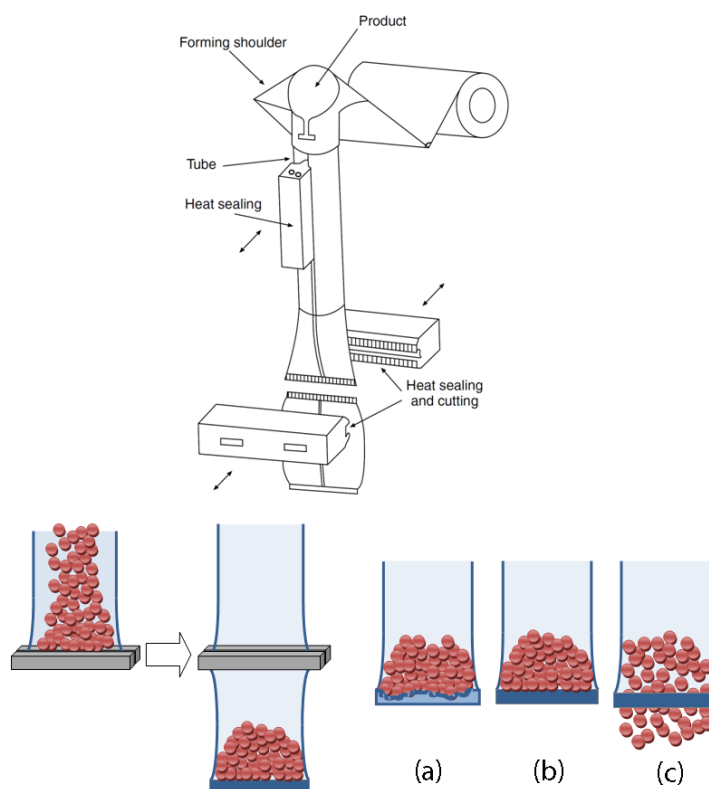


Figure 2-5 Schematic of a vertical form fill seal machine (VFFS) (Mesnil et al., 2000), and the schematic of weight bearing of seal in a VFFS process while it is still molten and hot (a) a deformed seal (b) a good seal (c) a weak seal with poor hot tack

In horizontal form-fill-seal (HFFS) applications, hot tack is also important, although the weight bearing burden is not present. In particular, sufficient hot tack strength is required to resist ‘spring-back’ forces in gusseted areas where the films are folded (Coles et al., 2003). High hot tack strength could avoid seal separation in gusset area which results in leakage. Figure 2-6 shows the schematic of horizontal form fill seal machine.

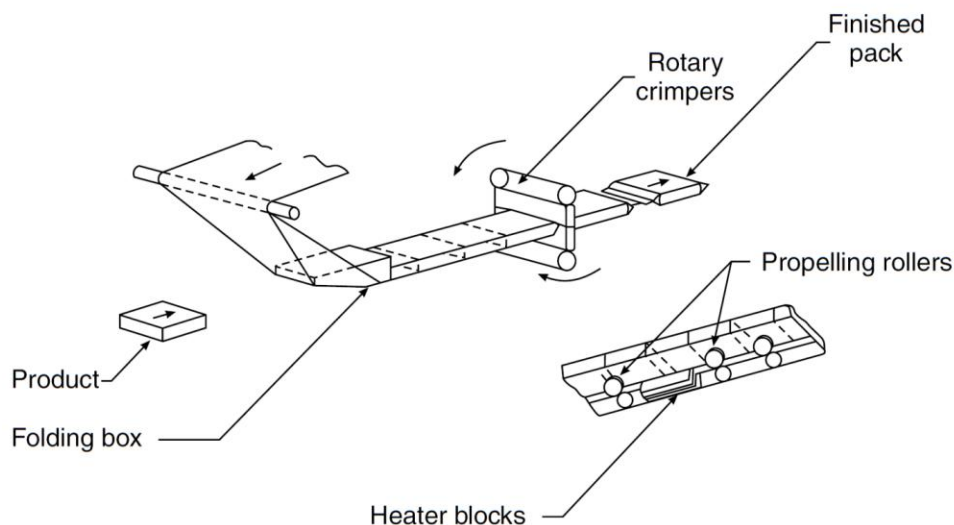


Figure 2-6 Horizontal form fill seal machine (HFFS) (Coles et al., 2003)

2.1.1.3 Failure mode

The failure of the joint is an important consideration in analyzing heat seals. If part of the seal structure fails, the whole package's integrity has failed. Many times, the failure does not occur within the actual seal area. Therefore, any thermo-mechanical factors which could cause weakening any of the regions adjacent to the seal area must be precisely monitored. When two materials are heat sealed together, three regions can be identified: the seal itself, the region adjacent to the seal, and the unaffected film region far from the seal interface.

Based on observations on monolayer and multilayer films, several types of failure have been reported in literature. The failure modes described in ASTM standard of seal strength and hot tack will be discussed here as the reference.

Peeling or adhesive failure mode (Figure 2-7a) in which the heat-seal bond peeled apart. The chains ends disentangle and withdraw from the opposite surface. This failure mode happens when the strength of the seal is poor. Cohesive failure mode (Figure 2-7b) occurs when the joint is strong enough to transfer the stress through the bulk of the seal layer. Delaminating failure mode (Figure 2-7c) involves tensile break of the seal layer, which is thin and weak, followed by the separation of seal from the support layer. Breakage failure mode (Figure 2-7d) is due to the weakening of the seal at its edge as the material fused, or the weakening of the laminate structures. Break or tear in film specimen (Figure 2-7e) distant from the seal area might happen

due to defects in structure or edges of film. Elongation (Figure 2-7f) occurs when the seal is strong but the other parts of the film are not strong enough to bear the stress and yield. Finally, a combination of these failure modes may also occur (Figure 2-7g).

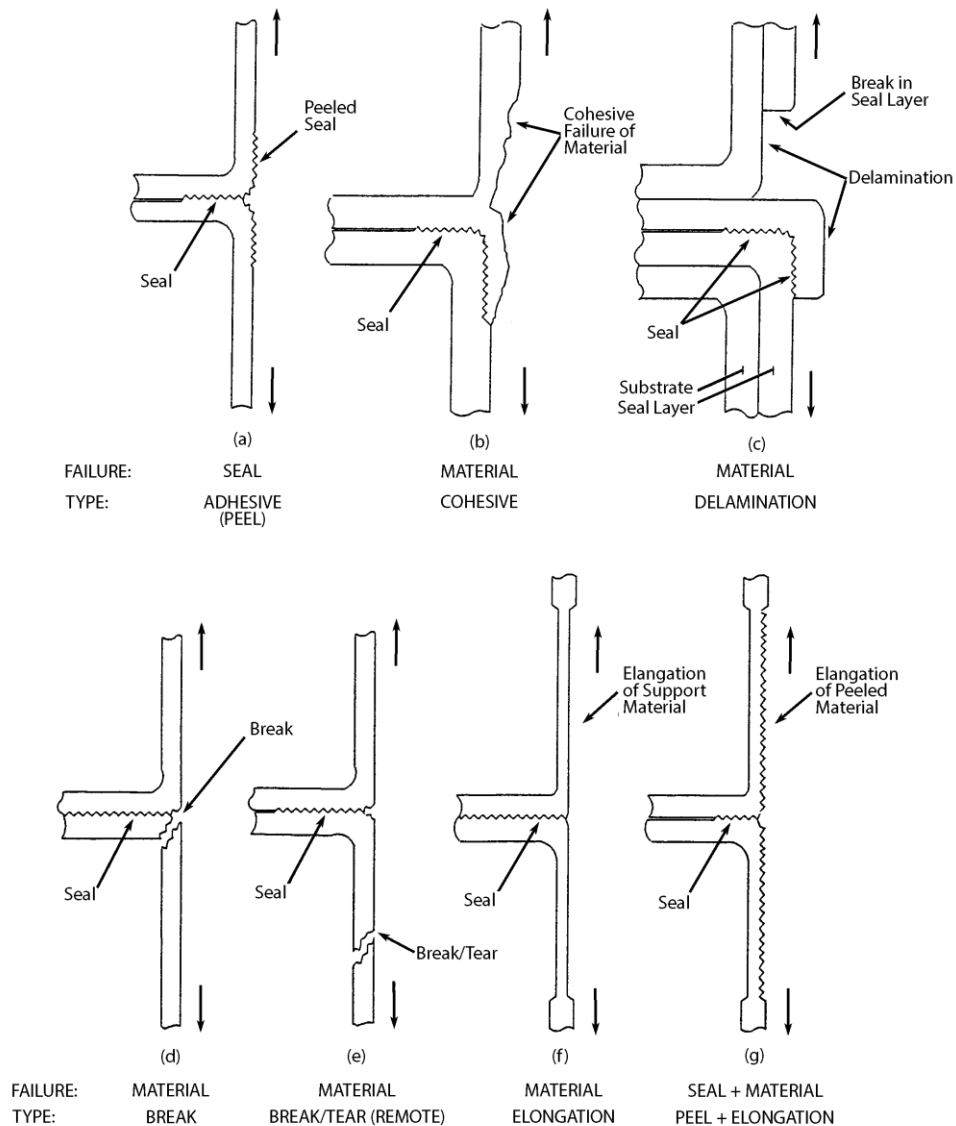


Figure 2-7 Failure modes of T-peel test for thermoplastic heat seal (F02 Committee, 2000, 1998)

2.1.2 Heat seal process parameters

Heat seal process parameters are the conditions of heat sealing. Temperature, dwell time, and pressure are recognized as the most important seal process variables (Gardon, 1963a, 1963b; Stokes, 1989). The adjustment of these three parameters controls the final seal properties.

2.1.2.1 Seal bar temperature

The strength of a heat seal is primarily determined by the maximum temperature achieved at the interface during heat sealing. For every semicrystalline polymer, the seal strength as a function of temperature was shown to behave similar to the curve presented in Figure 2-8. Ideally the temperature at the interface of the films should be considered as the controlling factor. However, measuring the value of temperature at the interface is practically challenging, and the required equipment for that is often unavailable and expensive. Therefore, the seal curve is commonly plotted as a function of bar temperature.

The schematic plot of seal strength curve, $SS_{(T)}$, introduced by Stehling and Meka (1994), can be approximately described by the following parameters illustrated in Figure 2-8:

- Seal initiation temperature T_{si} : the temperature at which measurable but weak seal strength is achieved.
- Plateau initiation temperature T_{pi} : the temperature where the plateau region begins.
- Final plateau temperature T_{pf} : the temperature where seal strength begins to drop off rapidly and extensive seal distortion sets in.
- Plateau seal strength SS_p : the ultimate seal strength value.

The temperature in which the ultimate seal strength could be achieved is thus a range from T_{pi} to T_{pf} which is referred to as temperature window of plateau seal strength.

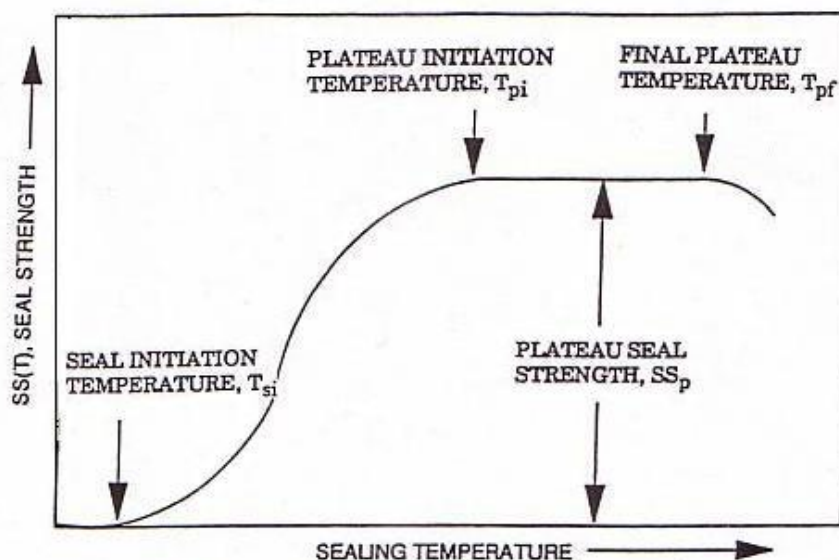


Figure 2-8 Schematic of general heat sealing curve, $SS(T)$, seal strength as a function of sealing temperature for semicrystalline polymers (Stehling and Meka, 1994)

2.1.2.2 Dwell time

Dwell time is the period of time that the two films are brought into intimate contact by the heated bars. The heated bars are in direct contact with films so heat could be transferred to the interface. The greater the heat flow rate, the shorter the dwell time required. Thus, the heat flow rate determines the dwell time of the process. Generally, the dwell time in modern flexible packaging industry is of the order of fractions of a second, or in some cases 1-2 seconds. The optimum dwell time ensures no excessive time is wasted, to keep up the production speed.

2.1.2.3 Pressure

Pressure is required to ensure the film surfaces are in intimate contact for interfacial penetration to occur. Excessive pressure, however, may result in the squeeze-out of the molten film from the seal area. For most heat seal materials, pressure is less important than either temperature or dwell time (Hassan, 2007; Stehling and Meka, 1994). Furthermore, in industrial production lines, higher pressure may be required if the pressure control is not precise, the film thickness is non-uniform, or the heated bars are poorly aligned (Selke et al., 2004).

2.1.3 The effect of process parameters on heat seal properties

Theller (1989), the pioneer in heat sealing variable investigations, examined the seal strength of LDPE film with serrated jaws. He showed that at 379°K, LDPE required 280 ms to reach a seal strength of 0.53 kN/m and at 383°K required 150ms. Regarding pressure, from 1 to 600psi he reported no effect of pressure on the seal strength of LDPE. Meka and Stehling (1994) applied a finite element (FE) model based on conduction heat transfer to predict the interfacial temperature of the seal as a function of dwell time. The results of the model were verified by a micro-thermocouple. The heat of fusion, conductivity, and film-thickness changes occurring during sealing were not considered in this model. Therefore, their model was not valid for temperatures above the final melting point of thermoplastics. Meka and Stehling (1994) also reported that the heat seal strength is primarily controlled by sealing temperature and dwell time, rather than pressure.

Tetsuya et al. (2005) studied oriented polypropylene (OPP)/cast polypropylene (CPP) laminate films. They reported that the tensile strength of the seal was affected by the orientation of the films. The SEM images from seal cross section revealed that the seal was not well formed at lower temperatures, while the laminates were totally fused together at high temperatures. In Figure 2-9, the interface of the CPP layers clearly indicated that the heat seal was not well formed. In Figure 2-10 the boundary is not clear and this essentially implies more complete heat sealing. At 170°C, melting and interdiffusion of polymer chains across the interface resulted in the formation of a good heat seal. This resulted in higher peel strength. In

Figure 2-11 the boundaries between the CPP-CPP heat seal in OPP/CPP laminated films are no longer distinguishable. The temperature of 250°C is much higher than the melting temperature of both OPP and CPP films, so both films melted during heat sealing.

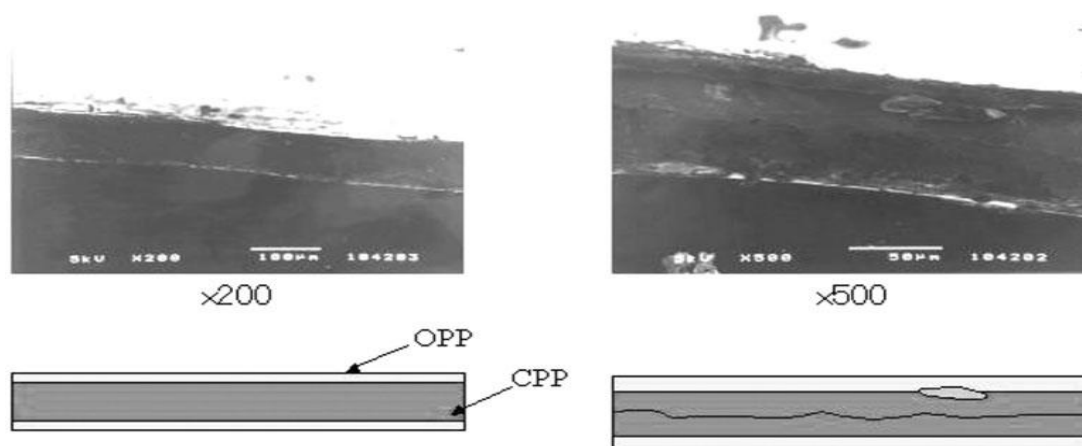


Figure 2-9 SEM micrographs and schematic diagrams of cross sections of OPP/CPP laminated films, heat sealed at 115°C in two magnifications (Tetsuya et al., 2005)

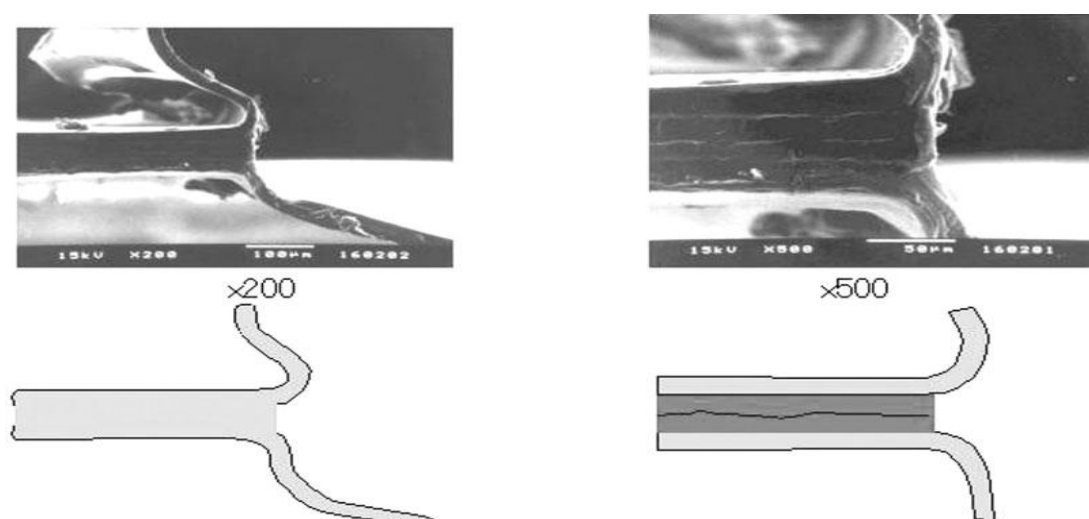


Figure 2-10 SEM micrographs and schematic diagrams of cross sections of OPP/CPP laminated films, heat sealed at 170°C in two magnifications (Tetsuya et al., 2005)

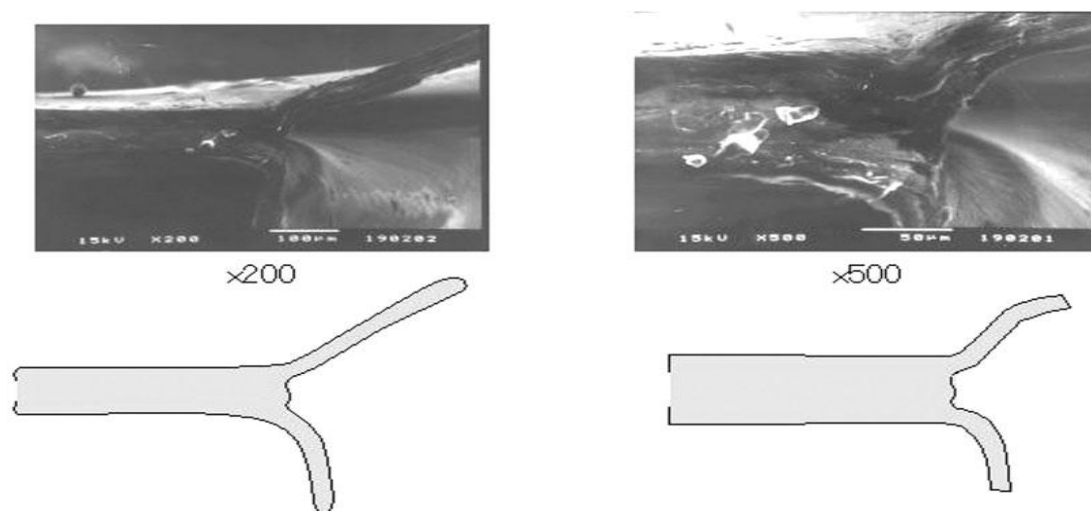


Figure 2-11 SEM micrographs and schematic diagrams of cross sections of OPP/CPP laminated films, heat sealed at 250°C in two magnifications (Tetsuya et al., 2005)

In another study Tetsuya et al., 2006 carried out investigations on the failure criteria of the OPP/CPP heat seals made by impulse heat sealing. They reported that heat seals were stronger at temperatures low enough not to destroy the level of crystalline orientation in OPP as support layer. The possibility of breakage in the heat seal edges because of the existence of the so-called weak spots and also the deformation of seal area was also emphasised (Hashimoto et al., 2006; Tetsuya et al., 2006).

Poisson et al. (2006) reported that blending ethylene vinyl acetate (EVA) to LDPE in the seal layer positively influence sealability. A decrease in melting temperature, reduction in crystallinity, and finer crystalline structure in EVA/LDPE shifted the seal initiation temperature to lower temperatures. (See Figure 2-12) Enhancement in seal strength was observed because of strong intermolecular forces appearing between the C=O polar groups of EVA. This mechanism is reinforced by the migration of the EVA to the surface of the seal layers (Poisson et al., 2006a, 2006b). The seal strength of EVA/LDPE as a function of pressure was also studied. As illustrated in Figure 2-13, by increasing pressure, the seal strength increases and then decreases. In spite of the minor effect of pressure mentioned in other studies, pressure might be influential in seal strength performance.

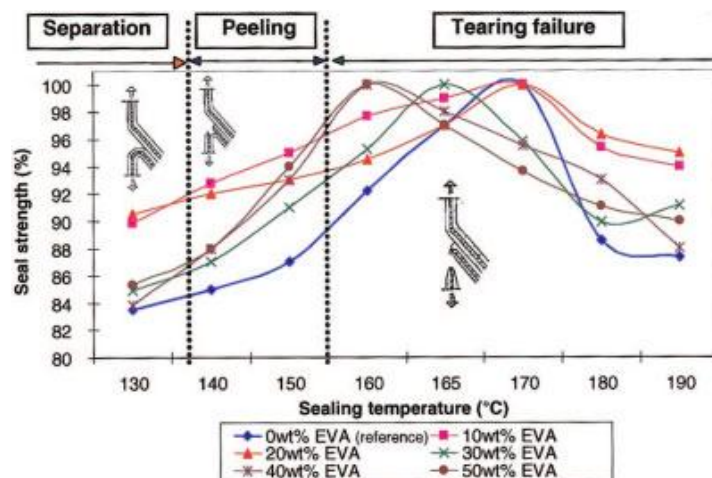


Figure 2-12 Effect of sealing temperature on the seal strength of EVA/LDPE film (Poisson et al., 2006a)

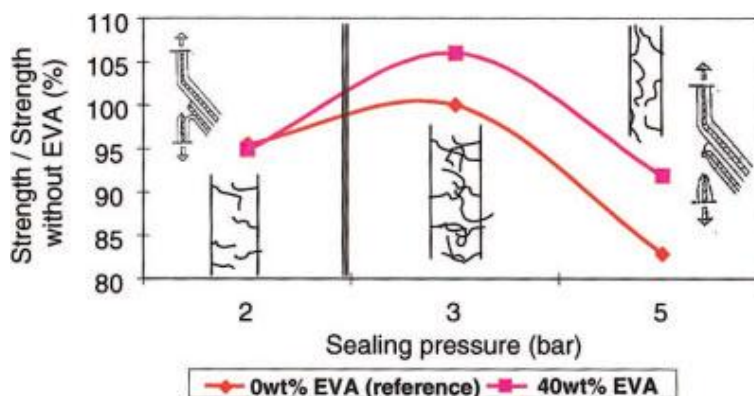


Figure 2-13 Effect of pressure on the seal strength for EVA/PE 170°C (Poisson et al., 2006b)

The interrelationship between seal bar temperature, dwell time, and package design on heat seal strength was studied for ionomers (Morris, 2002). It was shown that a thicker packaging structure required longer dwell times for a given seal bar temperature to reach the same heat seal strength compared to thinner structures. This was due to heat transfer considerations; it takes longer for the heat to transfer through thicker films.

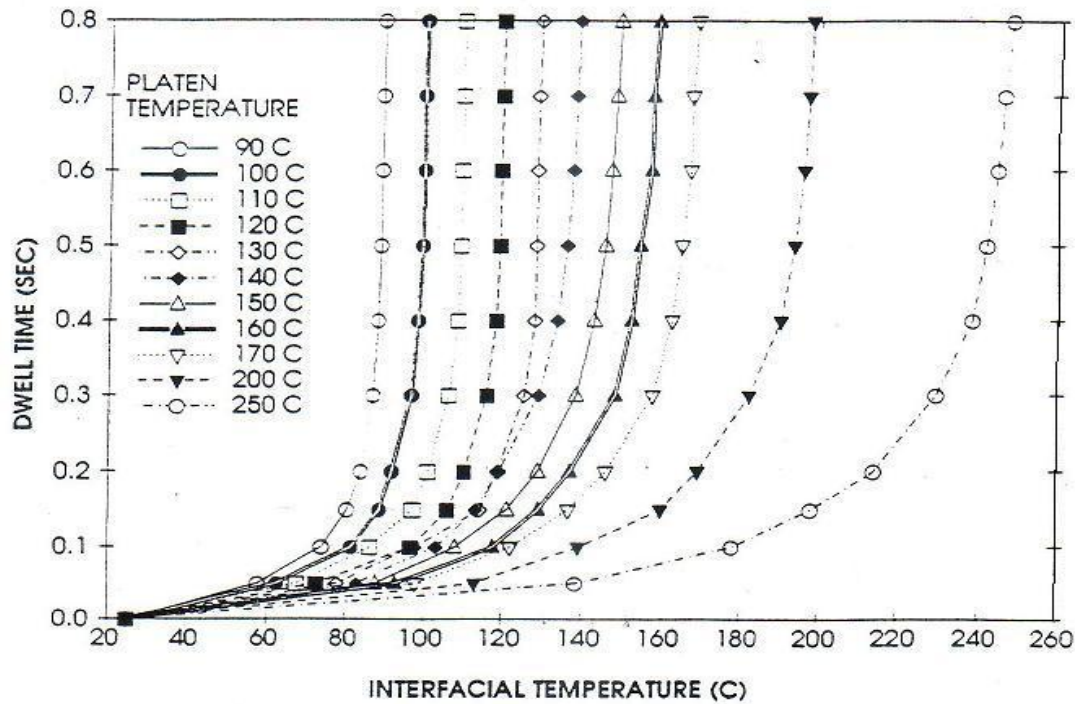


Figure 2-14 Interfacial temperature as a function of dwell time as calculated by the finite elements model under different platen temperatures

Meka and Stehling, (1994) used heat transfer finite elements model to obtain the temperature at interface and they showed that the interface temperature will reach to the jaw temperature after 0.3-0.5 s. They measured seal strength of several polyethylene resins based on the interface temperature which is not practically possible to measure in an industrial process. They concluded that the seal strength is strongly dependent to interface temperature (Stehling and Meka, 1994). In an industrial process however, the heat sealing occurs at a non-isothermal condition and temperature setting is based on the control temperature on the heated jaws.

Mueller et al., (1998) studied the time dependence of welding two films at long contact times of 1 - 1000 s and showed the linear dependence of peel strength to the square root of contact time. The contact time they had applied however was much longer than the short contact times usually used in heat sealing.

There have been several studies on the influence of heat seal variables on sealability. However, to our knowledge there has been no comprehensive study in open literature on the interactive effects of these variables, trends of their influences and the micro-mechanism involved in the process.

2.1.4 Lamination

Generally, the seal layer is a part of a multilayer film for which every layer provides a certain advantage to the whole structure. In heat sealing, these other layers protect the seal layer from sticking to the jaws and deformation, because direct contact to the jaws weakens the seal properties. Furthermore, a strong bond between seal layer and other layers is necessary for high hot tack and seal strength. Excluding the multilayer films produced in coextrusion, the combinations of two or more polymer films is known as lamination. Sometimes, a metal foil or paper web may also be used as substrates. There are several methods of lamination which can be categorized into two main types of extrusion and adhesive lamination. Hot-melt lamination and extrusion lamination are processes that apply a heated adhesive or thermoplastic to the substrates, and joining them as it cools down.

In adhesive lamination, adhesives are used to hold the films together into a single structure. There are several methods of adhesive lamination. Depending on the type of the adhesive, they can be solvent-based, water-based, or solvent less. Generally, it is required to modify the film surface in order to generate radicals and active groups through corona or plasma treatments. Reaction between the materials and the adhesive is necessary for strong lamination. The complete wetting of the surface to create flawless laminates is important, especially for non-polar materials such as polyolefins (Selke et al., 2004).

2.1.5 Seal layer material

A wide range of materials has been used as a seal layer for flexible packaging applications, either in laboratory or industrial scale. Polyolefins including polyethylene and polypropylene families, ionomers, ethylene vinyl acetate copolymers, biodegradable resins such as poly(lactic acid) and polycaprolactone (PCL), are the most utilized resins for seal layer applications (Hanlon et al., 1998; Wagner, 2009). In spite of many efforts carried out by raw material producers, no existing polymer is able to combine all the properties required by some challenging industrial packaging applications. Therefore, melt blending has been an alternative.

For the purpose of this study, a number of polyethylene resins have been used. Therefore, in this work we will focus on this group of polymers and describe variety of molecular architectures as well as their heat seal properties.

2.1.5.1 Low Density Polyethylene (LDPE)

Low-density polyethylene, LDPE was one of the first commercially produced polyolefins in the early 1940s by the free radical polymerization of ethylene using a high temperature and high pressure process (Brydson, 1999). The product of this polymerization is a broad MWD polyethylene with chains containing branches of many different lengths distributed non-uniformly throughout the chain backbone. (See Figure 2-15) The main advantage of LDPE is its high melt strength and good processability, especially in film production processes, e.g. film blowing and casting.

However, there are some disadvantages for LDPE in seal layer which makes it impractical for some applications. For instance, it can be difficult to open the package because of the way the film stretches without tearing (Hanlon et al., 1998). Because of the high elongation of LDPE, when opening a seal of a LDPE/PET laminated structure, the PET layer may break first. High crystallinity, and thus high T_{si} , is not desirable for many application and equipments (Selke et al., 2004). Therefore LDPE is used mostly in the form of blends with LLDPE, metallocene catalyzed PEs, or other copolymers such as EVA or ionomers (Nase et al., 2009; Poisson et al., 2006a). Blends of LDPE/LLDPE at several compositions could provide a good balance of processability, mechanical, optical, and heat seal properties (Colls et al., 2006).

2.1.5.2 HDPE

The development of highly active catalysts made polymerization possible at lower temperatures and low pressures. Then, the production of linear PE (or high-density polyethylene, HDPE) was developed in the mid-1950s (Brydson, 1999). HDPE chains are linear, containing neither short nor long chain branches. (See Figure 2-15) Application of HDPE in seal layer is very limited due to its high crystallinity, high T_m , difficult melt processing, and lack of flexibility. Therefore, it is used in the form of blends with other resins and in easy-open heat seals (Malsen et al., 2008; Miyata and Toshiyuki, 2011; Theller, 1989).

2.1.5.3 Linear low density polyethylene LLDPE

Further developments in catalyst technology led to the possibility of copolymerization of ethylene with small amounts of an α -olefin (Hosoda, 1988; Mirabella and Ford, 1987). This method of polymerization incorporates short side-chain branches to the ethylene backbone.

Linear low-density polyethylene (LLDPE), was first developed commercially in the late 1970s (Brydson, 1999). LLDPEs catalyzed by Ziegler-Natta type catalysts (Zn-LLDPE) exhibit broad molecular weight distributions (MWD) and short chain branching distributions (SCBD) because Ziegler-Natta catalysts have multiple active sites. The distribution of short chain branches is not uniform among and along the chains. (See Figure 2-15) Generally, the shorter chains have higher SCB content than longer chains (Wild et al., 1982). Thus, the Zn-LLDPEs has heterogeneous structure at both inter- and intramolecular levels. The Zn-LLDPEs are widely used as seal layer in both forms of neat and blends with other seal polymers.

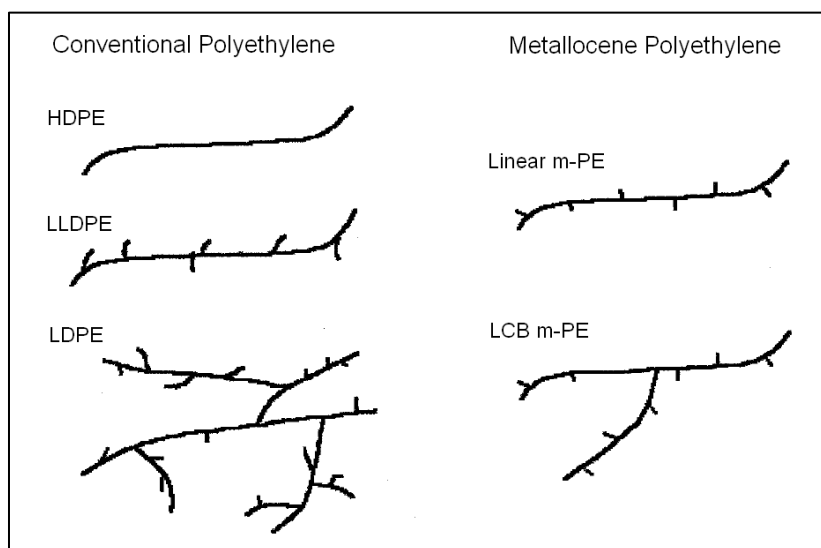


Figure 2-15 Schematic of the molecular structure of the different polyethylenes, (Wood-Adams, 1998)

2.1.5.4 Metallocene catalyzed ethylene α -olefin

One of the most recent major progress in the area of polyethylenes catalysis systems has arguably been single site catalysts (Feldman, 1996). Advances in organometallic chemistry resulted in the development of metallocene catalysts that have been used mainly in the production of ethylene and propylene polymers. Metallocene-catalyzed polyolefins, in particular, have been produced commercially since the late 1990s (Brydson, 1999). Although the use of metallocene-catalysts is well established, the full potential of this technology is probably yet to be fully realized. More recent advances in single-site metallocene-catalysts have resulted in the production of structurally superior PEs (Brydson, 1999). Metallocene-catalyzed PEs (m-PEs) are ethylene copolymers with

uniform incorporation of the comonomer and have a narrower molecular weight distribution MWD.

There are two subclasses within the general category of m-PEs: linear m-PEs which have no branches or only short chain branches but no LCB, and the branched m-PEs which containing precisely controlled low levels of LCB. These uniformly distributed LCB contain chains are produced using the constrained geometry catalysts. The terms, very low-density polyethylene (VLDPE) or ultra low-density polyethylene (ULDPE), are often used to describe metallocene-catalyzed PEs (Brydson, 1999; Halle, 2003; Khare et al., 2000; Manaure and Müller, 2000; Manaure et al., 1997; Shanks et al., 2000; Tanrattanakul and Udomkichdech, 2001).

Due to the inherent plastic and elastomeric features in some m-PE grades, they are often referred to as plastomers (Halle and Davis, 1995; Halle, 2003). Although, m-PEs are relatively new, the properties of these materials are well characterized and established in the literature (Chum et al., 2000; Jordens et al., 2000; Nitta and Tanaka, 2001; Nitta et al., 2000; Razavi-Nouri and Hay, 2001; Vega et al., 1996; Wang et al., 2004).

The unique design of chain structure for metallocene catalyzed polyethylene resins resulted in superior toughness, heat sealing, and clarity compared to the conventional PEs. These superior properties have led to their extensive use in stretch films, heavy duty sacks, and flexible food packaging such as frozen food films. As seal layer in all types of packaging, metallocene catalyzed resins have been used in both neat forms and as a minor phase in blends with other polyolefins. There are several patents on application of metallocene as variety of film structures and blends (Donovan et al., 1999; Farley et al., 1996; Van Loon et al., 2009).

In a paper presented in a TAPPI conference Halle, 2003 showed the superior hot tack strength of metallocene catalyzed PE resins compared to EVA. (See Figure 2-16) They also studied the blends of plastomers and LLDPE and reported the enhancement of hot tack by incorporation of plastomers to LLDPE (Halle, 1997). Shih et al. (1999) studied a blend of 30wt%LDPE and 70wt%m-PE. They reported that the film made of m-PE/LDPE blend has higher hot tack than m-PE. However, they measured the hot tack after 0.4 s delay time, where the film already cooled and partial crystallization occurred. They explained this result with the gap between crystallization temperatures of the two components, which caused reinforcement of hot tack by some crystal formation (Shih et al., 1999).

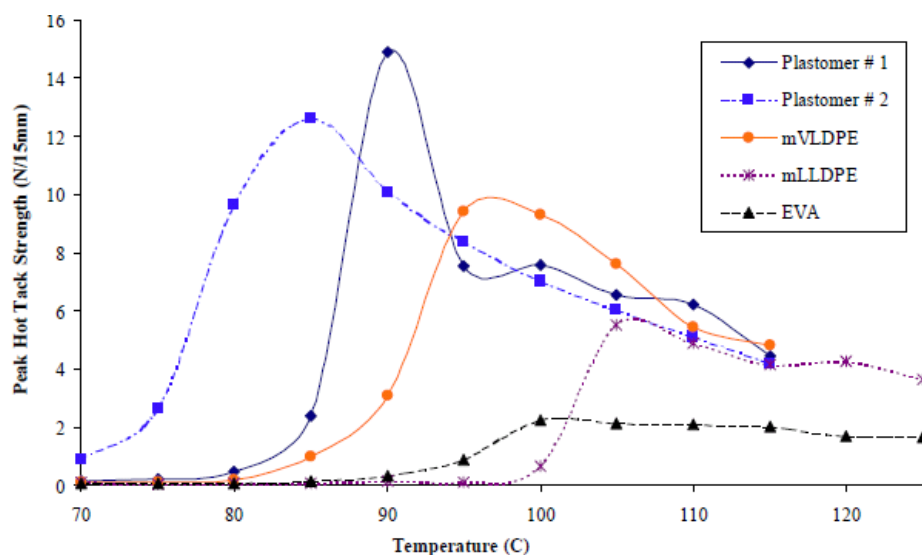


Figure 2-16 Hot tack measurements of the different types of m-PE compared to EVA (Halle, 2003)

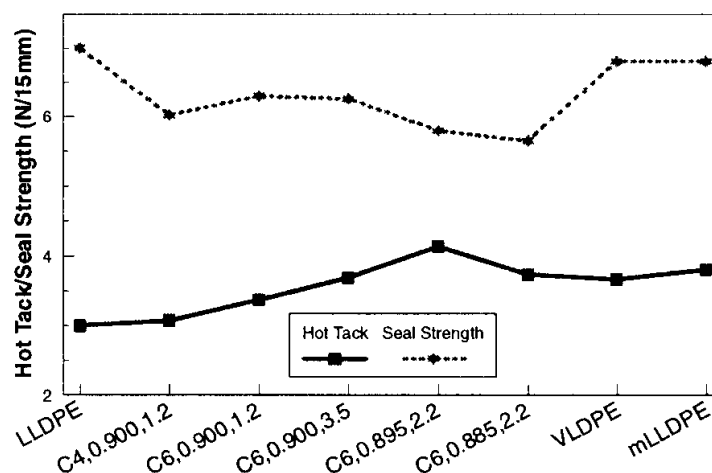


Figure 2-17 Peak hot tack and plateau seal strength of “25% metallocene plastomer-75% LLDPE” blends (Halle, 1997)

2.2 Adhesion between polymers at their interfaces

Polymer-polymer interfaces can be categorized into two general categories of symmetric (A/A) and asymmetric (A/B) interfaces (Wool, 1995). The symmetric interface occurs when the same polymer is on both sides of the interface. Asymmetric interfaces are the case where two dissimilar polymers are being blended, coextruded, laminated, recycled plastics, etc. The two polymers in asymmetric interfaces might be compatible or incompatible. They might be even

from the same chemical structure but with different molecular weight or chain architecture. Therefore, the number and complexity of asymmetric interfaces are far beyond symmetric interfaces (Boiko and Prud'homme, 1999; Brown, 1991a; Lee, 1991).

Because the focus of this study is the heat sealing process of polymer films in flexible packaging applications, the adhesion of symmetric interfaces will be reviewed in this section. The primary interest here is to describe adhesion between surfaces that had not been in contact previously. Thus, the literature and theories of crack healing and fractured surfaces are not of interest here. Thus, in the following section we will review the adhesion between pairs of polymeric materials, and more specifically the formation of interfaces between non-reactive polymers.

The majority of studies on polymer adhesion in this subject focused on the welding of polymers for a certain time (usually a few minutes or hours) (Brown, 1991a; Hamed and Shieh, 1983; Kim et al., 1994; Klein, 1990; Wool, 2006) in a certain temperature. In the studies on welding of polymers, the strength of the interface is examined using several types of geometries (Wool, 1995). However, the adhesive fracture energy is the main criteria to examine the strength at polymer interface. The work of adhesion or adhesive fracture energy is the irreversible fracture energy of interface. It is defined as the energy required for crack propagation through the material by a unit length in a specimen of unit width (Brown, 1991a; Zhang and Rong, 2012).

The adhesive fracture energy known as G_c , will encompass all forms of the energy losses incurred around the crack tip. Similar to ordinary fracture mechanics, the adhesive fracture energy is determined by the local dissipation energy at crack tip during fracture (Brown, 1989; Xu et al., 1991). At the microscopic scale, crack propagation is mainly led by the growth of a craze at the crack tip (Brown, 1991b; Miller et al., 1991). Crazing is a deformation mechanism which generally leads to a significant increase in fracture toughness (Wool, 1995). The craze corresponds to the plastic response of a material at the crack tip and the energy dissipation occurs mainly by crazing (Brown, 1991b; Kramer and Berger, 1990; Kramer, 1983; Miller et al., 1991). As polymer chains diffuse across the interface during welding, more material will be involved in the deformation process during the fracture test (Creton et al., 1992). Until the adhesive fracture energy reaches the fracture energy of bulk polymer. The main complication here is to connect the examined fracture energy at interface with microscopic diffusion parameters.

We will discuss the interdiffusion mechanism at polymer interfaces in more details in the following section. We will review the literature on adhesion of polymers, especially self-adhesion studies which is also the case encountered in our work.

2.2.1 Wetting

In the adhesion of polymer interfaces, several stages were described by Wool and O'Connor (1981): surface approach, wetting, diffusion, and finally randomization.

When two polymer surfaces are brought into contact, the level of proximity at the microscopic scale is determining in the formation of an adhesive bond. The term wetting implies that the two material surfaces should spread over each other in order to displace air or any other contamination that may be present between them (Cherry, 1981). The topography and roughness of the surfaces, rearrangement of the roughness after contact, how it modifies with time, temperature and pressure, should be considered important in the wetting stage.

Depending on pressure, temperature, chain orientation, presence of crosslinks between chains or crystal structure at the surfaces, wetting might be time dependent (Brown, 1991a; Frederix et al., 2013; Wool et al., 1989; Zhang and Rong, 2012). The wetting kinetic is a two dimensional nucleation and growth process. The wetted spots are nucleated at random locations at the interface and grow until a complete wetting is achieved.

It is known that wetting is a necessary but not a sufficient step for mechanical recovery and interdiffusion is necessary for strength development (Boiko and Lyngaae, 2005; Brown, 1991a). After the wetting stage, the chains are able to move across the interface in the subsequent stages of diffusion and randomization. However, the interdiffusion occurs in the wetted areas, so the wetting stage convolutes with the diffusion stage and affects the strength development at the interface (Wool, 2008, 1995). By applying a light pressure, the achievement of intimate contact is relatively fast for melted polymeric interfaces (Boiko and Prud'homme, 1998; Brown, 1991a).

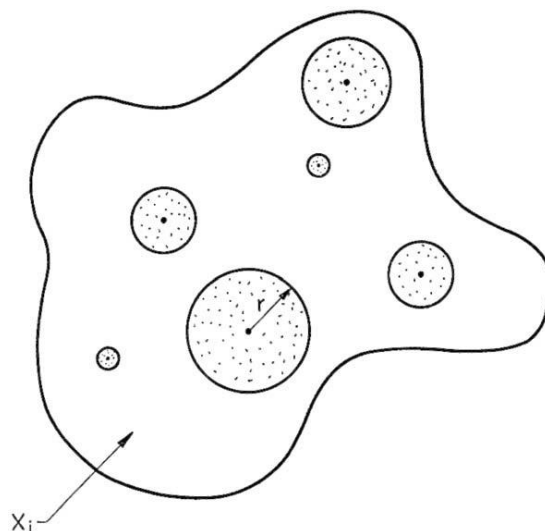


Figure 2-18 Partially wetted domains at interface, contacted regions of radius r grow and coalesce. Interdiffusion occurs only in the wetted regions while surface rearrangement occurs in the non-wetted areas (Wool and O'Connor, 1981; Wool et al., 1989).

2.2.2 Interdiffusion at polymer interfaces

The interdiffusion mechanism is principally based on the framework of the reptation motion model of polymer chains introduced by de Gennes (1971, 1979) and Doi and Edwards (1979, 1978a, 1978b, 1978c). This theory considers a single chain trapped in a network. In a melt, the chains can move by Brownian motion, but they cannot intersect each other. The chain is not allowed to cross any obstacle but can move in between in a wormlike fashion which is referred as reptation. The concept of tube was introduced by Edwards, the tube which contains the chain as represented in Figure 2-19. The chain goes back and forth along the centre line of the tube, change its conformation, and disengage itself from the tube that was defined at an earlier moment. The microscopic details of polymer chain dynamics have been examined by many researchers using experimental, theoretical, and computer simulation approaches (Basin, 1967; Kunz and Stamm, 1996; Russell et al., 1993; Stamm et al., 1991; Voyutskii et al., 1966; Zhao et al., 1993).

The interdiffusion of polymer chains across the interface is inherently an unsteady state process. At the earliest times of contact, the chain ends at the interface initiate interdiffusion, producing a fast but very thin broadening of the interfacial region (Foster and Wool, 1991; Schweizer, 1989). Then the rate of interdiffusion decreases and following reptation dynamics deep interdiffusion

occur. Experimental observations indicated a good correlation between the observed experimental time and the different time regimes given by the reptation theory (Kunz and Stamm, 1996; Stamm et al., 1991). Favorably, the studies on interdiffusion have progressed with the development of new techniques, such as neutron reflectivity (NR) or dynamic secondary ion mass spectroscopy (DSIMS). The development of the reptation model for the description of polymer self adhesion and better understanding of the fracture behaviour of amorphous polymers, initiated a variety of studies.

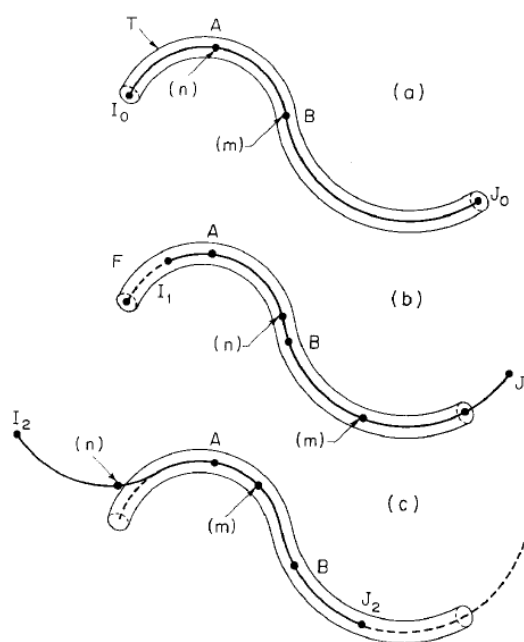


Figure 2-19 Reptation motion of a chain, (a) Initial position: the chain is restricted to a tube, (b) the chain moved along its tube to the right by reptation, (c) The chain moved to the left and exit its original tube, but a certain fraction of the chain is still trapped in the initial tube at stage. (de Gennes, 1971)

The buildup of strength at the interface of two polymers has been discussed by several researchers (de Gennes, 1983; Kausch and Tirrell, 1989; Kline and Wool, 1988; Schnell et al., 1999, 1998; Wool, 2008, 1995; Wool et al., 1989). Several models have been proposed to find correlations between the examined fracture energy of interface in amorphous polymers and the reptation model parameters. The bridge model developed by Prager and Tirrell (1981) and de Gennes (1983), was based on the assumption that the number of bridges crossing the interface, the so called “crossing density” determines the adhesive fracture energy. They discussed that the

crossing density is simply proportional to the adhesive fracture energy, which result the following scaling law:

$$G_a \sim M^{-3/2} t^{1/2}$$

where M is molecular weight and t diffusion time.

Another model is Wool's minor chain model (Wool et al., 1989) which assumes that the average monomer interpenetration distance, $X(t)$, determines the adhesive fracture energy. From the reptation theory, it is known that $X(t)$ scales with the average contour length, $\langle l(t) \rangle$, of those parts of the chains which have escaped the initial tube, referred as "minor chains", as $X(t) \sim \langle l(t) \rangle^{1/2}$. The adhesive fracture energy, G_a is simply assumed to be proportional to the average length of the minor chains $\langle l(t) \rangle$. Therefore, G_a exhibits the same scaling laws with respect to molecular weight and welding time as $\langle l(t) \rangle$:

$$G_a \sim M^{-1/2} t^{1/2}$$

Both welding models mentioned above predict that the adhesive fracture energy will increase with the square root of welding time. This scaling law is confirmed by experimental data on both glassy polymers (Bastien and Gillespie, 1991; Foster and Wool, 1991; Fowler et al., 1987; Kline and Wool, 1988; Yoo et al., 1991) and elastomers (Wool and O'Connor, 1981). The experimental data on the influence of molecular weight on the adhesion fracture energy was reported to favour the minor chain model (Wool and O'Connor, 1981).

2.2.3 Polymer adhesion between amorphous polymers

In this section, different studies on the fracture toughness of interfaces between amorphous polymers will be reviewed. In the early 90's, the advances made on the understanding of the micromechanisms of interdiffusion and fracture at interfaces and the development of neutron reflectivity as a technique to measure interfacial width between polymers, with angstrom resolution, reinforced progress of polymer adhesion.

In welding of glassy interfaces of amorphous polymers, the interfacial width and the average distance between entanglements are the parameters that control fracture toughness (G_c) (Brown, 1991a). It has been shown that the chains need to diffuse over a certain distance in order to

entangle and therefore transfer the stress between the two surfaces. The chains act as connectors and the presence of entangled connectors at the interface reinforce the interface.

It has been found by several research groups that the G_c is directly linked to the interfacial width (Benkoski et al., 2002; Brown, 2001; Schnell et al., 1999, 1998). In the welding of some monodisperse amorphous polymers at temperatures higher than T_g , the fracture toughness was measured as function of the range of interfacial widths. The direct measurement of the interfacial width was carried out by neutron reflectivity. The results are summarized in Figure 2-20. The variation of G_c with interfacial width is largely non linear. Three different regimes were introduced in terms of microscopic failure mechanisms. In regime I, G_c is low and presumably the failure mechanism is simple chain pull out or simple chain scission. In regime II, G_c increases sharply with interfacial width, suggesting a transition from pullout or scission to crazing. In regime III, the failure stress becomes independent of the interfacial width and G_c is that of the bulk polymers. The transition from regime II to III is the stage that the interface can no longer be distinguished from the bulk by its fracture mechanism. It is believed that the transition occurs from chain pullout to crazing at the point where the bridge chains could sustain the stress higher than the crazing stress.

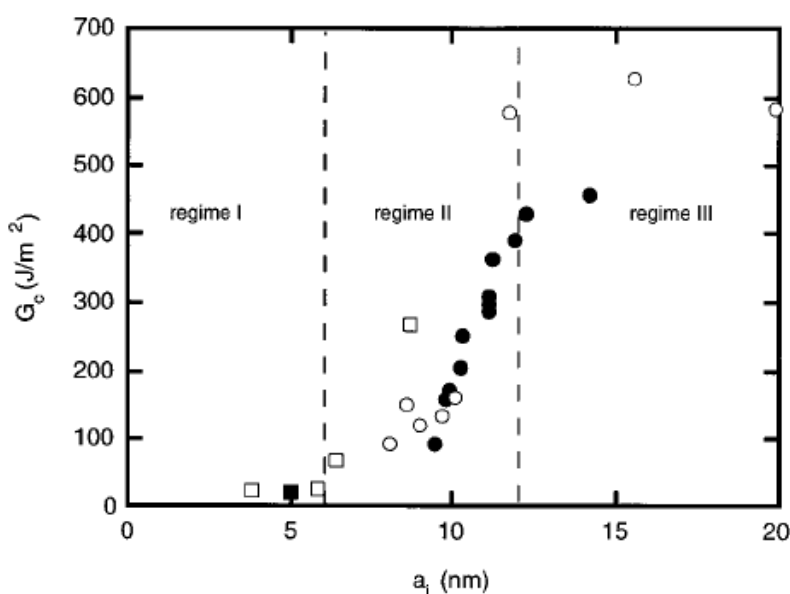


Figure 2-20 Fracture toughness, G_c , of different pairs as a function of the interfacial width a_i ;

□ poly(bromostyrene-styrene)/polystyrene (PBr_xS/PS) interfaces;

■ polystyrene/polymethylmethacrylate (PS/PMMA) interfaces (Brown, 1990) ;

• polystyrene/poly(p-methylstyrene) PS/PpMS interfaces; ○ polystyrene/polystyrene (PS/PS) interfaces. (Schnell et al., 1999)

2.2.4 Self-adhesion of semicrystalline polymers

Boiko et al. (2001) studied the self adhesion of polyethylene terephthalate (PET) in both amorphous and crystalline form. They quenched molten PET films to obtain the amorphous specimens. They annealed some samples at 180°C for 10 minute to prepare the semicrystalline specimens. The samples were bonded in a lap-shear joint geometry at temperatures varying from 64°C ($T_g-17^\circ\text{C}$) to 108°C ($T_g+17^\circ\text{C}$) for contact times from 5 minutes to 15 hours, cooled to room temperature, and submitted to a tensile loading at a cross head speed of 0.5 cm.min⁻¹. Shear strength was calculated as the measured force at break divided by the contact area. The shear strength for amorphous/amorphous PET interfaces in the vicinity of T_g was one order of magnitude higher than for crystalline/crystalline PET interfaces. In crystalline/crystalline interfaces the molecules are trapped in the crystals so the interdiffusion occurred only over a short distance.

In this study on crystalline/crystalline PET interface, the situation is similar to heat sealing at a temperature lower than T_m . However, the contact time is much longer than the common dwell time of heat seal adhesion. In spite of that, the low diffusion at temperatures lower than T_m was also reported in heat seal studies as mentioned earlier.

Xue et al. (1998) studied the welding of ultra high molecular weight polyethylene (UHMWPE) using T-peel test above and below the melting point. They prepared UHMWPE films with two different methods: melt crystallized films and dilute solution crystallized films. Due to the ability of UHMWPE to crystallize in a wide variety of morphologies, these two methods created different morphology of crystals. After welding above T_m (145°C) for 3-65 minutes, the adhesive fracture energy was measured at both room temperature and at 135°C. It was found that, irrespective of the initial morphology, the adhesive fracture energy measured at 20°C was comparable to the fracture energy of the bulk material after a contact time shorter than 3 minutes. However, when fracture adhesive energy was measured at 135°C, the initial crystalline morphology affected fracture energy. In the case of films prepared from solution, the increase of adhesive energy was almost instantaneous. The solution-crystallized films exhibited the so-called “chain explosion”, a very fast increase of the radius of gyration upon melting. The authors

suggested that for UHMWPE the crystallization from an extremely dilute solution results in “single crystal mats”, in which entire polymer chains are confined to one crystal that caused a large reduction of the radius of gyration. Then, the chains rapidly recovered their R_g by heating to T_m . They suggested that this phenomenon caused the instantaneous strengthening of interface in case of pre-wetted solution-crystallized films (Xue et al., 1998). In contrast to solution-crystallized films, for melt crystallized films, the peel force gradually increased with contact time and even after 65 hours of welding, it could not attain the fracture energy of the bulk material.

In the second part of their work (Xue et al., 2000), they studied the effect of cocrystallization at the interface on self-adhesion of UHMWPE. They used solution-cast films containing regularly stacked lamellae, which exactly doubled in thickness upon annealing for 15 min at 125°C. (See Figure 2-21) In amorphous polymers, the strengthening of the interface is due to the formation of entanglements upon diffusion. The deep chain diffusion, and consequently long welding times, is required for good adhesion. The results of peel force in co-crystallized specimens showed that, for semicrystalline polymers, large-scale chain diffusion is not a prerequisite for good welding performance. The strength at the interface could be developed by cocrystallization at relatively low welding time (Xue et al., 2000).

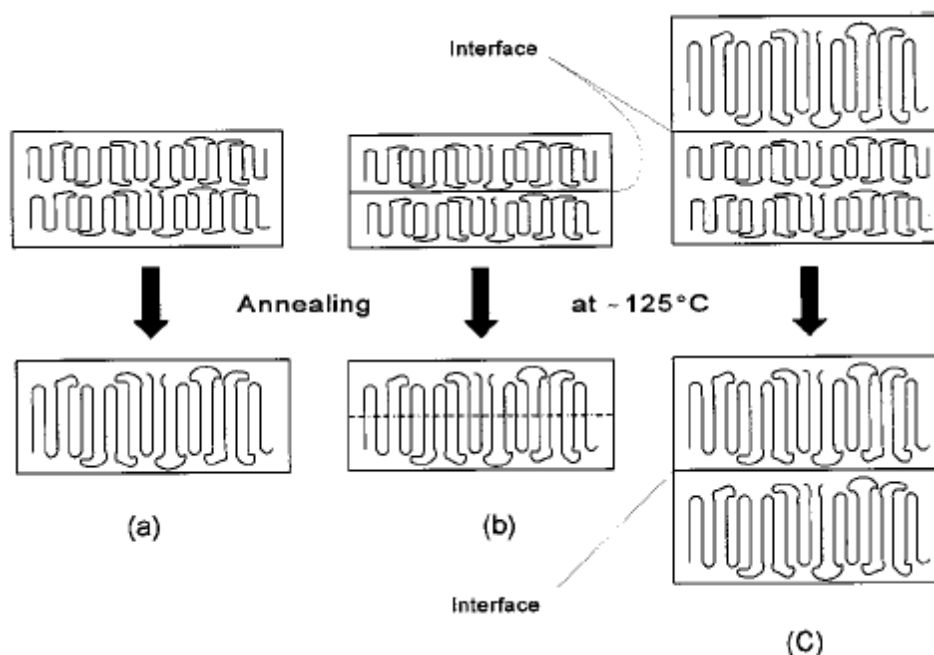


Figure 2-21 Schematic of the study plan by Xue et al. for investigation of cocrystallization at interface; (a) the lamellar doubling upon annealing at 125°C , (b) lamellar doubling by cocrystallization across the interface, and (c) pre-annealing one of the films prohibited cocrystallization across the interface. (Xue et al., 2000)

Cocrystallization was also studied in the welding between cross-linked high density polyethylene sheets. The very high adhesion force was interpreted to be the consequence of cocrystallization between polyethylene chains across the interface (Gent et al., 1997). The key role of cocrystallization was also illustrated by Smith et al. on welding of polypropylene in a range of temperatures from below to above T_m (Smith et al., 2001).

The literature review in this section highlighted the fact that the number and diversity of studies on interfacial adhesion of semicrystalline polymers are very limited. They are specifically limited to the welding process in which the contact time is commonly higher than few minutes. While the heat seal process is performed in either 1-2 seconds or a fraction of a second.

In summary, a few published studies in literature have investigated the effect of process parameters. However, an overall analysis on the control and optimization of process parameters is missing.

As mentioned earlier, in the work of Meka and Stehling (1994) the analysis was based on the interface temperature which is not practically possible to measure in industrial process. They used a finite elements model to obtain the temperature at interface and they showed that the interface temperature will reach to the jaw temperature after 0.3-0.5 s. In the industrial process however, the heat sealing occur in a non-isothermal condition and temperature setting is based on the control temperature on the heated jaws. They also introduced the heat sealing curve but did not discuss the trend of seal strength, plateau initiation temperature, and plateau broadness with regard to dwell time and pressure (Stehling and Meka, 1994).

Moreover, the seal strength of several polymers has been studied but there is not an overall fundamental investigation on seal strength of a resin at different process conditions (Mueller et al., 1998; Poisson et al., 2006a). Also, the analysis of the amorphous fraction of polyethylene resins has been done only in one process condition, where the amorphous fraction might be different for every process condition.

In studies on heat seal process parameters, the effect of pressure or dwell time were mostly discussed at one temperature. So the concurrent effects of dwell time-temperature and pressure-temperature have not been investigated before.

In addition, the models connecting the reptation parameters to the final interface strength has been verified in welding process where the contact time is much higher than the contact time of heat sealing in flexible packaging applications. So it needs to be verified in dwell times less than a second as the realistic dwell time.

A good hot tack performance and the influential parameters on it is still a major challenge. The reports on hot tack are limited to a few conference papers and patents comparing the hot tack performance of different polymers. To our best knowledge there is no study about the effect of molecular structure on hot tack properties. Also there is lack of fundamental knowledge about parameters affecting hot tack performances such as hot tack strength and also hot tack plateau broadness.

2.3 Originality of the work

According to the literature review above, the heat sealing process has not been well explored. There is a lack of knowledge about how the material structure or process parameters affect heat

seal quality. These are required to design optimal processing conditions, in order to maximise “seal performance” and minimise costs. Therefore, a comprehensive study is required to clarify the role of controlling factors of heat seal process, including process parameters and material characteristics.

The analysis of polymer adhesion from the microscopic point of view based on diffusion theories has been mainly used in polymer welding studies, where generally ideal condition of extended time and constant temperature exist at the interface. To our knowledge this approach has not been used in film heat sealing because it is a non-isothermal process and is usually performed in a fraction of a second. Then, a clear vision of the role of different molecular structures on heat sealing is still far beyond reach. The majority of existing studies on self adhesion of either semicrystalline or glassy polymers have been focused on measuring the interfacial adhesion after cooling. Also the focus has been mostly on amorphous polymers. In literature, very little has been reported on hot tack or interfacial self-adhesion of polymer films at the same temperatures as those of heat bonding.

2.4 Objectives of the current study

The main objective of this study is:

“To control and optimize the seal, in terms of material and process variables, for multilayer packaging films”

To achieve this main objective, these specific objectives were designated:

- The optimization of heat seal process parameters i.e. temperature, pressure, and dwell time by performing an extensive set of sealing experiments and using microscopic techniques to investigate seal microstructure
- To establish relationships between the molecular architecture of polyethylenes based sealant material and the final seal quality.

CHAPTER 3

ORGANIZATION OF ARTICLES

The main achievements of this research project are presented in the form of three scientific papers in the following three chapters:

Chapter 4 presents the results of the first paper: “*A novel approach toward the effect of seal process parameters on final seal strength and microstructure of LLDPE*” accepted to be published in the *Journal of adhesion science and technology*. In this work, we investigated the different active mechanisms in the heat seal process of a semicrystalline polymer. We have chosen to work on monolayer LLDPE film. The atomic force microscopy technique and the comparative roughness analysis were used to obtain information on the seal microstructure molecular interdiffusion.

Chapter 5 presents the results of the second paper: “*Role of Molecular Architecture in Interfacial Self-adhesion of Polyethylene Films*” submitted to *Macromolecules*. In this work, we investigated the role of molecular architecture of a polymer chain on final hot tack strength. The investigated molecular structures include: molecular weight (Mw), molecular weight distribution (MWD), amount and distribution of long chain branch (LCB), and short chain branch (SCB) distribution among and along polyethylene chains. A variety of polyethylene resins with different chain structures were utilized in this part.

Chapter 6 presents the results of the third paper: “*Interfacial Self-Adhesion of Polyethylene Blends: the Role of Long Chain Branching and Extensional Rheology*” submitted to *Rheologica Acta*. In this work, we focused on binary blends of two main categories of polyethylene, metallocene catalyzed and conventional polyethylene resins. Through different blend compositions, we studied the influence of blending of resins with different chain structure on hot tack properties. The correlation of melt elongational rheological properties with final hot tack strength was discussed in this part of the work.

CHAPTER 4

ARTICLE 1: A NOVEL APPROACH TOWARD THE EFFECT OF SEAL PROCESS PARAMETERS ON FINAL SEAL STRENGTH AND MICROSTRUCTURE OF LLDPE*

Zahra Najarzadeh, Abdellah Ajji

4.1 Abstract

The optimization of heat sealing process parameters, including time, temperature, and pressure, was performed on a monolayer linear low density polyethylene (LLDPE) film. The seal properties examined for each process condition were: Seal initiation temperature (T_{si}), plateau initiation temperature (T_{pi}), final plateau temperature (T_{pf}), plateau seal strength (SSp) and failure mode. Increasing dwell time enhanced seal strength. However, it was found that the rate of this enhancement is different for each interval of dwell time. In term of temperature window broadness at each specific dwell time, a narrow temperature plateau was observed for dwell times lower than 0.4 s and higher than 2 s. While in between a broad temperature window was observed. The pressure shows its influence up to the stage of wetting. And after providing the intimate contact between two film layers, additional increase in pressure does not enhance seal strength significantly. At high pressures, seal curves showed a narrower temperature window. Moreover, it was shown that the level of plateau seal strength in LLDPE changes slightly with process conditions. A 3D mapping of process safety zone was introduced for seal strength in the range of heat seal process variables for the very first time. The analysis of this 3D representation revealed that seal strength has a linear correlation with the square root of dwell time. In addition, the interfacial bond strength was shown to be proportional to the fraction of melted crystals. It was found that this fraction is determined by dwell time and temperature. Topography and morphology of surfaces after peeling revealed enlargement of fibrillar morphology to taller

* Accepted to publish in Journal of Adhesion Science and Technology, March 2014.

failure fracture complex shapes. Extensive roughness analysis on film surfaces after peeling found the much rougher surfaces after breakage of strong bonding.

4.2 Introduction

Heat sealing is the process of joining thermoplastic materials and is the most popular method for bonding two polymer films in package closures. It is performed by direct contact of films with heated bars and applying sufficient pressure. Factors dictating the ultimate performance of heat sealed films could be categorized as process related and film related. Process related factors are heat seal process conditions: the applied temperature, dwell time, and pressure. Previous studies have confirmed that seal strength depends mainly on temperature and dwell time, and less on pressure¹⁻⁴. However, the trend of this influence in various time intervals is not straightforward and has not been discussed yet.

The general heat sealing curve introduced by Meka and Stehling² describes seal temperature dependence of seal strength in semicrystalline polymers at a constant dwell time and pressure. Figure 4-1 shows the schematic of this curve and its important features: Seal initiation temperature (T_{si}) is the lowest temperature in which a seal is created with a minimum level of seal strength. After a few degrees increase in temperature from T_{si} , a sharp increase in seal strength occurs and stays almost constant for a temperature interval afterward. Plateau initiation temperature (T_{pi}) is the temperature in which the plateau seal strength begins, and the final plateau temperature (T_{pf}) is the terminal temperature of plateau.

The ability to design optimum processing conditions to maximise “seal performance” while minimising costs is certainly desirable. The term “Seal performance” encompass not only consistency of film’s joints but time and temperature dependence of seal strength, broadness of the temperature range which produce the highest seal strength, seal initiation temperature, seal failure mode, and seal uniformity. In high speed industrial sealing lines, fluctuations in temperature of heated bars require obtaining high seal strength in a broad range of temperatures. This is where the broadness of the plateau is required. Depending on the material and process, the temperature window of plateau seal strength varies from a few degrees to 15°C or more.

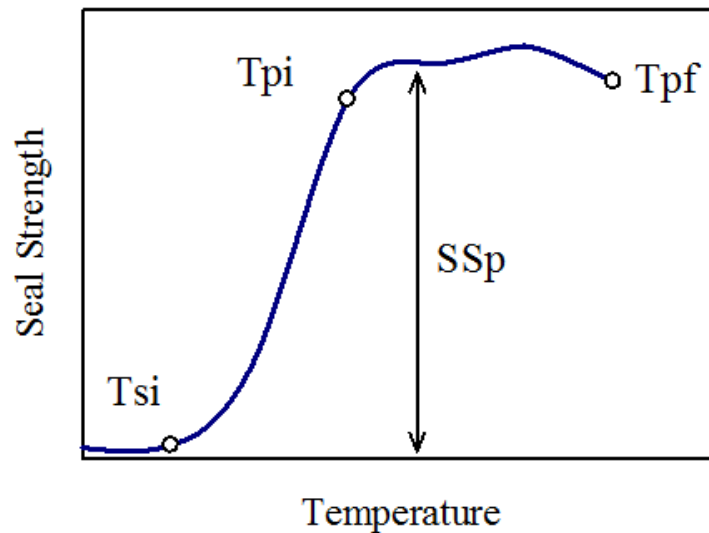


Figure 4-1 Heat seal curve for semicrystalline polymers introduced by Meka and Stehling ¹; T_{si} : Seal initiation temperature, T_{pi} : Plateau initiation temperature, T_{pf} : Final plateau temperature, SS_p : Plateau seal strength, Seal strength plateau is the interval of T_{pi} and T_{pf} ;

In the vertical and horizontal form fill seal (VFFS and HFFS) modern machines are generally required to operate at high production speed. The balance of sacrifice in either production speed or energy would be determined by optimization of process condition. Stehling and Meka ¹ introduced a model for the heat sealing mechanism at the molecular scale in semicrystalline polymer films. Figure 4-2 represents a schematic of it: crystal melting by applied heat, interdiffusion of polymer chains across the interface, making entanglements, and re-crystallization upon cooling. Based on this mechanism, the final seal performance depends also on crystallinity and polymer chain ability to interdiffuse as material characteristics.

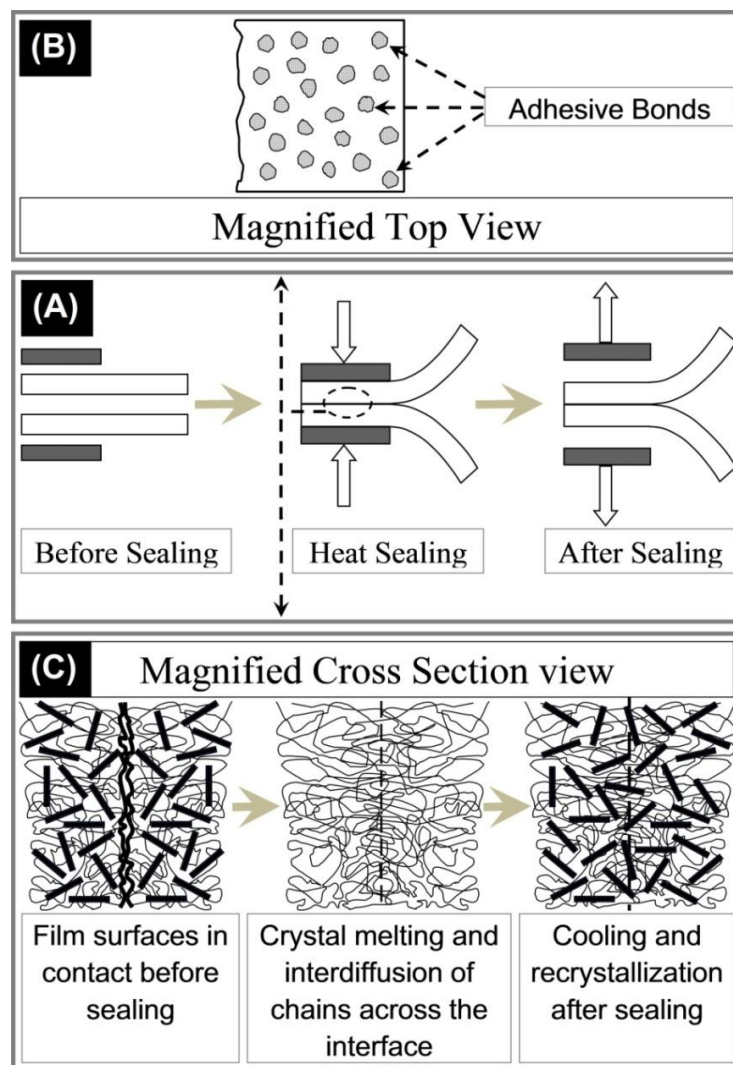


Figure 4-2 Heat sealing mechanism: (A) Heat sealing process steps, (B) Magnified top view of sealed area, (C) Magnified cross section view of sealed area showing the seal mechanism in molecular scale

In literature, there are two main stream approaches about heat bonding analysis of polymer interfaces. First, the analysis of polymer chain behaviour from microscopic point of view based on reptation theory. This is mainly used in polymer welding studies where ideal condition of extended time and constant temperature exist at the interface⁵⁻¹¹. However, to our knowledge this approach has not been used in heat sealing. Because film heat sealing is a non-isothermal process, and this process is usually performed in a fraction of a second which is a non-ideal condition.

Second approach is the conventional macroscopic analysis. This method empirically investigates the seal strength based on bulk properties of the polymer. Unlike the former, this approach does not consider chain motion and polymer chains interdiffusion. Through this approach, the macroscopic analysis, including mechanical and thermal properties, has been performed on some common sealant materials ^{1-3,12-16}. Several studies have been carried out on sealing properties of different materials and film structures. In a study on ionomers, Morris et al. ¹⁷ investigated the influence of thickness on the relation between applied temperature and dwell time. Also, by the use of impulse heat sealing on two laminated layers of oriented and cast polypropylene, the effect of orientation in machine and transverse direction on seal strength was studied ^{14,18}. Unfortunately there are not several previous studies on such a technically important topic and on LLDPE as the most widely used polymer in seal layer applications.

This study attempts to address both approaches by performing an extensive set of sealing experiments on a conventional LLDPE and using microscopic techniques to investigate seal microstructure.

Strength development at polymer-polymer interfaces was described by de Gennes, ^{5,19,20} Prager and Tirrell ²¹, and by Wool ²² through polymer chain diffusion based on reptation motion theory ⁷. They illustrated power law dependence of chain displacements to molecular weight and time. For ideal welding condition, the argument of non-Fickian diffusion was utilized to generate empirical equations for the number of chains crossing the interface and the average diffusion distance. These arguments have been utilized in researches on variety of subjects of polymer interfaces such as welding, adhesives, emulsions, polymer blends, etc ²³⁻²⁹.

In this work, seal experiments are conducted in order to optimize the sealing parameters and to perceive the effect of concurrent change of “temperature - dwell time” and “temperature - pressure” on seal strength. The atomic force microscopy technique was used to determine surface roughness of the seal interface after peeling in order to obtain information on the seal microstructure. The comparative roughness was utilized as representative of the level of film surfaces’ contact area and molecular interdiffusion.

4.3 Materials and methods

4.3.1 Film production

A hexane based ethylene α -olefin copolymer, produced with a conventional Ziegler-Natta catalyst for seal applications, was supplied by ExxonMobil Chemicals in the form of granules with MFI=2 g/10min and density 0.918 g.cm⁻³. Monolayer films were produced using a laboratory scale cast line comprised of a 45 mm Killion single screw extruder and an 8 inch cast film die. Cooling was performed using an air knife and calendar. The films were produced at constant throughput without slip or antiblock additives with a nominal film thickness of 2 mils (50 μ).

4.3.2 Differential scanning calorimetry (DSC)

Thermal analysis was performed using a TA Instruments Q1000 DSC with 10°C/min heating rate in first heating cycle. The melting temperature of the LLDPE film was determined as 120°C. Heat of fusion of a theoretically 100% crystalline polyethylene, which is 290 Jg⁻¹, was used to determine the crystalline and amorphous fractions of film³⁰. The final melting temperature (T_{mf}) was also considered as the temperature where all the crystals are melted.

4.3.3 Heat seal measurements

Heat seal experiments were performed on a SL10 LakoTool laboratory hot-tack and seal tester purchased from “Lako Tool & Manufacturing Inc.” at a determined temperature, pressure, dwell time and peel rate. Heat seals were made using flat seal bars covered with a layer of Teflon coating. Seal samples were cut in 2.54 cm×33 cm strips specimens parallel to the machine direction and a BOPP tape was used as the back-layer to protect the seal films from fusion and sticking to the hot seal bars. Seal strength experiments were performed based on ASTM F2029 and ASTM F88. T-peel tests were done at a constant peel rate (3.3 mm.s⁻¹) and peel strength was taken as the average plateau peel strength of 5 specimens.

4.3.4 Atomic force microscopy (AFM)

Escope AFM was operated in the tapping mode at room temperature using nanosensor tapping etched silicon probes (TESP) with single beam cantilevers. The amplitudes of the drive signal

used to set the cantilever oscillation were in the range between 2.8 and 4.2 V. The samples were attached to the AFM magnetic disk sample holders using double-sided tape. Height and phase images were collected simultaneously. Surface topography was performed before and after seal and peeling at the scan rate of 1 Hz. They were processed using NanoScope software by flattening to remove background shapes. Then the surface roughness parameters of height images were calculated.

4.3.5 Scanning electron microscopy (SEM)

In order to investigate the morphology of surfaces SEM was used. The surface of original films before adhesion and the peeled surfaces after sealing and the subsequent T-peel test were examined. They were gold sputtered and observed under a HITACHI S-4700 SEM.

4.4 Results and Discussion

4.4.1 Effect of Heat seal process parameters

Time and temperature dependence of seal strength is presented in a 3D graph shown in Figure 4-3. To see the effect of dwell time, the applied pressure was kept constant at 0.5 Nmm^{-2} and dwell time was varied from 0.1-3 s. In the interval of very low dwell times (0-0.5 s) sealing was done at every 0.1 s while for dwell times higher than 0.5 s it was done in steps of 0.5 s. In this 3D graph, the level of seal strength at each single point is represented by a color. Considering red/orange area as adequate seal strength (above 2000 g/25.4mm), at very low dwell time (lower than 0.4 s), a narrow plateau can be seen. While for higher dwell times, the orange area covers a wider range of temperature meaning a broader plateau, and become narrower again at very high dwell time. The broadest plateau is obtained in the range of 0.5-1.5 s. The lowest dwell time with broadest plateau is 0.5 s.

The border of the lower blue area represents T_{si} . This border shows that an increase in dwell time decreases T_{si} . Similarly the breakage zone is the area for which the time-temperature combinations leads to seal distortion due to very high sealing temperature or long dwell time. This seal distortion means a brittle seal area with the possibility of defects which may break the seal edge and threaten the integrity of seal and package.

The trend of T_{pi} , located in the area inside the green border (the yellow and orange area), represents seal strength higher than 1500 or 2000 (g/25.4mm). Based on this criterion for seal strength the parameter setting inside this area practically results in adequate seal strength. So, the user can decide between either time or temperature. This boundary reveals the optimum “production rate-energy” condition which indicates the lowest temperature requirement for each dwell time to obtain satisfactory seal strength.

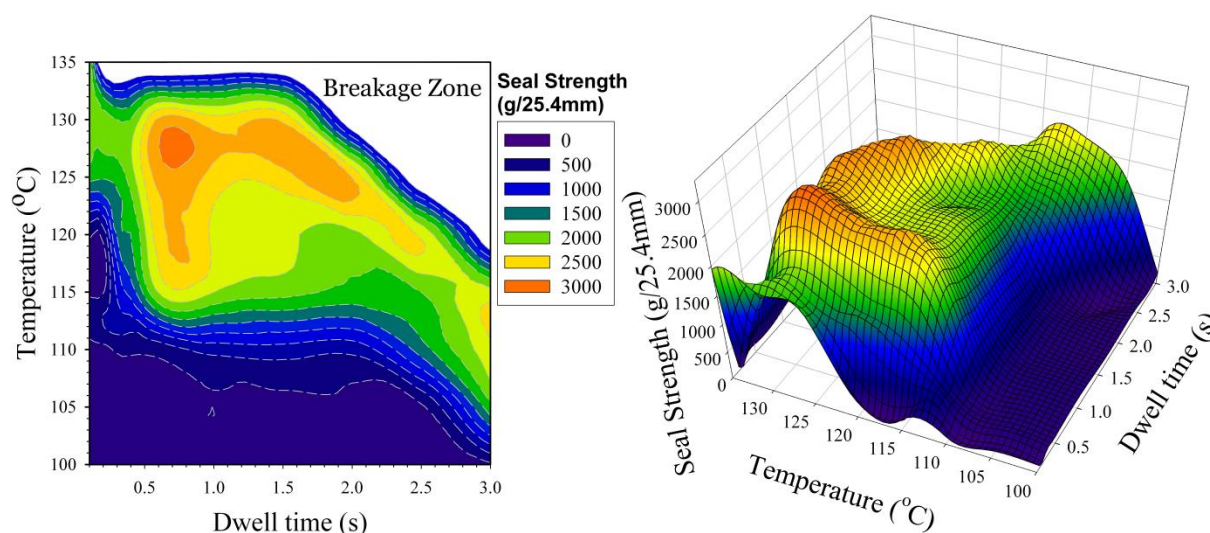


Figure 4-3 Dwell time-temperature dependence of seal strength in 3D and contour plots

This strong time and temperature dependence of interfacial strength was reported by other researchers as well ^{3,4,18,31,32}. Dwell time dependence of seal strength can be explained based on the higher amount of heat that reaches film interfaces at longer time and gradual change of film's surface from crystalline to partially and then fully melted surface. As mentioned earlier, diffusion of chains across the interface and entanglement are required to create a strong seal. The number of chains capable of diffusion is determined by the fraction of amorphous phase. Also the unmelted crystals act like obstacles hindering chains' free diffusion ³¹.

To evaluate the effect of pressure, sealing was performed at a constant dwell time of 0.5 s for pressures of 0.1, 0.5, 1, 1.5, 2, and 3 N.mm⁻². (See Figure 4-4) Again, considering red/orange color area as adequate seal strength (above 2000 g/25.4mm) the seal curves show narrow plateau for low pressures (0.1 N.mm⁻²). It becomes broader at higher pressures and remains almost constant by further increase of pressure. It then becomes narrow again for very high pressures.

The trend observed for low pressures is because of reduced contact at film interfaces, so even if temperature goes up and melt the crystals, it cannot provide enough interdiffusion. In addition, at very high temperatures, the film surfaces are totally molten and there is a possibility of shear flow under pressure which cause seal distortion and results in a decrease of seal strength. By increasing pressure from 0.1 to 0.5 N.mm⁻², the plateau initiation temperature (T_{pi}) shifts to lower temperatures. However, it remains constant after that. Thus, the effect of pressure is not as significant as the effect of dwell time as long as the films are in sufficient contact. Although some previous researchers³ doubted the possibility of producing a good seal at pressure as low as 1 bar (= 0.1 N.mm⁻²), adjusting temperature allowed us to obtain a plateau of high seal strength.

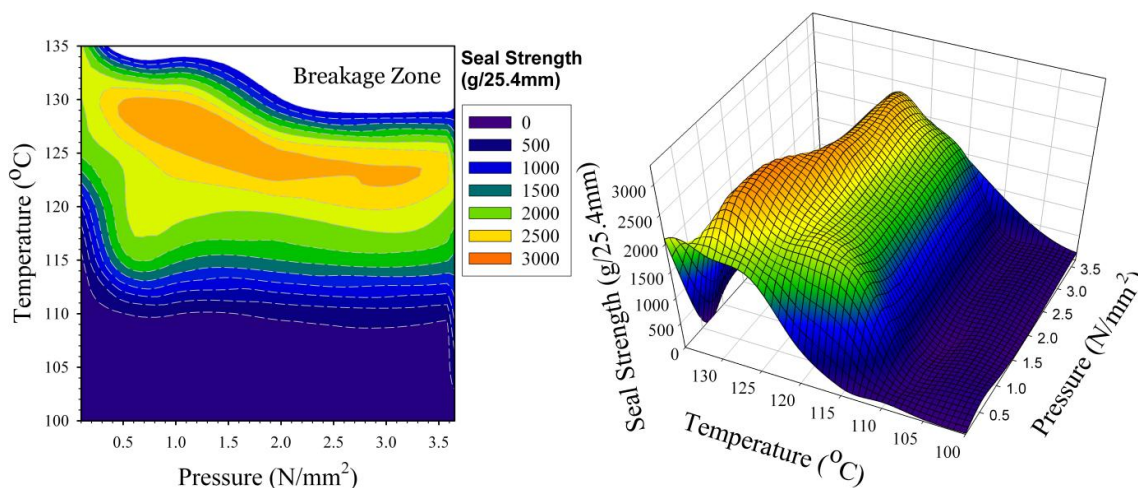


Figure 4-4 Pressure-temperature dependence of seal strength in 3D and contour plots

Compared to the effect of dwell time on seal strength, pressure influences seal through an entirely different mechanism. The role of pressure is to provide intimate contact at interface. Establishment of this contact at the molecular scale is known as “wetting” at interface. Wetting is sensitive to surface topography and hydrostatic pressure^{10,33}. With an increase in pressure, wetting is enhanced at interface and provides improved contact for polymer chains at interface. The next step after wetting is chain diffusion by reptation motion. Generally high hydrostatic pressure decreases free volume, thus retards segmental motion and hence reduces the diffusion coefficient. This effect is not prominent for hydrostatic pressures lower than 100 N.mm⁻². Consequently, in seal process, pressure would not influence significantly chains’ interdiffusion²².

Then pressure has no significant effect on interdiffusion stage which controls the development of mechanical properties during healing.

It is concluded that seal strength is dependent on pressure to some extent, because pressure promotes the wetting step by making better contact. However, it is independent of pressure afterward in interdiffusion stage.

4.4.2 Amorphous fraction

In the previous section, through discussion on seal strength as a function of sealing temperature, it was argued that seal strength strongly depends on the amount of available amorphous phase at the interface. The plot of weight fraction of amorphous phase as a function of temperature ($f_a(T)$) is known as melting distribution². For a semi-crystalline polymer, the final melting point (T_{mf}) of a film is the temperature where $f_a(T)=1$.

$$f_a(T) = 1 - \left(\frac{\Delta H_S}{\Delta H_U} \right) + \left(\frac{\Delta H_T}{\Delta H_U} \right)$$

Where ΔH_S is Heat of fusion of sample, ΔH_U is Heat of fusion for a theoretically 100% crystalline polyethylene, which is 290 Jg^{-1} ,³⁰ and ΔH_T is the Cumulative heat of fusion at temperature T .

Figure 4-5 represents the melting distribution curve of LLDPE and Figure 4-6 shows the part of melting distribution curve corresponding $T_{si} - T_{pi}$ interval for each dwell time. Figure 4-6 explains how the seal strength plateau correlates with amorphous fraction. At lowest dwell time (0.1 s) the amorphous fraction is 0.93 to 1, for $T_{si}=120^\circ\text{C}$ and $T_{pi}=125^\circ\text{C}$, indicating that this high amorphous portion of material is necessary to provide adequate chain mobility to make good seal in this short time. Whereas for dwell time= 0.5 s the amorphous fraction at $T_{si}=110^\circ\text{C}$ is 0.82 and at $T_{pi}=115^\circ\text{C}$ is 0.87, and for dwell time= 1 s the amorphous fraction at $T_{si}=106^\circ\text{C}$ is 0.83 and at $T_{pi}=112^\circ\text{C}$ is 0.80. It can be concluded that at higher dwell times (0.5, 1, 2 s), lower amount of amorphous phase is adequate to cross the interface, make entanglements and consequently create a good seal. This argument support the interrelated effect of time and temperature as mentioned earlier.

Practically it is important to be able to predict T_{pi} and T_{si} of sealant by knowing its melting point. In literature, there is disagreement about the relation of T_{pi} and T_{si} with melting point for different materials. Stehling and Meka reported that the fraction of amorphous phase is almost constant

(equal to $77\pm3\%$) at the seal initiation temperature and the T_{pi} value corresponds closely to T_{mf} for all the unsupported polyethylene films they studied^{1,2}. For Oriented PP/Cast PP films, Tetsuya et al. and Yuan et al. obtained T_{pi} lower than T_{mf} while Morris reported T_{pi} higher than T_{mf} for ionomers^{3,17,18}. Our results shows that the T_{pi} - T_{mf} relationship is strongly influenced by the heat seal process parameters, dwell time and pressure. This effect explains the mentioned conflict that in previous studies sealing in various conditions resulted to various values. It also depends on the microstructure, mobility of polymer chains and their functionality. This subject will be further discussed in our future works.

It should be mentioned that the amorphous fraction in this work and other studies in literature have been obtained using DSC melting thermographs at the heating rate of $10^{\circ}\text{C}/\text{min}$. However, the heating rate in the heat sealing process is two orders of magnitude higher than that, which is not achievable in conventional DSC equipments. The amorphous fraction might be influenced by the heating rate. Then using DSC to obtain amorphous fraction could only assist estimation of this parameter during the heat seal process.

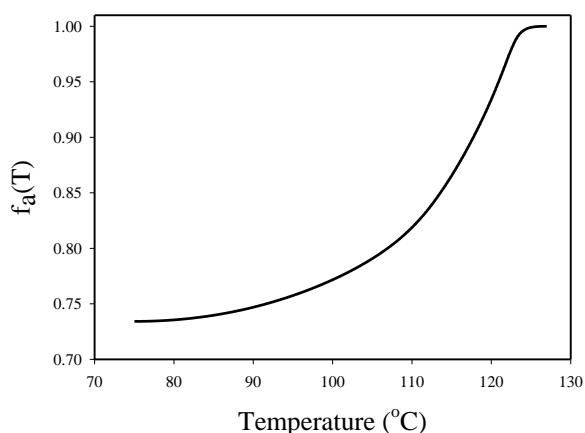


Figure 4-5 Melting distribution of LLDPE film as a function of temperature considering $T_{mf}=125^{\circ}\text{C}$

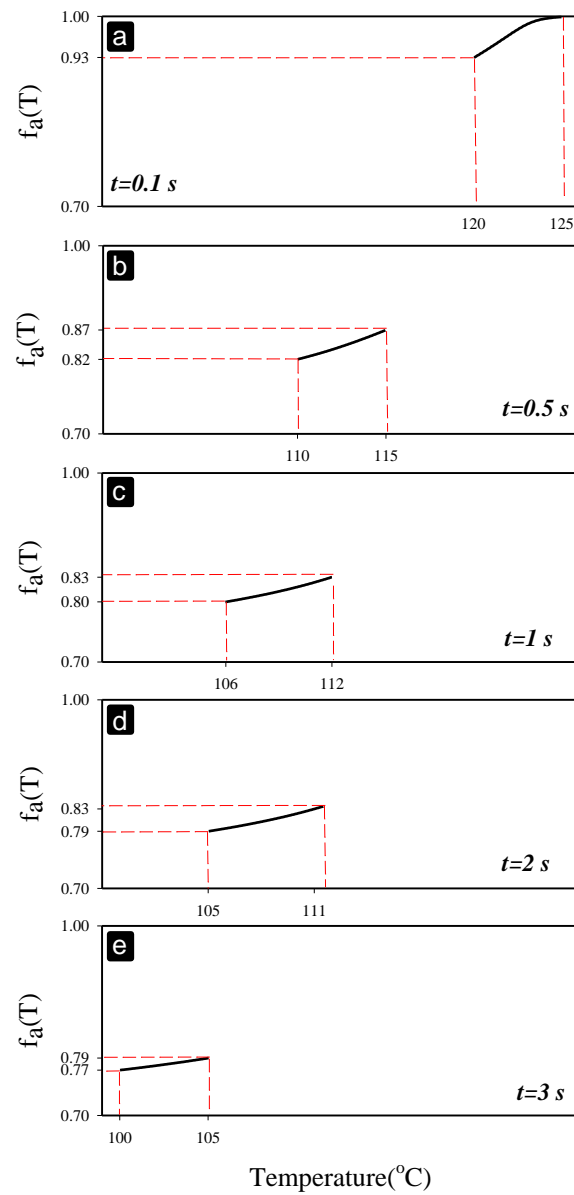


Figure 4-6 Melting distribution of LLDPE as a function of temperature, the corresponding amorphous fraction at T_{si} and T_{pi} are indicated at: (a) dwell time=0.1 s, (b) dwell time=0.5 s, (c) dwell time=1 s, (d) dwell time=2 s, (e) dwell time=3 s

4.4.3 Time dependence of seal strength

The development of a heat seal at the microscopic scale involves joining two polymer surfaces by the action of time, temperature and pressure. To create a weld that is indistinguishable from bulk,

the polymer chains need to diffuse across the interface. The temperature needs to be high enough for free chain ends to have reptation motion in process time.

As discussed by de Gennes⁵ in reptation theory, chain ends located at the interface are going to relinquish their initial tube, cross the interface, and create junctions at the other side. With time, the number and the length of interdiffused chains increase. Over time the chain ends could diffuse deeper to the other polymer layer. This part of chain ends which escaped from their initial tube is called “minor chain”. The lengths of minor chains increase by time until they are long enough to make entanglements. Therefore, the number of entanglements also increases over time. At a critical time, known as saturation time (τ), the interface is going to disappear and the properties of bulk material would be reached. This is the time in which the maximum interface strength is reached²².

In order to relate microscopic properties of the interface to macroscopic properties of polymer bulk; the healing theory was introduced³⁴:

$$S_{(t)} = f(H_{(t)})$$

In which $S_{(t)}$ represents interface strength (as macroscopic property) and $H_{(t)}$ represents the microscopic property (such as minor chain length). Both $H_{(t)}$ and $S_{(t)}$ are time dependent, thus they are known as dynamic terms of healing. An increase in macroscopic strength of interface ($H_{(t)}$) is a function of increase in chain interdiffusion ($S_{(t)}$). In a fully healed interface which has the strength of bulk material, S_{∞} and H_{∞} were introduced. They are defined as the equilibrium values of $S_{(t)}$ and $H_{(t)}$. S_{∞} and H_{∞} are known as static terms of healing. S_{∞} represents minor chain length (or any other microscopic property at interface) after $t = \tau$ when the chains at interface are at the equilibrium state of bulk material, and the gap at interface is disappeared^{22,34}.

At polymer-polymer interface the scaling law^{22,35} was introduced to show the relation of dynamic ($H_{(t)}$) and static (H_{∞}) terms of microscopic property in healing:

$$H_{(t)} = H_{\infty} \left(\frac{t}{\tau} \right)^{r/4}$$

Which τ is saturation time (as mentioned earlier), and $r=1, 2, 3 \dots$ is a power law factor. The value of r is determined by the type of microscopic property (H). Assuming that the

interpenetration chain length, as H , plays a major role in controlling the strength of polymer interface it was shown that $r=2$ ³⁴.

In this work the peel rate is low then disentanglement is dominant failure in peeling and, as mentioned earlier, the minor chain length plays the major role in the time dependence of overall strength development²². Using the scaling law, the time dependence of interface strength would be:

$$S(t) = S_{\infty} \left(\frac{t}{\tau} \right)^{1/2}$$

According to this analysis, the strength of jointed film surfaces would have a linear relation with square root of time. The scaling law presented, offered a convenient framework for researchers evaluating molecular behaviour in terms of the static and dynamic properties of the polymer chains²². This model was established by several researchers theoretically^{7,10,25} and experimentally^{7,34}.

The interface strength power dependence of $t^{1/4}$ was also reported in some cases of healing in polymer interfaces. However it is reported to be applicable in case of the segregation of chain ends at the surface, which could occur due to prior fracture of surfaces^{7,26,36,37}.

Figure 4-7 shows the $t^{1/2}$ dependence of seal strength before complete healing of interface. As it can be seen in the graph, seal strength has linear dependence to $t^{1/2}$. It confirms the linear prediction of healing theory. The increase in the slope of the square root dependence of seal strength depicts the enhancement in seal strength's time dependency by increasing temperature. In Figure 4-7, the last point of seal strength at any temperature represents the point the maximum strength of interface is reached beyond that.

In the healing theory discussed above, the presence of crystals was neglected and surfaces in contact were considered to be amorphous. In semicrystalline polymers, the bridges across the interface could participate in crystals on either sides of the interface. Consequently, in addition to entanglement, these minor chains act as tie molecules and the small crystals created upon cooling act as anchors for the tie. Even though, the linear dependence of LLDPE seal strength to $t^{1/2}$ in this work is in agreement with the healing theory.

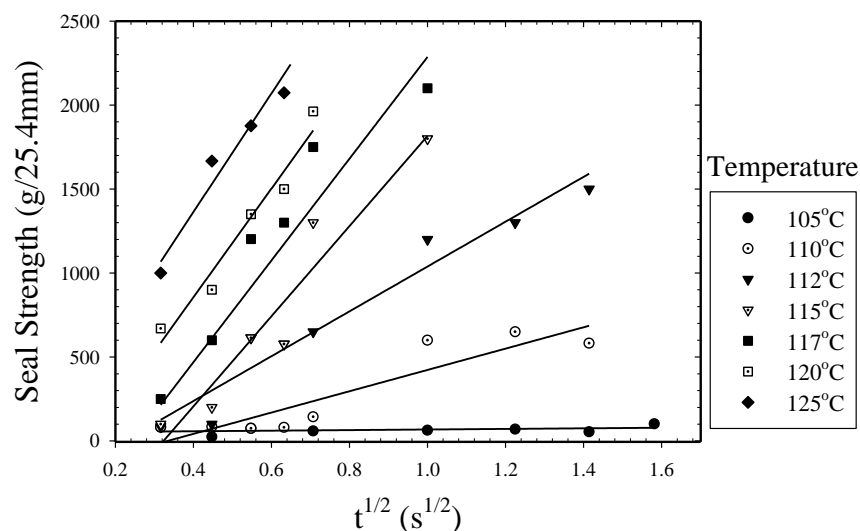


Figure 4-7 Seal strength at different temperatures versus (dwell time)^{1/2} Linear regression's line slope= a (g/25.4mm.t^{0.5}); 105°C: $a=17.5$; 110°C: $a=634.8$; 112°C: $a=1334.8$; 115°C: $a=2684.3$; 117°C: $a=3029.8$; 120°C: $a=3234.8$; 125°C: $a=3534.4$;

4.4.4 Failure mode analysis

In failure mode analysis, three temperature zones were observed, representing failure mode categories. As illustrated in Figure 4-8, peeling failure zone is the region in which the seal area peels apart at the film interface because of a limited chain interdiffusion and molecular entanglement across the interfacial zone. Plateau zone, in which delamination and elongation failure occur, is the range of elevated temperatures capable of melting crystals providing adequate number of chains available for diffusion and entanglements such that the seal area is well-built. Breakage zone is where high temperature, dwell time, or pressure cause merging of film surfaces which make adhesive debonding impossible. In these circumstances, the strength of the sealed interface exceeds the film strength at seal edge which, along with distortion in seal area, causes the arms to neck or break. In general the upper limit of the measurable seal strength locates at the onset of this zone.

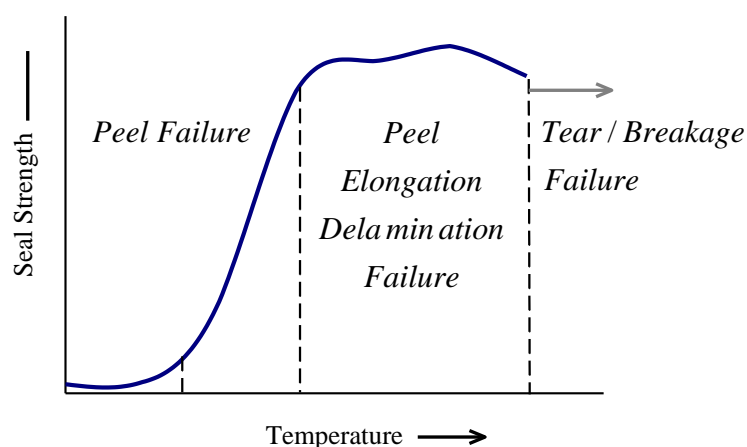


Figure 4-8 Representation of failure modes observed in seal curve of LLDPE films supported with BOPP tape

4.4.5 Topographic and morphological analysis by AFM and SEM

In the debonding failure mode, the crack propagates through the sealed interface and it leaves numerous fractured craze fibrils on the peeled surfaces. Depictive AFM 3D height micrographs of virgin film and peeled surfaces are shown in Figure 4-9. To capture all topographic aspects and tiny features of the examined surfaces in 3D, height imaging of 5 μ m scan sizes was found the most efficient.

The origin of these fibrils is the bridges formed during sealing by diffusion of chains across the interface and entanglements they made. These bridges constructed from moveable amorphous parts of polymer chains act as stitches attaching two film layers. Fibrils density and length depend on the depth of diffusion and number of chains available for motion at every definite sealing condition. At sealing temperatures below T_{pi} (115°C) the chain motion is more confined and, consequently, fewer bridges cause lower peel strength and leave smaller fibrils at peeled surface. By increasing temperature, the broken stitches are stronger thus the fibrils on the peeled surface are noticeably longer and thicker.

In peeled surfaces from sealing at temperatures above T_{pi} , a network of connected larger peaks and valleys is more prominent than isolated fractured thin fibrils. This shows a better adhesion in higher temperatures. Hiltner et al. have reported similar connections between fibrils as sealing time was increased to 1000-3500 s in welding of LLDPE films. They called it a three-

dimensional cellular structure. The alteration in fractal morphology of peeled surfaces suggests the presence of larger number of stitches and broader bridges across the two surfaces. In addition to this visual conclusion from AFM 3D height images, this change was monitored through roughness parameters.

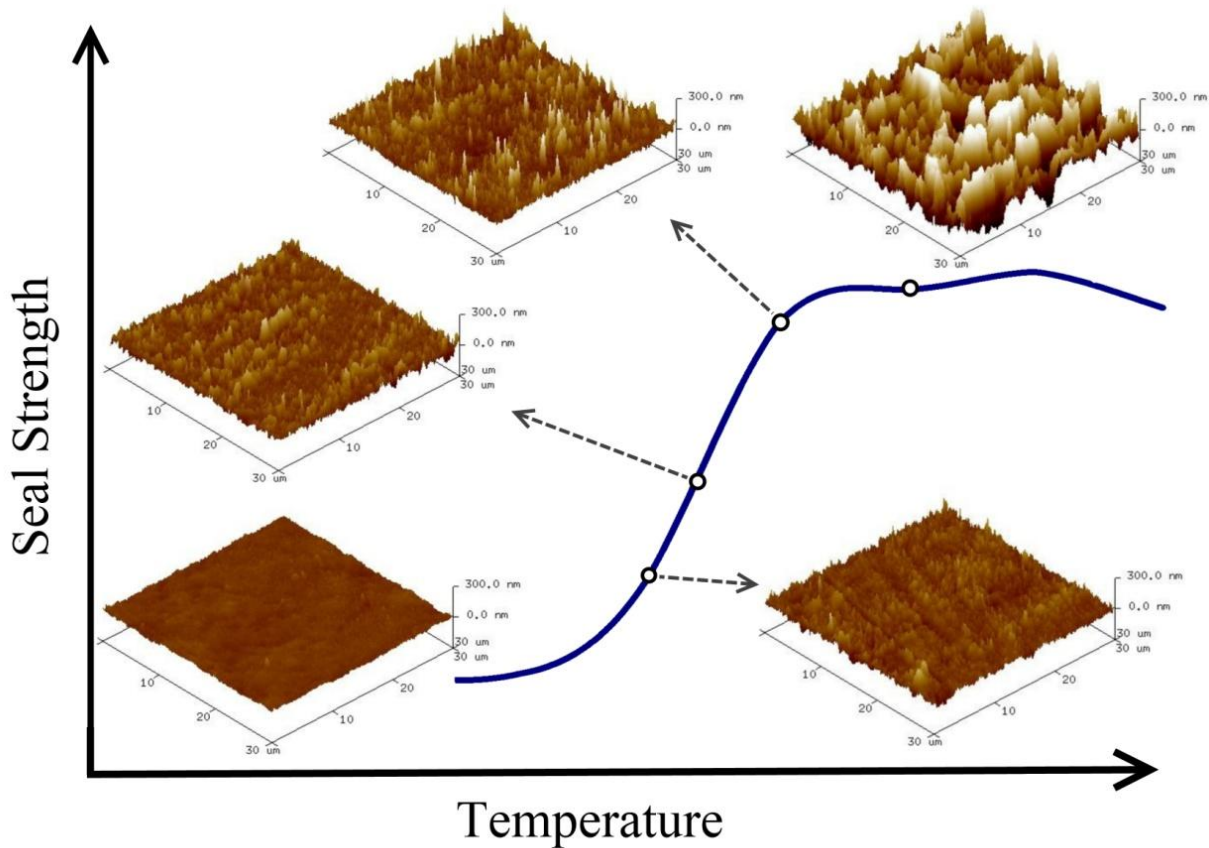


Figure 4-9 AFM images of peeled surface of samples sealed at $t=0.5$ s and $P=0.5$ Nmm⁻²

The roughness parameters (RMS and R_a), using various statistical measurements, provide information about both absolute and relative features of surface topography³⁸. These parameters, for different scanned samples, provide adequate information and a clear idea about the characteristics of a peeled surface on one hand, and produce a base for comparison among peeled surfaces on the other hand. This analysis was carried out only on the peeled surfaces resulting from interfacial failure.

In AFM experiments, it is well known that the image size affects surface roughness measurements³⁹⁻⁴¹. In order to compare peeled surfaces, roughness calculations have to be made

from a fixed image size. Therefore, the largest available high quality images ($30 \times 30 \mu\text{m}^2$) were selected to obtain the roughness values.

AFM variables were optimized to get precise topography with high clarity. In this case, AFM was capable of producing phase image and three-dimensional surface image with spatial resolution of a few nanometres.

Regarding basic roughness measurements, the roughness analysis parameters to consider are: height range, *RMS* and R_a . Considering Z as height from mean plane of the image, average roughness R_a is the arithmetic average of the absolute values of the surface height deviations measured from the mean plane:

$$R_a = \frac{1}{N} \sum_{i=1}^N |Z_i|$$

The Image *RMS* is the root mean square average of height deviations taken from the mean data plane, it is expressed as:

$$RMS = \sqrt{\frac{\sum_{i=1}^N Z_i^2}{N}}$$

Which N is the number of points within the image; Z_i is the current Z value. Image Surface area is the three-dimensional area of the entire image. The image height range indicates the maximum vertical distance between the highest and lowest data points in the image.

Figure 4-10(b) shows a comparison between peeled surfaces of samples which were sealed at the same dwell time and pressure but different temperatures regarding the different roughness parameters: height range, surface area, *RMS*, and R_a . The height range value has wider range for high sealing temperatures. Surface area as well as *RMS* and R_a increased with sealing temperature and diffusion improves consequently. Compared to the original film surface, much rougher surfaces were observed after sealing and the subsequent T-peel test.

All roughness parameters for samples sealed at 120°C reveal drastically higher values in comparison to the first four samples which show relatively high specimen surface roughness. It is clear that when strong adhesive bonds are broken, the resulting fracture surfaces are often

extremely rough. After peeling, the film surfaces in sealed parts are opaque. This is believed to be due to scattering of visible light, which become effective when *RMS* roughness is over 50 nm.

Figure 4-10(a) shows the measured roughness parameters from peeled surfaces of the films which were sealed for different dwell times (0.1, 0.5, 2.5 s) at their plateau initiation temperature. All the measured roughness parameters are in the same range, which illustrates that the diffusion reached to a certain point at T_{pi} regardless of the process conditions. Although T_{pi} is different at each dwell time, the roughness parameters are similar for the various sealing temperature and dwell time.

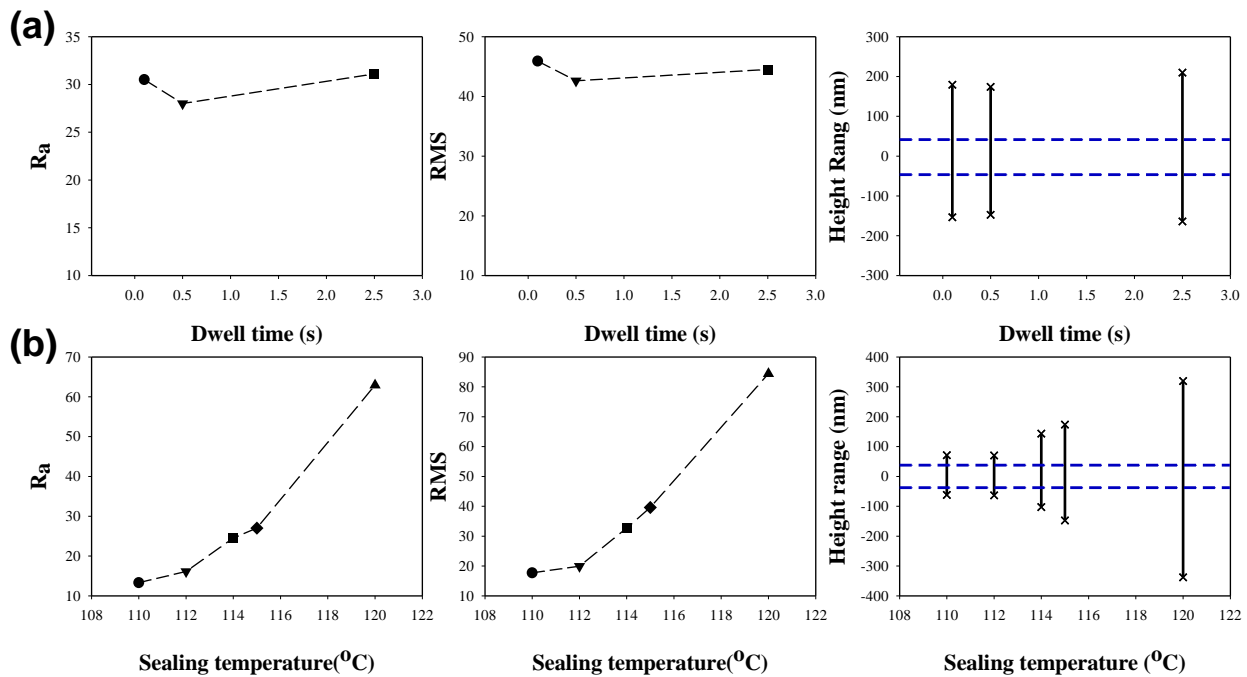


Figure 4-10 Roughness parameters of (a) peeled surface of samples sealed 0.5 Nmm⁻² at $t=0.1$, 0.5, 2.5 s at $T=T_{pi}$; Blue dash lines in height rang graph are the height range of film before seal, (b) peeled surface of samples sealed at $t=0.5$ s and $P=0.5$ Nmm⁻² and $T= 110, 112, 114, 115, 120^{\circ}\text{C}$

Figure 4-11 shows SEM images of the original films before adhesion and the peeled surfaces after sealing and the subsequent T-peel test. In SEM image of peeled surfaces, the fractured fibrils are visible. In specimens sealed at higher temperature (120°C) the fibrillar morphology developed into larger, higher fracture points and became more complex. They thus appear more like a network.

The changes in craze morphology and the significant increase in seal strength correspond to the increased number of chain stitches and entanglements across the interface. The number of stitches bonding the interface depends on the breadth and integrity of the interfacial region, which is determined by the number of available amorphous chains ⁴².

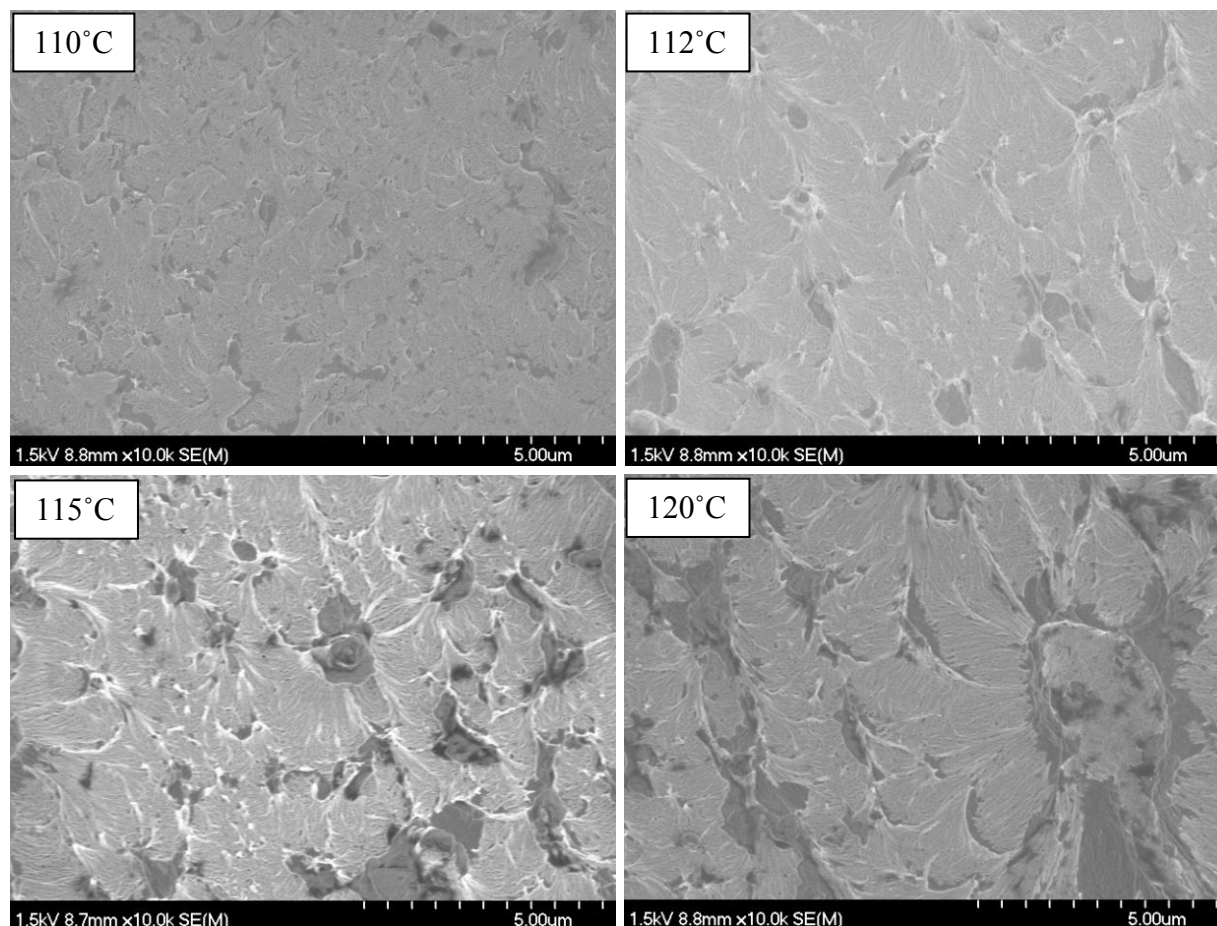


Figure 4-11 SEM micrographs of peel surfaces from films sealed at $t=0.5$ s, $P=0.5$ Nmm⁻² and $T=110, 112, 115, 120^{\circ}\text{C}$

It has been suggested that the fibrillar morphology is originated from interlamellar tie molecules. It bears the applied stress of peeling and form tight stretched links between crystals. Local yielding and microvoiding pull the material into highly oriented craze fibrils ²².

In Partially molten state some chains diffuse across the interface, while non molten crystals remain at both film surfaces unaffected and immobile. Chain mobility is strongly inhibited by crystals in this case, so the interdiffusion is limited and the adhesion strength is low. Sealing at higher temperature enhances the breadth of movable interfacial areas and transform this “micro-

spot adhesion” into a more integrated adhesion. Then it produces an interconnected texture in the fractured area of peeled surfaces ⁴³.

4.5 Conclusions

In this work, the main variables of the heat sealing process were evaluated in term of their influence on the development of seal strength. Interrelated influence of “dwell time - sealing temperature” and “pressure - sealing temperature” was established simultaneously. This optimization was performed on LLDPE as the most widely used polymer in seal application. It was shown that dwell time and temperature affect seal strength through the same mechanism of crystal melting and interdiffusion. However, pressure affects seal strength through wetting as a very different microscopic scale mechanism. It was concluded that the time and temperature dependence of seal strength is a consequence of the amount of heat available in the interface of films.

It was found that the seal strength has a linear correlation with square root of sealing time. This was shown to be in agreement with healing theory. Moreover, the slope of this linear correlation increases with temperature.

Analysis of melting distribution revealed that the amorphous fraction of film required for high seal strength is not a unique value for all the process conditions. And depending on the process variables including temperature and dwell time, it may locate within a certain range.

Peeled seal surfaces roughness analysis revealed that its topography strongly depends on the seal creation process variables. The fibrillar structure was observed by AFM and SEM from the peeled surfaces of the samples sealed at low temperatures. In specimens sealed at high temperature, the fibrillar morphology developed into larger, higher fracture points and became more complex in shape.

4.6 Acknowledgement

Financial support from 3S Pack NSERC/Saputo/Prolamina industrial research chair is gratefully acknowledged.

4.7 References

1. Meka, P. & Stehling, F. C. Heat sealing of semicrystalline polymer films. I. Calculation and measurement of interfacial temperatures: Effect of process variables on seal properties. *J. Appl. Polym. Sci.* **51**, 89–103 (1994).
2. Stehling, F. C. & Meka, P. Heat sealing of semicrystalline polymer films. II. Effect of melting distribution on heat-sealing behavior of polyolefins. *J. Appl. Polym. Sci.* **51**, 105–119 (1994).
3. Hassan, A. Effect of bar sealing parameter on OPP/MCPP heat seal strength. *Express Polym. Lett.* **1**, 773–779 (2007).
4. Theller, H. W. Heatsealability of Flexible Web Materials in Hot-Bar Sealing Applications. *J. Plast. Film Sheeting* **5**, 66–93 (1989).
5. De Gennes, P. G. Reptation of a Polymer Chain in the Presence of Fixed Obstacles. *J. Chem. Phys.* **55**, 572–579 (1971).
6. Frederix, C., Beauchene, P., Seguela, R. & Lefebvre, J. M. Kinetics of the non-isothermal fusion-welding of unlike ethylene copolymers over a wide crystallinity range. *Polymer* **54**, 2755–2763 (2013).
7. Kim, K. D., Sperling, L. H., Klein, A. & Hammouda, B. Reptation Time, Temperature, and Cosurfactant Effects on the Molecular Interdiffusion Rate during Polystyrene Latex Film Formation. *Macromolecules* **27**, 6841–6850 (1994).
8. Klein, J. The Interdiffusion of Polymers. *Science* **250**, 640–646 (1990).
9. Tashiro, K. & Gose, N. Diffusion and aggregation of hydrogeneous and deuterated polyethylene chains at their interfacial boundary as studied by time- and space-resolved FTIR microscopic measurements. *Polymer* **42**, 8987–8998 (2001).
10. Wool, R. P. & O'Connor, K. M. A theory crack healing in polymers. *J. Appl. Phys.* **52**, 5953–5963 (1981).
11. Zhao, R. & Macosko, C. W. Polymer–polymer mutual diffusion via rheology of coextruded multilayers. *AIChE J.* **53**, 978–985 (2007).
12. Gardon, J. L. Peel adhesion. II. A theoretical analysis. *J. Appl. Polym. Sci.* **7**, 643–665 (1963).
13. Gardon, J. L. Peel adhesion. I. Some phenomenological aspects of the test. *J. Appl. Polym. Sci.* **7**, 625–641 (1963).
14. Hashimoto, Y., Ishiaku, U. S., Leong, Y. W., Hamada, H. & Tsujii, T. Effect of heat-sealing temperature on the failure criteria of oriented polypropylene/cast polypropylene heat seal. *Polym. Eng. Sci.* **46**, 205–214 (2006).
15. Poisson, Hervais, V., Lacrampe, M. F. & Krawczak, P. Optimization of PE/Binder/PA extrusion blow molded films. I. Heat sealing ability improvement using PE/EVA blends. *J. Appl. Polym. Sci.* **99**, 974–985 (2006).
16. Dowling, D. P. *et al.* Atmospheric pressure plasma treatment of amorphous polyethylene terephthalate for enhanced heatsealing properties. *Int. J. Adhes. Adhes.* **35**, 1–8 (2012).
17. Morris, B. A. Predicting the Heat Seal Performance of Ionomer Films. *J. Plast. Film Sheeting* **18**, 157–167 (2002).

18. Tetsuya, T., Ishiaku, U. S., Mizoguchi, M. & Hamada, H. The effect of heat sealing temperature on the properties of OPP/CPP heat seal. I. Mechanical properties. *J. Appl. Polym. Sci.* **97**, 753–760 (2005).
19. De Gennes, P.-G. Entangled polymers. *Phys. Today* **36**, 33–39 (1983).
20. Gennes, P.-G. de. *Introduction to Polymer Dynamics*. (CUP Archive, 1990).
21. Prager, S. & Tirrell, M. The healing process at polymer–polymer interfaces. *J. Chem. Phys.* **75**, 5194–5198 (1981).
22. Wool, R. P., Yuan, B.-L. & McGarel, O. J. Welding of polymer interfaces. *Polym. Eng. Sci.* **29**, 1340–1367 (1989).
23. Anderson K.L., Wescott J.T., Carver T.J. & Windle A.H. Mesoscale modelling of polymer welding. *Mater. Sci. Eng. A* **365**, 14–24 (2004).
24. Ezekoye, O. A., Lowman, C. D., Fahey, M. T. & Hulme-Lowe, A. G. Polymer weld strength predictions using a thermal and polymer chain diffusion analysis. *Polym. Eng. Sci.* **38**, 976–991 (1998).
25. Wool, R. P. Adhesion at polymer–polymer interfaces: a rigidity percolation approach. *Comptes Rendus Chim.* **9**, 25–44 (2006).
26. Boiko, Y. M., Guérin, G., Marikhin, V. A. & Prud'homme, R. E. Healing of interfaces of amorphous and semi-crystalline poly(ethylene terephthalate) in the vicinity of the glass transition temperature. *Polymer* **42**, 8695–8702 (2001).
27. Xue, Y.-Q., Tervoort, T. A. & Lemstra, P. J. Welding behavior of semicrystalline polymers. 1. The effect of nonequilibrium chain conformations on autoadhesion of UHMWPE. *Macromolecules* **31**, 3075–3080 (1998).
28. Xue, Y.-Q., Tervoort, T. A., Rastogi, S. & Lemstra, J. Welding Behavior of Semicrystalline Polymers. 2. Effect of Cocrystallization on Autoadhesion. *Macromolecules* **33**, 7084–7087 (2000).
29. Yu, W., Peri, S. R., Akgun, B. & Foster, M. D. Manipulation of Polymer/Polymer Interface Width from Nonequilibrium Deposition. *ACS Appl. Mater. Interfaces* **5**, 2976–2984 (2013).
30. Wunderlich, B. & Czornyj, G. A Study of Equilibrium Melting of Polyethylene. *Macromolecules* **10**, 906–913 (1977).
31. Mueller, C., Capaccio, G., Hiltner, A. & Baer, E. Heat sealing of LLDPE: relationships to melting and interdiffusion. *J. Appl. Polym. Sci.* **70**, 2021–2030 (1998).
32. Kim, S. *et al.* Enhanced Interfacial Adhesion between an Amorphous Polymer (Polystyrene) and a Semicrystalline Polymer [a Polyamide (Nylon 6)]. *ACS Appl. Mater. Interfaces* **3**, 2622–2629 (2011).
33. Buckley, C. P., Wu, J. & Haughie, D. W. The integrity of welded interfaces in ultra high molecular weight polyethylene: Part 1—Model. *Biomaterials* **27**, 3178–3186 (2006).
34. Wool, R. P. *Polymer Interfaces: Structure and Strength*. (Hanser-Gardner Publications, 1995).
35. Kim, Y. H. & Wool, R. P. A theory of healing at a polymer–polymer interface. *Macromolecules* **16**, 1115–1120 (1983).
36. Wool, R. P. Self-healing materials: a review. *Soft Matter* **4**, 400–418 (2008).
37. Bousmina, M., Qiu, H., Grmela, M. & Klemberg-Sapieha, J. E. Diffusion at Polymer/Polymer Interfaces Probed by Rheological Tools. *Macromolecules* **31**, 8273–8280 (1998).

38. Marcott, S. A., Ada, S., Gibson, P., Camesano, T. A. & Nagarajan, R. Novel Application of Polyelectrolyte Multilayers as Nanoscopic Closures with Hermetic Sealing. *ACS Appl. Mater. Interfaces* **4**, 1620–1628 (2012).
39. Kamal, M. R., Tang, Z. & Huang, T. Morphological characterization of PE blown films by atomic force microscopy. *Int. Polym. Process.* **16**, 376–387 (2001).
40. Teichert, C., Haas, A., Wallner, G. M. & Lang, R. W. Nanometer scale characterization of polymer films by atomic-force microscopy. *Macromol. Symp.* **181**, 457–466 (2002).
41. Simpson, G. J., Sedin, D. L. & Rowlen, K. L. Surface Roughness by Contact versus Tapping Mode Atomic Force Microscopy. *Langmuir* **15**, 1429–1434 (1999).
42. Cole, P. J., Cook, R. F. & Macosko, C. W. Adhesion between Immiscible Polymers Correlated with Interfacial Entanglements. *Macromolecules* **36**, 2808–2815 (2003).
43. Yang, H., Ward, T. C. & Zhang, W. Interfacial Interpretation of Autohesion of Ethylene/1-Octene Copolymers by Atomic Force Microscopy. *J. Adhes.* **83**, 1043–1068 (2007).

CHAPTER 5

**ARTICLE 2: ROLE OF MOLECULAR ARCHITECTURE IN
INTERFACIAL SELF-ADHESION OF POLYETHYLENE FILMS***

Zahra Najarzadeh, Abdellah Ajji

5.1 Abstract

The influence of molecular architecture on interfacial self-adhesion above melting temperature of polyethylene films was examined in this study. The investigated molecular structures include molecular weight (M_w), molecular weight distribution (MWD), amount and distribution of long chain branch (LCB) and short chain branch (SCB) distribution among and along polyethylene chains. The amount of long and short chain branches was quantified using gel permeation chromatography (GPC) and nuclear magnetic resonance (NMR) techniques, respectively. The adhesion strength was measured immediately after melt bonding using T-Peel test. The results showed that increasing M_w resulted in higher adhesion strength in linear metallocene ethylene α -olefins. The presence of even a low amount of LCB hinders the reptation motion and diffusion, and resulted in lower adhesion strength in the metallocene ethylene α -olefins. In addition, highly branched chains of low density polyethylene (LDPE) yielded to a very low self-adhesion. A drastic difference in adhesion strength between metallocene and conventional linear low density polyethylene (LLDPE) was observed and was attributed to the homogeneity versus heterogeneity of composition distribution, including MWD and SCB distribution. The low interfacial self adhesion in the conventional polyethylene was concluded to be due to enrichment of highly branched low molecular weight chains at the surfaces of the films. These segregated chains at the interface diffuse before the high molecular weight chains located in the bulk.

* Submitted to Macromolecules, March 2014.

5.2 Introduction

Interfacial adhesion of polymer surfaces, which is categorized into either symmetric or asymmetric types, is an immensely complicated subject. Self adhesion is a symmetric type of adhesion having critical importance in some specific applications such as flexible packaging industry. It happens through inter-diffusion of chains at the interface at temperatures high enough to allow chains to perform segmental motion through reptation in a reasonable process time. Knowing that interface strength development depends on the total molecular crossing density ¹, chain diffusion has the main role in strength build up at interface.

The foundation for our current understanding of chain dynamics in polymer diffusion is reptation model. Many researchers have tried to explain the reptation mechanism associated with the influence of several parameters such as molecular weight, repeat unit, long and short chain branches etc. ¹⁻⁴.

Former studies on interfacial adhesion were concerned with the strength of the interface at room temperature ^{2,5-7}. To eliminate the additional complication due to crystallization upon cooling, the focus has been mostly on amorphous polymers ⁸. Thus, the molecular characteristics for effective interfacial reinforcement are relatively well established for glassy polymers. But there is less information on the interfacial strength of semicrystalline polymers ⁹. Studies on self adhesion of semicrystalline polymers are limited to a few cases, in which self adhesion occurred at temperatures higher than the melting point and adhesion strength measurements performed at room temperature ¹⁰⁻¹⁴.

Another reason for limited fundamental studies on self adhesion in the melt state is that the measurement of interfacial adhesion between two polymer melts is practically challenging. Recently, Schach et.al ⁹ used a custom-designed probe test (known for pressure sensitive adhesives) to measure the tack of SBR random copolymers. In this method two polymer layers with different thicknesses of 1 and 200 μm were bonded to silicon and glass substrates, respectively ⁹. The thinner layer of SBR (1 μm) was attached to a moving probe. Adhesion occurred when the probe approached the other layer for a certain contact time and pressure. Then by recording the required force for debonding in the last step, the adhesion strength was measured.

Although this method might apply for adhesive applications at the industrial scale, it is not realistic for flexible packaging. In this work, the T-Peel test was used by a machine capable of measuring the adhesion strength immediately after melt adhesion while interface is still molten. This technique allows performing adhesion in a fraction of a second and the measurement of interface strength in a few milliseconds. This creates a unique opportunity to study self adhesion at very short time which is very scarce in literature.

Since the development and commercialization of metallocene single site catalysts polyolefins, these have found significant applications and use in packaging industry. They have been used as sealant layer in various types of packaging structures and their performance was observed to be far better than that of conventional polyethylenes. However, a clear understanding of the mechanisms of action and interpretation of this high performance is still to be developed. Single site catalysts polymerization enabled a great opportunity for the control of polydispersity, type and distribution of branch, and branch content. These structural features at the molecular scale have modulated the physical properties of metallocene PE in the solid and melt states, such as crystallinity, bulk mechanical properties, melt rheology and flow behaviour. The general term of homogeneous molecular structure used for m-PE, refers to narrow molecular weight distribution, homogeneous distribution of branches among the chains and along one chain.

To our knowledge, there is no study on the role of molecular architecture of a polymer chain on the self adhesion strength at the melt state. This work is aimed at investigating the micro-mechanisms and molecular structure features involved in this property for m-PEs. The investigated molecular structures include: molecular weight (M_w), molecular weight distribution (MWD), amount and distribution of long chain branch (LCB), and short chain branch (SCB) distribution among and along polyethylene chains. To reach this goal, the materials were selected from a wide range of the polyethylene family from highly branched to linear, high molecular weight to low, and heterogeneous to homogeneous composition distribution.

5.3 Experiments

5.3.1 Materials and films preparation

Five commercial polyethylene plastomers (mSC1, mSC2, mSC3, mSC4, and mSC5) produced with Exxpol™ metallocene catalyst were supplied by ExxonMobil. Two commercial long chain

branched metallocene polyethylene were selected from Enable™ metallocene polyethylene supplied by ExxonMobil (mLC1), and from AFFINITY™ metallocene polyethylene supplied by Dow chemical company produced with INSITE™ technology (mLC2). Conventional LDPE, LLDPE, and HDPE polymers were also supplied by ExxonMobil Company. Table 5-1 shows the α -olefin comonomer type, the molecular weight (determined by gel permeation chromatography), and other characteristics of the polymers. Films of 50 μm in thickness were prepared using a cast film line. A 45 mm Killion single screw extruder followed by 2 mm opening and 20 cm width slit cast die were used. The extrusion temperatures for every polymer were adjusted according to their melting and processing temperature profiles recommended by the suppliers. The extrusion was carried out at 40 rpm and the distance between the die exit and nip rolls was 10 cm. An air knife and calendar cooling system were used to cool the films after the die exit. Extrusion speed and the collecting speed were kept the same for all resins in order to control draw ratio and have uniform final thickness.

Table 5-1 Main characteristics of polyethylene resins

Nomenclature	MFI (190°C/2.16kg) (g/10min) ^a	α -olefin comonomer ^a	Density ^a (g.cm ⁻³)	T _m ^a (°C)	M _w ^b (kg.mol ⁻¹)	PDI ^b
LDPE	2	-	0.923	110	160	8.756
Zn-LLDPE	2	butene	0.918	123	133	5.448
HDPE	2.5	-	0.965	134	187	9.492
mSC1	2	hexene	0.918	118	102	2.545
mSC2	2.2	hexene	0.895	88	111	2.156
mSC3	1.2	hexene	0.900	96	123	2.329
mSC4	3.5	hexene	0.900	95	96	1.869
mSC5	7.5	hexene	0.900	94	73	2.794
mLC1	0.5	hexene	0.920	114	115	2.636
mLC2	1	octene	0.902	99	115	2.12

^a Provided by the manufacturer. ^b Obtained from high temperature GPC.

5.3.2 Differential scanning calorimetry (DSC)

In order to determine crystallinity and melting temperature of the various PEs, differential scanning calorimetry (DSC) was used with the help of a TA Instruments Q1000 calorimeter. The melting properties of films are needed to determine the range of temperature for adhesion

experiments. The melting temperature and degree of crystallinity of the films were determined from the first heating ramp. The samples were heated from room temperature to 200°C at a heating rate of 10°C/min. In order to eliminate initial thermal history, samples were equilibrated for three minutes at 200°C, then cooled down to -90°C by cooling rate of 10°C/min to obtain the crystallization point. For crystallinity determinations, a value of 290 J.g⁻¹ was taken as the enthalpy of fusion of a 100% crystalline PE¹⁵.

5.3.3 Nuclear magnetic resonance spectroscopy (NMR)

NMR spectra were recorded on Varian Inova 600 (Agilent, Santa Clara, CA, USA) operating at frequencies of 150.87 MHz for ¹³C, using a double resonance 5mm broadband probe. Spectra were recorded following the ASTM protocol D-5017 – 96 for the determination of composition of low density polyethylene. Briefly, the quantitative ¹³C NMR spectra were recorded at 120 °C with 10 s recycle delay, 90° pulses (16 μs long), an acquisition time of 2 s, full decoupling and a spectral width of 250 ppm. Spectra were accumulated until a signal to noise ratio of 2500:1 was obtained using 2 Hz exponential apodization.

Samples were dissolved in 1,1,2,2-tetrachloroethane-d₂ with concentrations of more than 35%wt in 10 mm sample tubes to get the much needed sensitivity. Figure 5-1 gives a typical spectrum of a hexene based ethylene-α-olefin sample and Table 5-2 shows the assignment of the different chemical shifts according to Randall¹⁶.

5.3.4 Gel permeation chromatography (GPC)

Molecular weight and molecular weight distribution were determined on a Viscotek HT-GPC Triple Detection. The samples were dissolved in 1,2,4-trichlorobenzene (TCB) and the measurements were carried out at 150°C. The OmniSEC software was used to determine the number of long chain branches in 10⁴ carbon atom. The calculations were based on tri-functional Zimm-Stockmayer equation¹⁷⁻¹⁹.

5.3.5 Interfacial self-adhesion measurements

Adhesion experiments were performed using SL10 Lako-Tool equipment based on T-Peel test method. Heat bonding was made with two metallic jaws of 19.1 mm×25.4 mm covered by Teflon. The film samples were cut in 2.54 cm×33 cm strips specimens parallel to machine

direction. This method consists in two stages of bonding and peeling: in the first stage, two layers of film locate between two jaws, then the jaws approach to provide the contact between the two sides of the film with a controlled pressure for 0.5 s. In the second stage, the jaws were removed and simultaneously two arms of the machine pull the films to measure the adhesion strength, while still hot (also called hot tack). To avoid film distortion due to direct contact to heated jaw, the PE films were laminated to a layer of PET support film using a polyurethane based solventless adhesive. To obtain reliable results for adhesion in the molten state, the PE films must be strongly attached to the support film. The adhesion experiments were performed in a range of jaw temperatures from 50°C to 140°C at a constant pressure of 0.5 MPa and dwell time of 0.5 s. The values reported in adhesion graphs are averages of at least five tests.

5.3.6 Extensional rheology

The melt extensional behaviour of the polymers was determined using the SER geometry of the ARES Rheometric Scientific rheometer. The data were recorded at a Hencky strain rate of $\dot{\epsilon}=1 \text{ s}^{-1}$ on samples having dimensions of 13mm×18mm×0.8mm. Since the adhesion measurements were performed at different temperatures, the melt strength measurements should be also be performed at the corresponding temperature for every polymer. Thus, in order to achieve coherent results, $T_m+10^\circ\text{C}$ was selected to be the reference temperature for each resin.

5.4 Results and discussion

5.4.1 Material characterization

Molecular weight and molecular weight distribution, long and short chain branching (LCB and SCB) are the most important structural variables influencing the properties of polyethylene. Therefore, it is necessary to obtain qualitative and quantitative information on the nature and number of chain branches. High resolution ^{13}C NMR spectroscopy was used to determine the number of short chain branches per 10^3 carbon atoms.

Figure 5-1 shows the spectrum of mSC3 as a sample of hexene based ethylene α -olefin copolymers and the assignment of important chemical shifts. A tabulation of chemical shifts and assignments are listed in Table 5-2. The nomenclatures and chemical shifts assigned to different carbonyl groups were introduced by Randall and later developed by others¹⁶. The Greek

alphabets (α , β , γ , δ^+) are used to denote the positions of a given backbone carbon site relative to the methane carbons and side-chain carbons. For SCB the label has the format mB_n , where n represents the length of the side chain and m refers to the position of the carbon. In the sequence assignments, H and E represent the hexene comonomer and ethylene monomer, respectively. The $\delta^+\delta^+$ peak was set at 29.98 ppm as the reference peak. The tertiary carbon atom of the branch point is seen at 38.2, 35.9, and 34.2 ppm. The SCB quantification was obtained by the ratio of integrals associated with a branch site to that of the CH_2 bulk sites:

$$SCB \text{ mole}\% = \frac{A_{\alpha}}{2A_{Total}}$$

Through this general equation, the mole% of SCB was obtained using the detailed collective assignments method. The calculations can be found in ASTM-D5017-96²⁰⁻²². The results are reported along with the number of branches per 1000 carbon atoms in Table 5-3 for ethylene α -olefin copolymers. In accordance with the categorization of comonomer content and density of ethylene α -olefin copolymers proposed by Bensason et al.²³, the calculated comonomer mole% of resins correlates with their density and crystallinity.

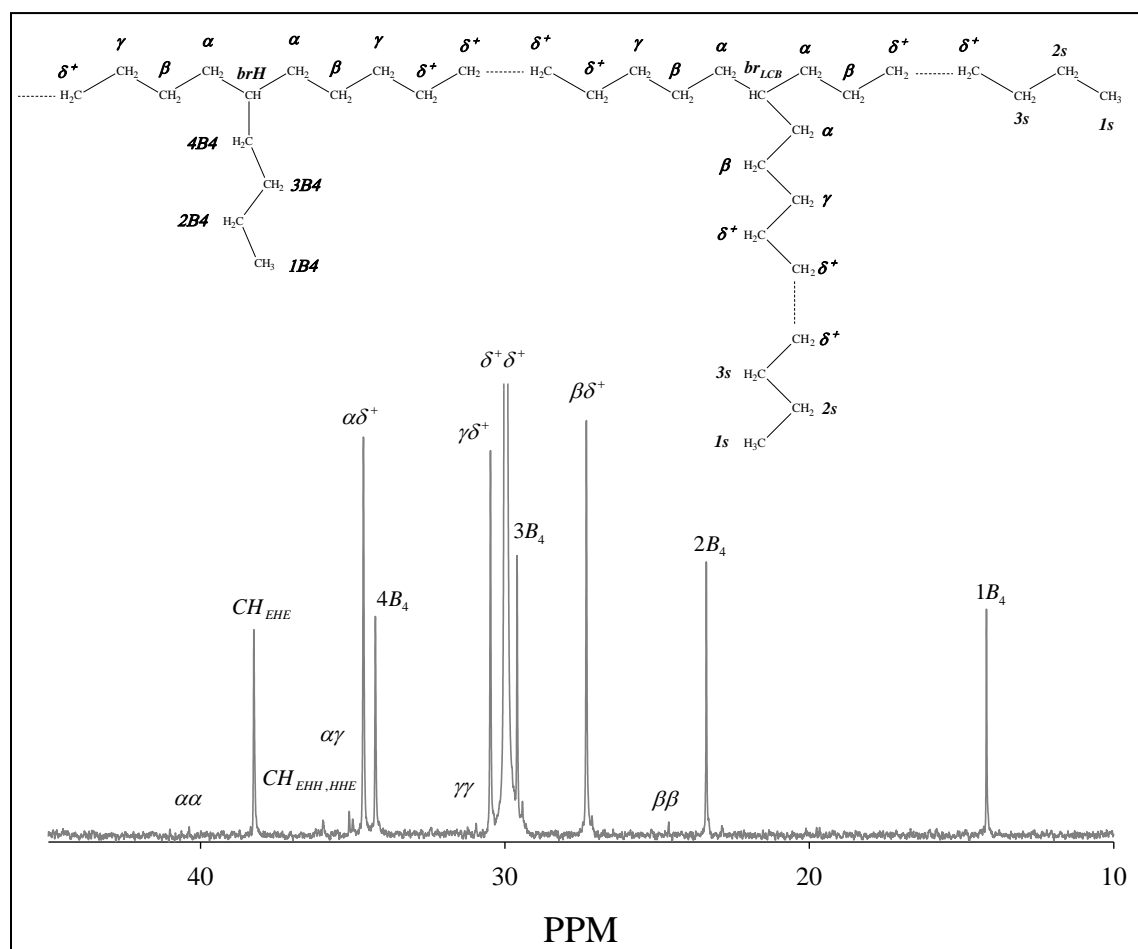


Figure 5-1 The nomenclature and peak assignments example of ethylene-1-hexene copolymer with the hypothetical presence of LCB, and ^{13}C NMR spectrum of a metallocene ethylene-1-hexene copolymer at 120°C using *1,1,2,2-tetrachloroethane- d_2* as solvent.

Table 5-2 NMR spectrum peaks and peak assignments observed for an ethylene-1-hexene copolymer

Chemical shift (ppm)	Carbon assignment	Sequence assignment
38.22	Methine	EHE
35.92	Methine	EHH+HHE
35.10	$\alpha\gamma$	HHEH+HEHH
	$\alpha\gamma$	EHEH+HEHE
35.02	$\alpha\delta+$	HHEE+EEHH
	4B4	EHH+HHE
34.62	$\alpha\delta+$	EHEE+EEHE
34.21	4B4	EHE
	Methine	HHH
30.94	$\gamma\gamma$	HEEH
30.48	$\gamma\delta+$	HEEE+EEEH
29.98	$\delta+\delta+$	(EEE) _n
29.58	3B4	EHE
29.41	3B4	EHH+HHE
29.24	3B4	HHH
27.31	$\beta\delta+$	EHEE+EEHE
27.13	$\beta\delta+$	HHEE+EEHH
24.60	B β	EHEHE
	B β	EHEHH+HHEHE
	B β	HHEHH
23.39	2B4	EHE+EHH+HHE+HHH
14.21	Methyl	EHE+EHH+HHE+HHH

Table 5-3 Mole % comonomer and number of SCBs per 1000 carbon atom of PE resins

Nomenclature	Mole% comonomer	Br/1000 C atom
LDPE	-	-
Zn-LLDPE	5.31	42.45
HDPE	-	-
mSC1	2.03	10.09
mSC2	7.71	32.71
mSC3	5.68	25.41
mSC4	5.48	24.6
mSC5	6.72	29.26
mLC1	3.050	24.12
mLC2	6.037	37.5

The HDPE and LDPE are excluded from this calculation because of the linear structure of the former and the highly branched structure of the latter. LDPE structure consists of several kinds of LCB and SCB, which is due to the backbiting mechanism in free radical polymerization of ethylene in high pressure reactors. Therefore, obtaining the quantitative level of branches would be practically difficult and unreliable ²⁴.

In this work, GPC was used to obtain the quantitative amount of the long chain branches in the resins containing LCB. Although NMR spectroscopy is used as another experimental technique for measuring LCB in polyethylene, it has the disadvantage of treating the alkyl branches in an equal manner and defines branches longer than C₆ as long branches. From the rheology and diffusion perspective, LCB corresponds to the branches longer than critical entanglement molecular weight (M_c), hence are capable of making entanglements. Thus, in this application, the use of NMR continues to be problematic and still a subject of debate in literature ²⁵⁻²⁷. For the purposes of the present work, LCB refers to the one defined from the rheology and diffusion perspective, and it was determined from GPC.

Table 5-4 shows the DSC results of PE films. The nominal crystallinity (X_C), melting temperature (T_m), and crystallization temperature (T_c) as determined by DSC were observed to be almost identical functions of mole percent comonomer content for LLDPE polymers. The metallocene polymers (mSC2, mSC3, mSC4, mSC5, mLC2) with a high comonomer content of 5.4-7.7 mol% show the lowest T_m (T_m=90-102°C) and X_C (X_C=12.7-14.6%). While mSC1 and mLC1 with the

comonomer contents of 2.03 and 3.050 mol%, respectively, show $T_m=114^\circ\text{C}$. Although mSC1 and mLC1 have the lowest SCB content among the metallocene resins, their T_m and X_C are still lower than Zn-LLDPE. Zn-LLDPE which has quite high 5.31 mol% comonomer content shows $T_m=121^\circ\text{C}$ and $X_C=20\%$. This is due to the method of SCB incorporation in the molecular structure of Zn-LLDPE and the heterogeneity in comonomer composition distribution.

Table 5-4 Crystallinity properties of films

Nomenclature	T_m ($^\circ\text{C}$)	T_c ($^\circ\text{C}$)	ΔH (J.g^{-1})	X_C
LDPE	107	96	71.1	24.5
Zn-LLDPE	121	107	59.4	20.5
HDPE	132	119	187.4	64.6
mSC1	114	102	65.9	22.7
mSC2	90	72	36.9	12.7
mSC3	96	80	38.0	13.1
mSC4	95	79	38.3	13.2
mSC5	96	79	37.8	13.3
mLC1	114	104	73.8	25.4
mLC2	102	88	42.2	14.6

According to the classification proposed by Bensason et al.²³ and stated by Bubeck²⁸ on crystalline morphology of homogeneous ethylene α -olefin copolymers. The crystalline structure of mSC1 and mLC1 are speculated to be a mixed morphology of small lamellae and bundled crystals. Therefore these materials may form very small spherulites. The crystalline structure of mSC2, mSC3, mSC4, mSC5, mLC2 are speculated to be fringed micellar or bundled crystals which is implied to low crystallinity and melting temperature.

As stated by Hosoda et al.²⁹ the vast research on various kinds of ethylene α -olefin showed that the crystalline morphology of LLDPE is determined by three structural factors: the intermolecular comonomer composition distribution (CCD), the intramolecular comonomer distribution, and the identity of the comonomer. In Ziegler-Natta LLDPE, the multi active sites on the catalyst surface cause a heterogeneous intermolecular CCD. It creates high SCB concentration in low molecular weight chains and relatively low SCB concentration in high molecular weight chains. In addition to the broad MWD in Ziegler-Natta LLDPE, this non-uniformity facilitates the crystallization and results in a non homogeneous microstructure²⁸.

While in single site-metallocene catalyst LLDPE, intermolecular CCD and intramolecular CCD are uniform. Then comonomer sequence length distribution is uniform as a result of intramolecular CCD²⁸⁻³⁰.

SCB can be incorporated in LLDPE by copolymerisation of ethylene with α -olefin monomers, or by isomerisation reactions during ethylene homopolymerisation. They are incorporated into ethylene backbone as side branches typically at concentrations of 1- 19 mol%. Presence of SCB is particularly critical in morphology and solid state properties. Commonly they do not incorporate into crystal cells and act as structural defects during crystallisation^{28,29} and thus strongly affect size and amount of crystals, ultimately crystallinity, melting temperature, crystallisation rates, and consequently other solid state bulk properties. As discussed in connection with Table 5-4 and Table 5-3, DSC is very sensitive to the presence of branching. Through crystallinity, it provides qualitative insight into the degree of branching^{23,28}.

5.4.2 Effect of molecular weight on self-adhesion strength

Experimental investigation of molecular weight influence on self-adhesion strength requires a careful polymer selection. In order to eliminate the influence of other parameters (such as MWD, LCB and SCB content) polymers are selected from a group of linear metallocene α -olefin copolymers. These resins (mSC3, mSC4, mSC5) have narrow MWD and they are produced through the same polymerization method using the same metallocene catalyst technology. As seen in Table 5-1 and Table 5-3, they have the same short chain branches length, amount and distribution.

Figure 5-2 shows interfacial strength of ≈ 6000 , ≈ 5000 , and 3500 N.m^{-1} for mSC3, mSC4, and mSC5, respectively. It implies that the higher M_w results in a stronger interface. This is in agreement with what was reported in literature. It was found that long chains are more effective than shorter ones in strengthening the interface. The long molecules are more efficient to increase the energy of separation between polymer films than short chains^{3,31,32}.

On the other hand, in adhesion measurement using the peeling method, the melt strength of the material is determinant^{8,31,33} because the extensional field/force at the head of crack tip associates the adhesion strength to viscoelastic response of the material³³. In melt adhesion, interface strength is developed by the diffusion of chains across the interface. By peeling, the

applied force for separation of the films at the interface is determined by dis-entanglement or deformation of the entangled network of polymer chains at the interface. Peeling pulls out the diffused spots across interfaces which were developed as stitches between two surfaces. If v designates the peel rate and is 33 mm.s^{-1} and h the film thickness= $50 \text{ }\mu\text{m}$, the strain rate could be estimated roughly from $\dot{\gamma} = \frac{v}{h}$ relation. The strain rate would be $\approx 660 \text{ s}^{-1}$. So peeling is equivalent to applying an extensional force on entangled chains at very high strain rates. Therefore, the peeling of interface correlates the adhesion strength with melt strength of polymers^{34–36}. This was confirmed for self-adhesion of un-crosslinked elastomers, polyethylene and other thermoplastics^{8,31,37,38}. In the case of mSC3, mSC4, and mSC5, since the molecular architecture is similar, the reptation dynamic occurs at a similar mechanism. Then the melt strength is the distinct feature between them. The transient elongational viscosity of the resins at 1 s^{-1} strain rate is illustrated in Figure 5-3. It is clear from those results that the melt strength curves have the same sequence as adhesion strengths in Figure 5-2.

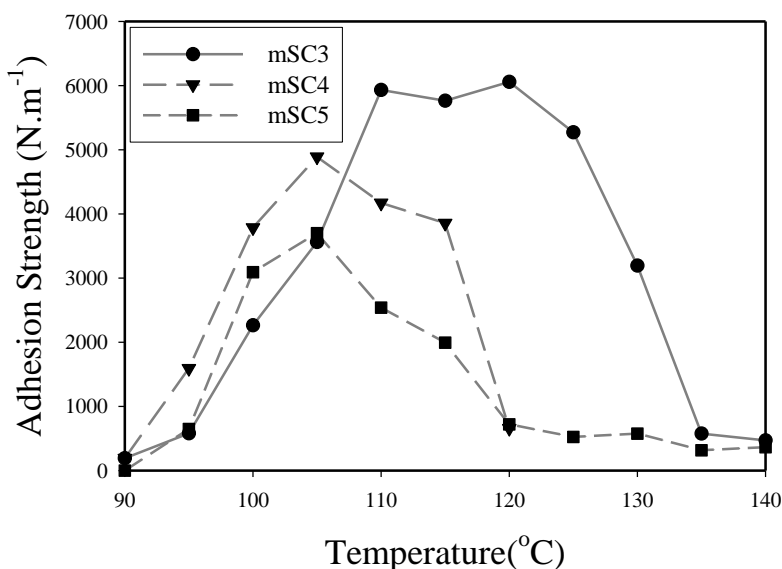


Figure 5-2 Self adhesion strength of mSC3, mSC4, and mSC5 films as a function of temperature, obtained from T-peel tests with peel rate of 33 mm.s^{-1}

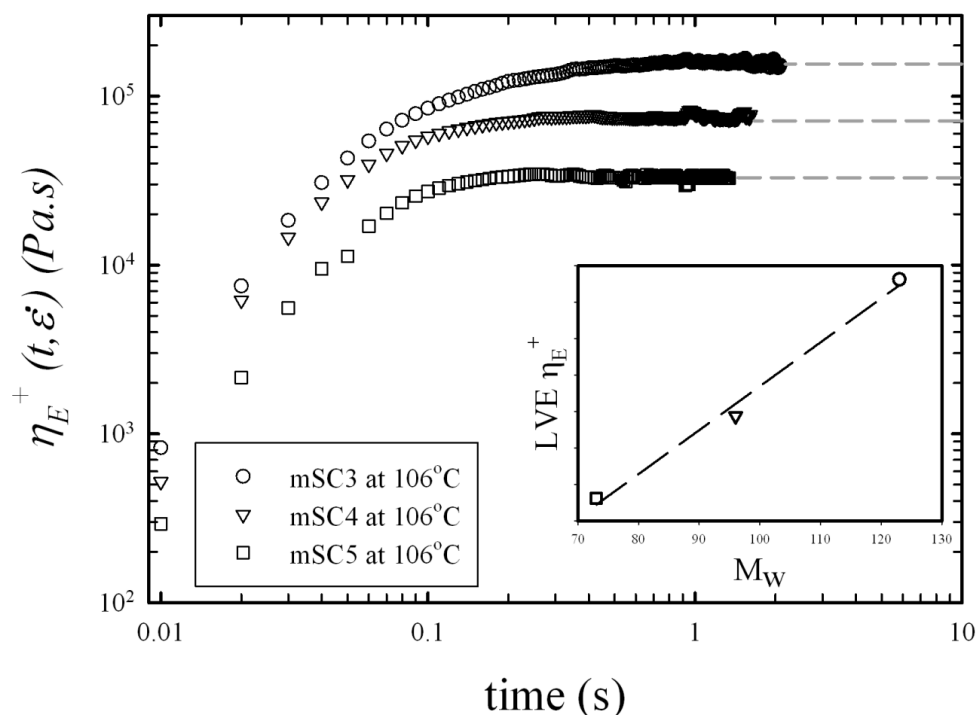


Figure 5-3 Transient elongational viscosities of mSC3, mSC4, and mSC5 at Hencky strain rate $\dot{\epsilon}=1 \text{ s}^{-1}$, the dash-lines represent the linear behaviour where the transient elongational viscosity is equal to three times of the zero shear viscosity in shear flow

Creton et al.³³ also explained the adhesion energy at the soft interface of polydimethylsiloxane elastomers in terms of energy dissipation mechanisms at molecular and macroscopic scales. At the molecular scale, the connector chains which are attached and well integrated into the interface resist against being extracted from the bulk polymer. At the macroscopic scale, because of the large extensional deformations involved in peeling, the bulk deformation in the viscoelastic material determines the fracture toughness. Thus the fracture energy strongly depends on the viscoelastic properties of material. Connector chains with higher molecular weight have higher toughness during peeling of the interface³³.

Assuming the temperature range of 110-125°C for mSC3, 100-115°C for mSC4, and 100-110°C for mSC5 as the range of temperatures in which they have reached a plateau of their maximum adhesion strength. This maximum adhesion strength is displayed as a function of molecular weight of the polymer in Figure 5-4. It shows a linear dependence of the maximum adhesion

strength of mSC3, mSC4, and mSC5 to molecular weight. To our knowledge, there is no report on the direct and linear dependence of self adhesion strength to molecular weight.

In the measurement of self-adhesion strength at temperatures higher than T_m , it is practically not possible to separate the influence of interdiffusion and melt elasticity because both of them are affected by molecular weight. According to the reptation model, the self diffusion coefficient D_{self} is proportional to M^{-2} . However, the definition of D_{self} is based on the total displacement of center of mass of a chain or diffusion of the order of R_g of the chain¹. In fact, it was reported that the full strength at the interface could be developed by interdiffusion of the chains in the order of a fraction of R_g ($\approx 0.81R_g$)^{1,39}. Therefore the diffusion of long chains is sufficient to strengthen the interface as long as they are able to make entanglements on the other side of the interface. This length is a chain length of the order of the critical molecular weight of entanglements (M_c). For polyethylene, this molecular weight was reported to be $\approx 4000 \text{ g.mol}^{-1}$ ⁴⁰⁻⁴². Therefore, as presented in Figure 5-3 and Figure 5-4 for the resins with similar molecular architecture investigated in this study, the influence of melt elasticity is dominant.

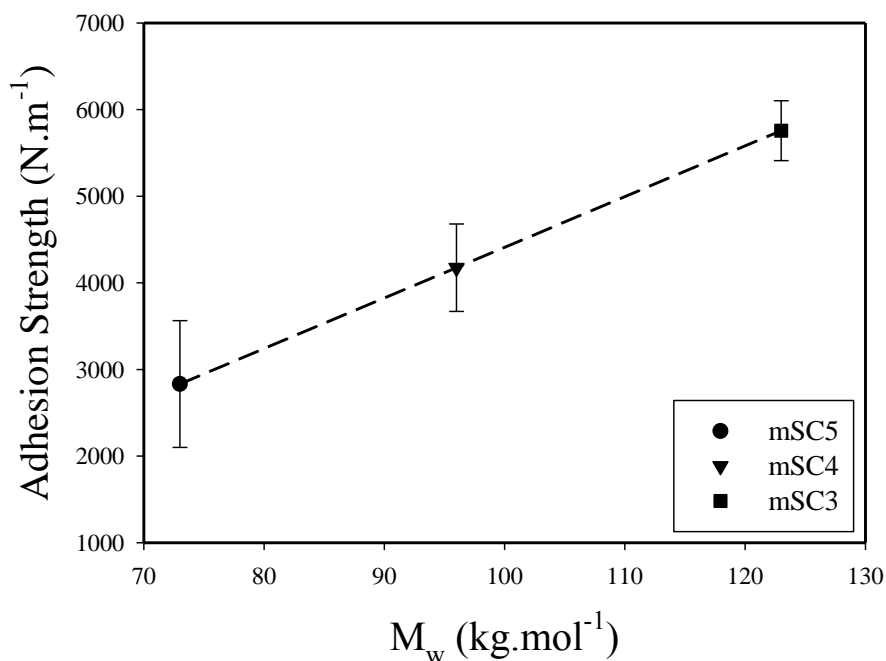


Figure 5-4 Linear dependence of maximum self adhesion strength of mSC3, mSC4, and mSC5 films to molecular weight

In the following sections, the focus will be on the influence of PEs chain structural characteristics such as long chain branches, molecular weight distribution and heterogeneity of composition distribution on adhesion strength. In order to eliminate the effect of molecular weight, polymers were selected from the same range of M_w . In addition, because of the slight differences in M_w and to exclude its potential minor effect, the adhesion strengths were normalized by dividing by M_w , since they are linearly related as shown above. The normalized ultimate adhesion strengths for these three resins overlapped at the value $\approx 0.048 \text{ N.m}^{-1}/\text{g.mol}^{-1}$. Therefore, dividing by M_w removed the gap between adhesion strengths of mSC3, mSC4, and mSC5. Following this result, the measured adhesion strength of polymers in this work would be normalized by their M_w .

5.4.3 Effect of long chain branching on self-adhesion strength

There are two kinds of long chain branch (LCB) containing polymers in the polyethylene family: Conventional LCB polymers (such as LDPE) synthesized by free-radical polymerization at high temperature and pressure which have very broad molecular weight distribution, heterogenous LCB length, and distribution; Sparsely long chain branch ethylene α -olefin copolymers with narrow MWD, homogenous branch composition, and distribution (such as metallocene ethylene α -olefin copolymers containing sparsely LCB). The advent of metallocene catalysts has offered this great opportunity for producing sparsely LCB content PEs. It improved the control of the final ethylene backbone and branching microstructure through synthesis. In this work, the effect of LCB on self-adhesion will be evaluated in these two categories.

As mentioned earlier, long-chain branching has a profound effect on melt rheology and processability of polyethylenes. Even at very low branching densities, such as 1 LCB/ 10^4 carbon atoms, melt elasticity and strain hardening was reported to be influenced by the presence of the long branches⁴³. This is due to the involvement of these side chains in entanglements and playing an anchor role in polymer melt.

As mentioned above, LCB densities were determined quantitatively (albeit averaged over all molecules present) using high temperature GPC. Table 5-4 shows LCB contents of LDPE, mLC1, and mLC2 determined by GPC. Generally, the presence of LCB changes the dynamics of chain diffusion from simple reptation to arm retraction, which retards the movement of chains along their backbone.

Table 5-5 LCB contents of long chain branch resins obtained from GPC

Nomenclature	LCB/ 10^4 C atom
LDPE	5.2
mLC1	0.19
mLC2	0.3

Figure 5-5 shows the comparison between interfacial adhesion strength of the LCB containing resins and their linear counterparts. The ultimate interfacial strength is $4.6 \times 10^{-2} \text{ N.m}^{-1}/\text{g.mol}^{-1}$ for mLC1 and $1.7 \times 10^{-2} \text{ N.m}^{-1}/\text{g.mol}^{-1}$ for mLC2, compared to $4.9 \times 10^{-2} \text{ N.m}^{-1}/\text{g.mol}^{-1}$ value for mSC3. Since they all have narrow MWD, this could be explained by the retarded diffusion caused by hindrance of reptation motion at branch points, which is more drastic for higher LCB content. It is surprising to see such a strong effect of LCB on adhesion considering that both of the samples have relatively low LCB densities.

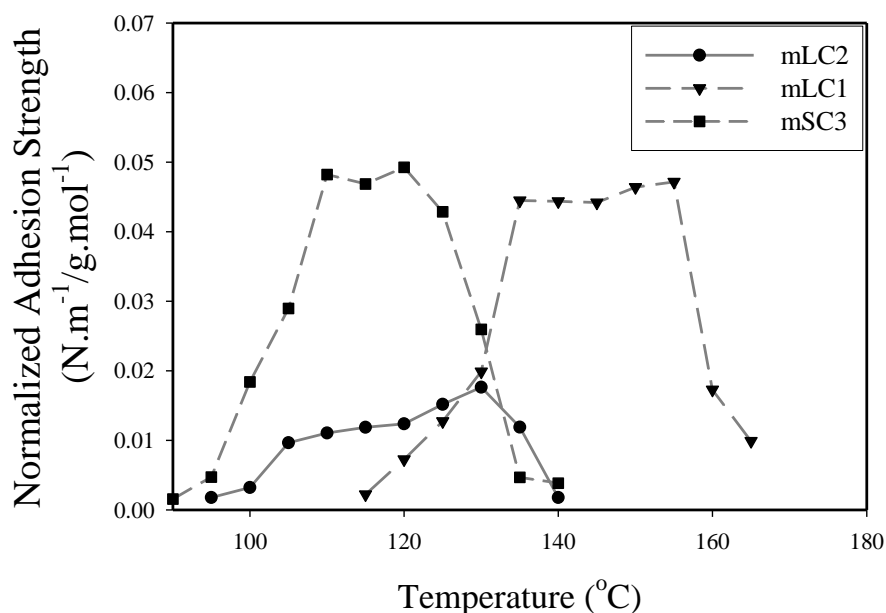


Figure 5-5 Self adhesion strength of mLC1, mLC2, and mSC3 films as a function of temperature obtained from T-peel tests with peel rate of 33 mm.s^{-1}

Figure 5-6 illustrates the comparison between adhesion of LDPE, LLDPE, and HDPE. For LDPE, the adhesion strength is lower than that obtained for LLDPE and HDPE. It can be clearly noted that the presence of large amount of LCB drastically diminished interdiffusion, which

resulted in drastic decrease in interfacial strength. Figure 5-7 represents the normalized adhesion strength of all three LCB containing resins in order to compare the magnitude of discrepancy between their adhesion strength. Moreover, the adhesion strength of these resins versus their LCB content is illustrated in Figure 5-8. The high level of irregular LCB in addition to broad MWD causes LDPE to have the very lowest level of adhesion strength. Moreover, this intrinsic molecular architecture of LDPE causes the separate investigation of the effects of MWD and LCB practically impossible.

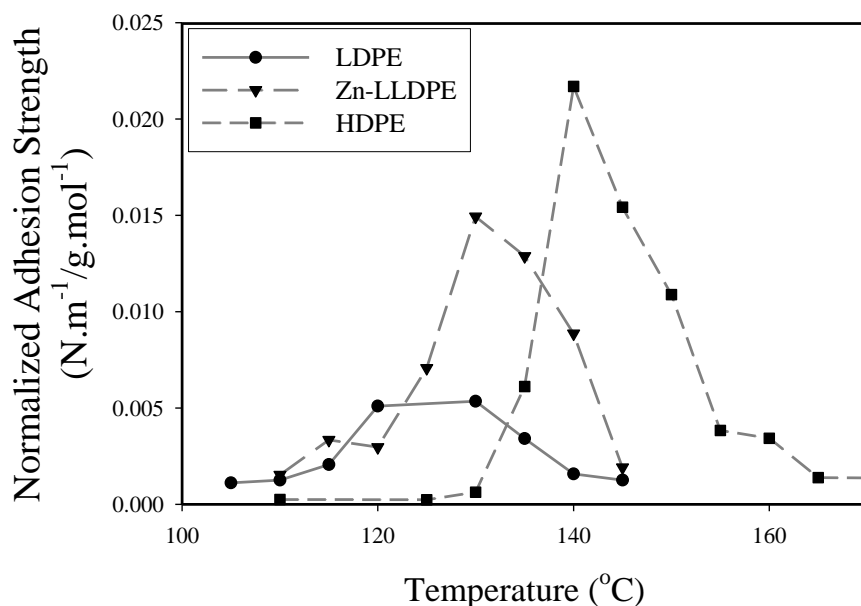


Figure 5-6 Self adhesion strength of LDPE, Zn-LLDPE, and HDPE films as a function of temperature obtained from T-peel tests with peel rate of 33 mm.s⁻¹

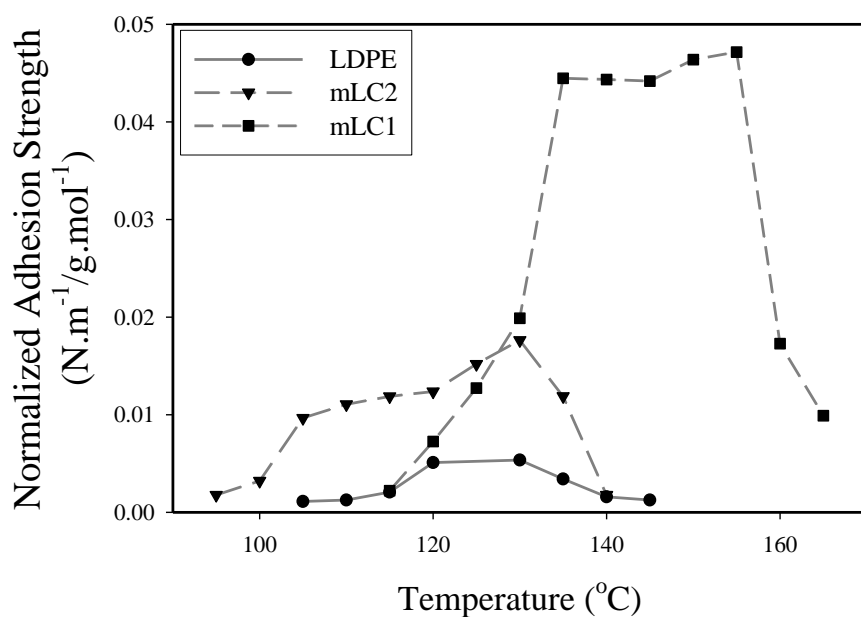


Figure 5-7 Self adhesion strength of films as a function of temperature obtained from T-peel tests with peel rate of 33 mm.s⁻¹

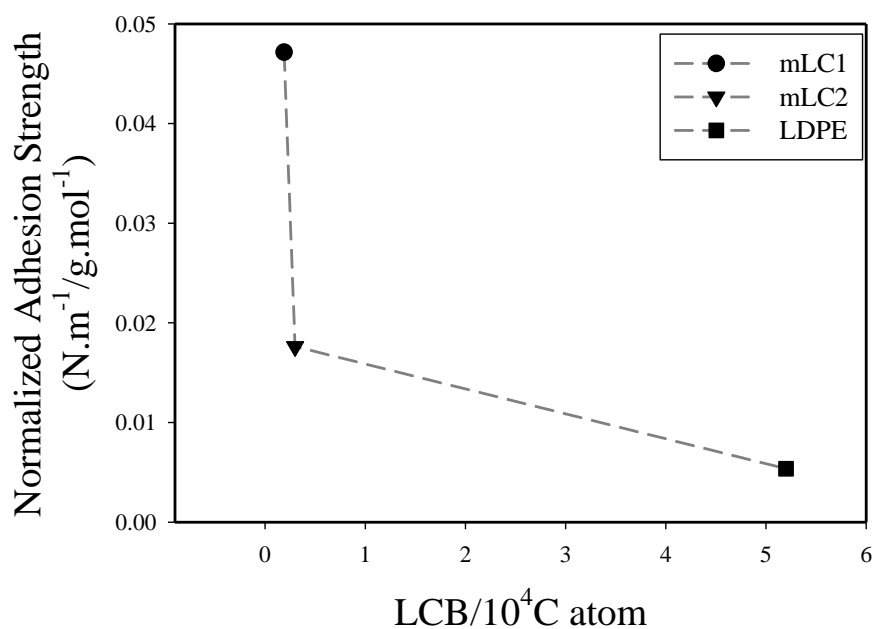


Figure 5-8 Normalized adhesion strength of versus LCB content of LDPE, mLC1, and mLC2

Figure 5-9 shows the elongational behaviour of LCB containing polymers compared to their linear counterparts, from both metallocene and conventional groups. Generally, presence of LCB

in polyethylene appears strikingly in extensional flow in the form of strain hardening behaviour. Strain-hardening in uniaxial extensional flow is seen for LDPE, mLC1, and mLC2. This differs qualitatively from the behaviour of un-branched or SCB melts. In Figure 5-9, LDPE which has multiple, irregularly spaced and long side branches, shows a sharper increase in the extensional viscosity curve and a deviation from linear viscoelasticity at a shorter time. The strain hardening behaviour of LCB resins causes high melt strength but it would not be of any help in increasing the interfacial adhesion strength. Because the adhesion at the interface happens sequentially, interdiffusion happens first and then peeling. The melt strength of polymer is determinant in peeling stage. Chain interdiffusion is pre-requisite for self-adhesion.

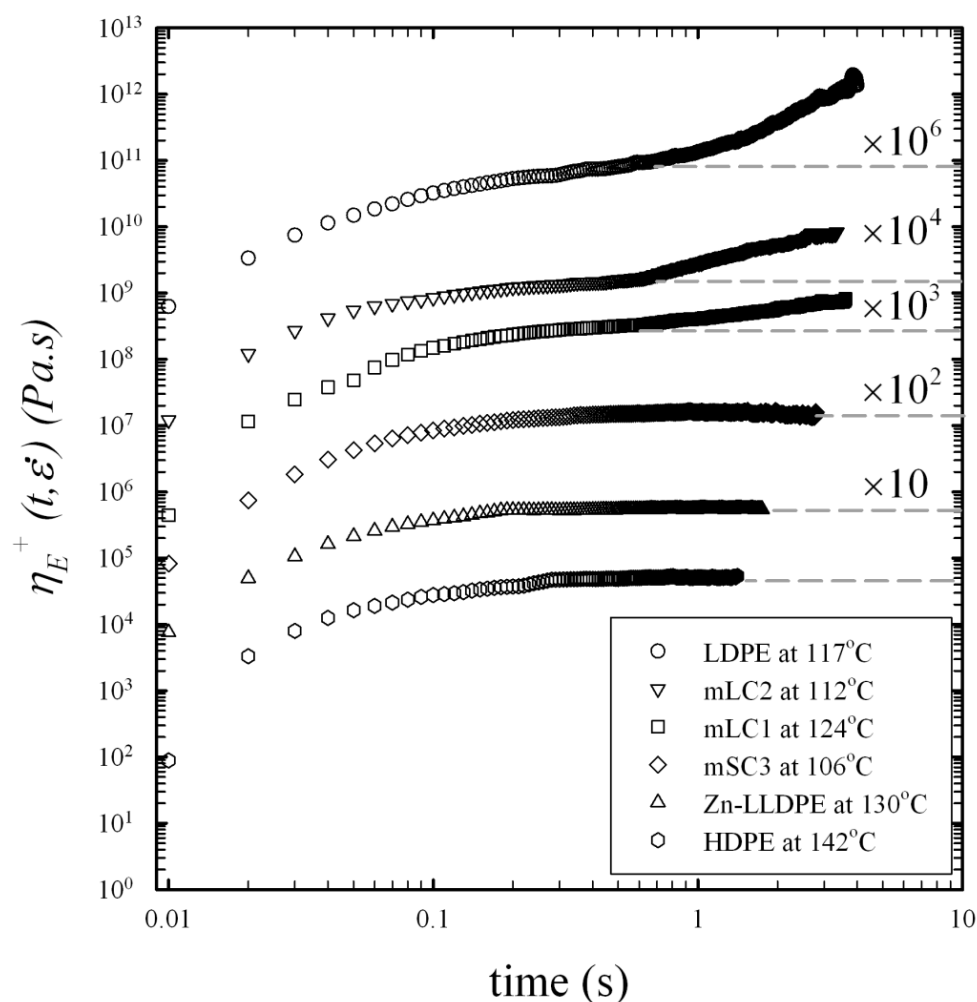


Figure 5-9 Transient elongational viscosities at Hencky strain rate $\dot{\epsilon}=1 \text{ s}^{-1}$, the dash-lines represent the linear behaviour where the transient elongational viscosity is equal to three times of

the zero shear viscosity in shear flow, (to facilitate the comparison between data, the curves have been shifted by a multiplication factor as indicated)

5.4.4 Effect of heterogeneity and homogeneity of molecule structure on self-adhesion strength

In this section, polymers were selected from the same range of M_w in order to eliminate the effect of molecular weight. In addition, to exclude any potential minor effects, the adhesion strength was normalized dividing by M_w as mentioned above. The interfacial adhesion strength of this group of linear metallocene ethylene α -olefins and conventional PEs is shown in Figure 5-10.

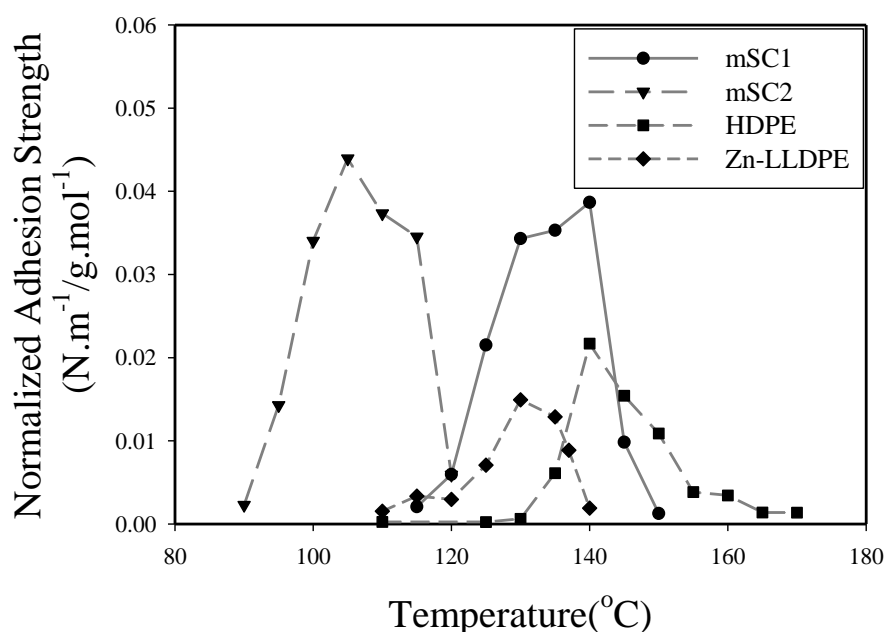


Figure 5-10 Self adhesion strength of films as a function of temperature obtained from T-peel tests with peel rate of 33 mm.s^{-1}

The normalized adhesion strength of mSC1 ($\approx 4.3 \times 10^{-2} \text{ N.m}^{-1}/\text{g.mol}^{-1}$) and mSC2 ($\approx 3.8 \times 10^{-2} \text{ N.m}^{-1}/\text{g.mol}^{-1}$) are much higher than HDPE ($\approx 2.1 \times 10^{-2} \text{ N.m}^{-1}/\text{g.mol}^{-1}$) and Zn-LLDPE ($\approx 9.1 \times 10^{-3} \text{ N.m}^{-1}/\text{g.mol}^{-1}$). Superior self adhesion of metallocene polyolefin polymers compared to conventional PEs has already been reported⁴⁴⁻⁴⁷. Also, as seen in Figure 5-11, none of the polymers have the advantage of higher elongational melt strength or strain hardening compared to the others.

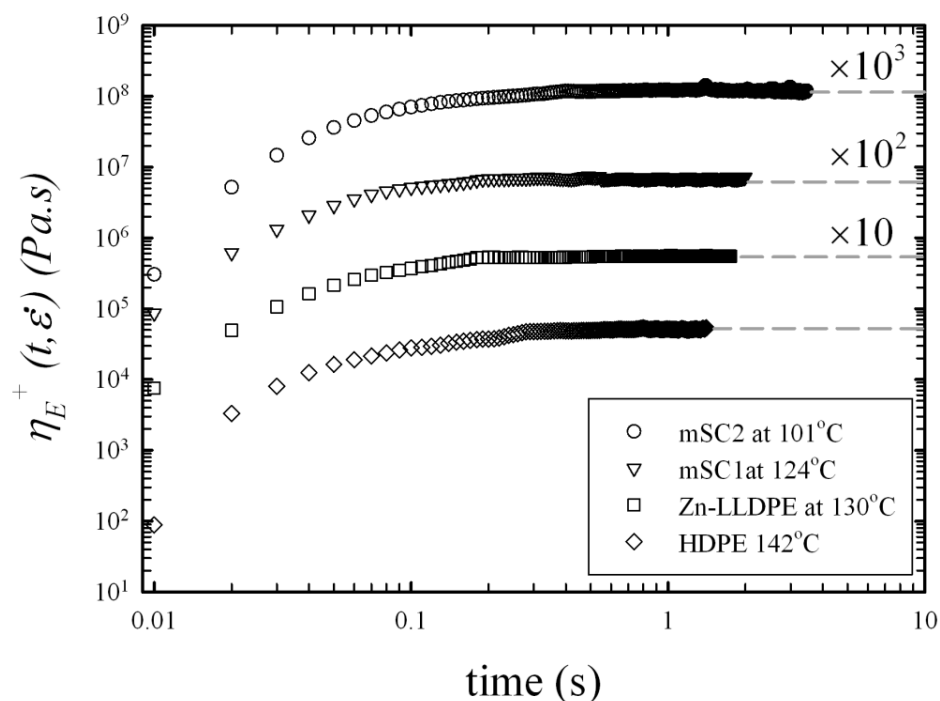


Figure 5-11 Transient elongational viscosities at Hencky strain rate $\dot{\epsilon}=1 \text{ s}^{-1}$, the dash-lines represent the linear behaviour where the transient elongational viscosity is equal to three times of the zero shear viscosity in shear flow, (to facilitate the comparison between data, the curves have been shifted by a multiplication factor as indicated)

Therefore, the dissimilarity in adhesion strength is related to the diffusion and the role of short chain branches and/or composition distribution. The amount of SCB in mSC2 and mSC1 is higher and lower than in LLDPE, respectively. However, they both show higher adhesion strength compared to Zn-LLDPE (See Figure 5-12). In addition, HDPE containing no SCB shows lower adhesion strength than metallocene PEs. Moreover, in the reptation dynamics of polymer chains, the role of short chain branches has been widely neglected^{31,48,49}. Therefore the influential parameter seems not to be the presence or amount of SCB but rather its distribution, and basically the composition distribution of polymer chains. Figure 5-13 shows the interfacial adhesion of Zn-LLDPE as well as mSC resins and HDPE as a function of polydispersity index (PDI).

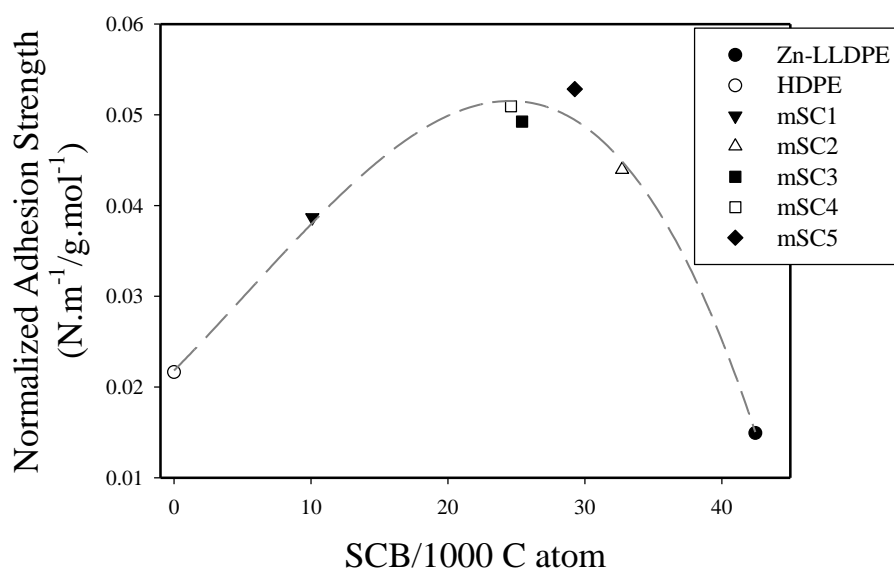


Figure 5-12 Normalized adhesion strength of LLDPEs and HDPE as a function of SCB content, assuming SCB content of HDPE equal to zero

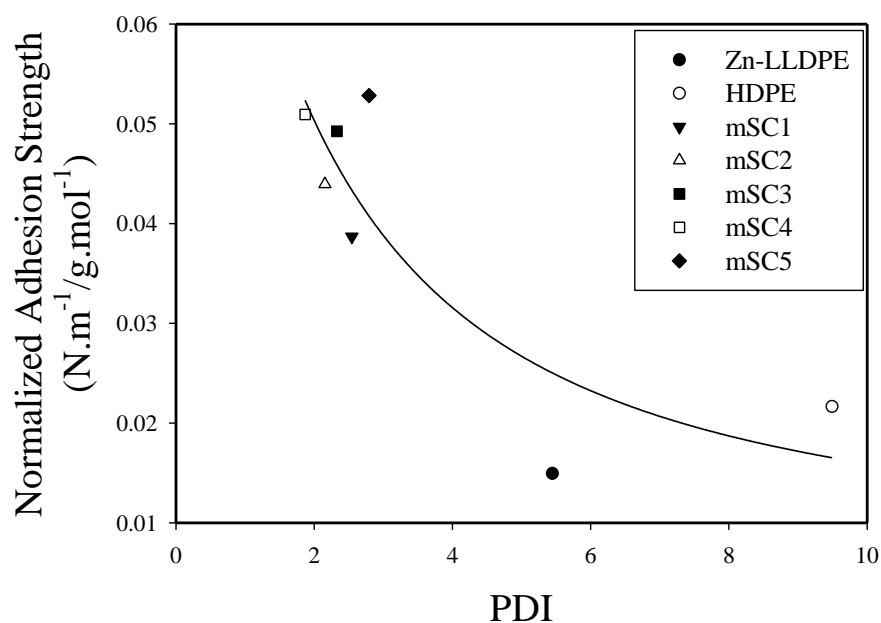


Figure 5-13 Normalized adhesion strength of LLDPEs and HDPE as a function of PDI

There are some strong indications in the literature for an enhanced concentration of low molecular weight chains at the interface with a low-energy surface, such as air or vacuum⁵⁰⁻⁵². Several researchers confirmed this segregation and the formation of an amorphous layer at the

surface. It has a composition dissimilar from bulk chain composition of the polymer and might form after its solidification from the melt or solution^{50,51,53}. This segregation of low M_w chains to the surface was explained thermodynamically by enthalpy and entropy driving forces^{38,50,51}. In very thin films prepared from model blends of polyolefins, dramatic surface enrichment by a highly branched component was confirmed^{54,55}. Also, migration of low molecular weight fractions of LDPE and PP to the surface was confirmed at their interface with an EVA phase⁵³. In another study, the amplitude and length scale of surface enrichment for variety of chain architectures was examined, and the role of degree of branching on surface enrichment was highlighted⁵¹.

For conventional polyethylene films, the presence of such a low molecular weight chain enriched surface layer was examined using atomic force microscopy^{50,56}. The layer thickness was reported to be 100 nm in Ziegler-Natta LLDPE⁵⁰. Employing the same AFM technique, such a segregated layer in metallocene copolymers with uniform CCD and narrow MWD was not found. In polymers with a homogenous structure, the same densely packed lamellar morphology as the bulk was reported to exist at the interface⁵⁰.

As discussed earlier, composition distribution includes molecular weight distribution and short chain branch distribution among and along PE chains. Non-uniform intermolecular and intramolecular CCD in Zn-LLDPE is due to the concentrated SCB in the low molecular weight chains. Therefore, in the case of Zn-LLDPE the segregated layer in the surface consists of highly branched low molecular weight chains. The low self-adhesion of LDPE also could be attributed to this segregated layer, in addition to the presence of LCB in LDPE chains. In the case of homopolymers (such as HDPE) which have linear chains but broad MWD, the segregated layer consists just of low molecular weight chains^{51,52}.

In interfacial adhesion of polymers, the strength development at the interface is the result of chain diffusion across the interface. The chains that diffuse across the interface could make entanglements and make bridges between the two film sides. These bridges act as stitches attaching them. Diffusion of low molecular weight chains is known to occur in shorter time scale than for longer chains, so they diffuse first⁵⁷. However, they would not build a strong adhesion^{3,32}. As mentioned previously, the high molecular weight chains are more effective in interfacial adhesion than low molecular weight chains³¹.

In the case of heterogeneous PEs, the influence of segregated low molecular weight chains in interfacial adhesion is that they hinder creating a strong interface. As illustrated in Figure 5-14, long chains in the bulk have to pass through the segregated layer in the surface (which is doubled upon sealing) and reach the long chains in the bulk of the other side of the interface. In other words, it takes longer time for high molecular weight fractions to become involved in interdiffusion. Investigating the kinetics of adhesion in heterogeneous ethylene α -olefins confirmed that the time dependence of interfacial strength was two orders of magnitude longer than the expected time for chains to make entanglement ³⁸. In contrast, for metallocene ethylene α -olefins, the melt adhesion reached the maximum strength instantaneously in the experimental time scale ⁵⁰. Moreover, in self adhesion of very narrow MWD polystyrene, no delay was reported in diffusion across the interface ⁵⁸.

As mentioned, in reptation dynamics of polymer chains, the role of short chain branches has been widely neglected ^{31,48,49}. However, a few studies reported that the occasional concentration of branches in one portion of the chain can act as an anchor to increase local friction and hamper reptation ⁵⁹. This could explain the poor adhesion results of Zn-LLDPE compared to HDPE.

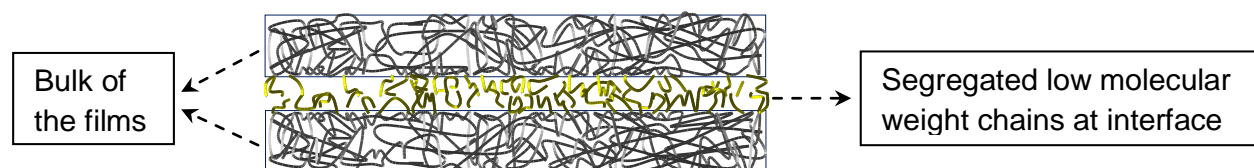


Figure 5-14 Schematic of the cross section of a polydisperse polymer in molecular scale showing the segregated layer of low molecular weight chains at interface

The focus of this study was to measure the self adhesion of polyethylene films while they are still molten and hot. However, it should be mentioned that, even if it is a slight possibility, strain induced crystal formation could not be disregarded. Chain orientation is the origin of strain induced crystallization at high strain rates. Due to the limitations in present techniques, it was not possible to verify the occurrence of this crystallization during the peel test. However, the hypothesis of a segregated layer at the interface weakens the probability of crystal formation in the peel test. The presence of highly SCB concentrated low molecular weight chains at the interface will hinder crystallization and would not undergo high orientation (due to their very low relaxation time). In metallocene PEs of this study, high SCB content in entire polymer chains and low X_C could also undermine the chances for crystallization.

5.5 Conclusions

In this work, the influence of chain architecture on interfacial self-adhesion of polyethylene above melting temperature was investigated. It was particularly pointed out that the level of interfacial self-adhesion strength is the outcome of chain interdiffusion across the interface and melt strength of polymer. The following conclusions can be drawn:

1. For stereo regular and narrow molecular weight distribution metallocene ethylene α -olefins containing equal amount of SCB, higher adhesion strength was observed for the polymer with higher Mw. Having all the other characteristics similar, a linear dependence of adhesion strength to Mw was observed. Interfacial self-adhesion at temperatures higher than the melting point was closely related to their melt elasticity.
2. The presence of even low amount of long chain branches hinders the reptation motion and diffusion, and results in lower adhesion force in metallocene ethylene α -olefins. In addition, highly branched chains cause LDPE to have very low self-adhesion.
3. Heterogeneity in composition distribution of conventional polyethylene results in poor self-adhesion performance. In these resins long chains in the bulk have to pass through the segregated layer at the surface, which is doubled upon sealing, and reach the long chains in bulk of the other side of the interface, in order to make entanglements and strengthen the interface.

5.6 Acknowledgement

We would like to thank A. Arnold in Université du Québec à Montréal (UQAM) for performing NMR experiments, his excellent technical assistance, and helpful advices on NMR analysis. Financial support from 3S Pack NSERC/Saputo/Prolamina industrial research chair is gratefully acknowledged.

5.7 References

1. Kim, K. D., Sperling, L. H., Klein, A. & Hammouda, B. Reptation Time, Temperature, and Cosurfactant Effects on the Molecular Interdiffusion Rate during Polystyrene Latex Film Formation. *Macromolecules* **27**, 6841–6850 (1994).

2. Wool, R. P., Yuan, B.-L. & McGarel, O. J. Welding of polymer interfaces. *Polymer Engineering & Science* **29**, 1340–1367 (1989).
3. Kausch, H. H., Nguyen, T. Q. & Petrovska-Delacrétaz, D. Chain interdiffusion: macromolecules between Rouse and de Gennes. *Phys. Scr.* **1991**, 57 (1991).
4. Wool, R. P. & O'Connor, K. M. A theory crack healing in polymers. *Journal of Applied Physics* **52**, 5953–5963 (1981).
5. Jud, K., Kausch, H. H. & Williams, J. G. Fracture mechanics studies of crack healing and welding of polymers. *J Mater Sci* **16**, 204–210 (1981).
6. Prager, S. & Tirrell, M. The healing process at polymer–polymer interfaces. *The Journal of Chemical Physics* **75**, 5194–5198 (1981).
7. Kausch, H. H. & Tirrell, M. Polymer Interdiffusion. *Annual Review of Materials Science* **19**, 341–377 (1989).
8. Wool, R. P. *Polymer Interfaces: Structure and Strength*. (Hanser-Gardner Publications, 1995).
9. Schach, R. & Creton, C. Adhesion at interfaces between highly entangled polymer melts. *Journal of Rheology* **52**, 749–767 (2008).
10. Boiko, Y. M., Guérin, G., Marikhin, V. A. & Prud'homme, R. E. Healing of interfaces of amorphous and semi-crystalline poly(ethylene terephthalate) in the vicinity of the glass transition temperature. *Polymer* **42**, 8695–8702 (2001).
11. Gent, A. N., Kim, E.-G. & Ye, P. Autohesion of crosslinked polyethylene. *Journal of Polymer Science Part B: Polymer Physics* **35**, 615–622 (1997).
12. Smith, G. D., Plummer, C. J. G., Bourban, P.-E. & Manson, J.-A. E. Non-isothermal fusion bonding of polypropylene. *Polymer* **42**, 6247–6257 (2001).
13. Xue, Y.-Q., Tervoort, T. A. & Lemstra, P. J. Welding behavior of semicrystalline polymers. 1. The effect of nonequilibrium chain conformations on autoadhesion of UHMWPE. *Macromolecules* **31**, 3075–3080 (1998).
14. Xue, Y.-Q., Tervoort, T. A., Rastogi, S. & Lemstra, J. Welding Behavior of Semicrystalline Polymers. 2. Effect of Cocrystallization on Autoadhesion. *Macromolecules* **33**, 7084–7087 (2000).
15. Wunderlich, B. & Czornyj, G. A Study of Equilibrium Melting of Polyethylene. *Macromolecules* **10**, 906–913 (1977).
16. Randall, J. C. A Review of High Resolution Liquid ¹³carbon Nuclear Magnetic Resonance Characterizations of Ethylene-Based Polymers. *Journal of Macromolecular Science, Part C: Polymer Reviews* **29**, 201–317 (1989).
17. Zimm, B. H. & Stockmayer, W. H. The Dimensions of Chain Molecules Containing Branches and Rings. *The Journal of Chemical Physics* **17**, 1301–1314 (2004).
18. Zimm, B. H. & Kilb, R. W. Dynamics of branched polymer molecules in dilute solution. *Journal of Polymer Science* **37**, 19–42 (1959).

19. Yu, Y., DesLauriers, P. J. & Rohlffing, D. C. SEC-MALS method for the determination of long-chain branching and long-chain branching distribution in polyethylene. *Polymer* **46**, 5165–5182 (2005).
20. Hsieh, E. T. & Randall, J. C. Monomer sequence distributions in ethylene-1-hexene copolymers. *Macromolecules* **15**, 1402–1406 (1982).
21. RANDALL J. C. & HSIEH E. T. in *NMR and Macromolecules* **247**, 131–151 (American Chemical Society, 1984).
22. D20 Committee. *Test Method for Determination of Linear Low Density Polyethylene (LLDPE) Composition by Carbon-13 Nuclear Magnetic Resonance*. (ASTM International, 2009).
23. Bensason, S. *et al.* Classification of homogeneous ethylene-octene copolymers based on comonomer content. *Journal of Polymer Science Part B: Polymer Physics* **34**, 1301–1315 (1996).
24. Bovey, F. A., Schilling, F. C., McCrackin, F. L. & Wagner, H. L. Short-Chain and Long-Chain Branching in Low-Density Polyethylene. *Macromolecules* **9**, 76–80 (1976).
25. Klimke, K. *et al.* Optimisation and Application of Polyolefin Branch Quantification by Melt-State ¹³C NMR Spectroscopy. *Macromolecular Chemistry and Physics* **207**, 382–395 (2006).
26. Pollard, M. *et al.* Observation of Chain Branching in Polyethylene in the Solid State and Melt via ¹³C NMR Spectroscopy and Melt NMR Relaxation Time Measurements. *Macromolecules* **37**, 813–825 (2004).
27. Shroff, R. N. & Mavridis, H. Assessment of NMR and Rheology for the Characterization of LCB in Essentially Linear Polyethylenes. *Macromolecules* **34**, 7362–7367 (2001).
28. Bubeck, R. A. Structure–property relationships in metallocene polyethylenes. *Materials Science and Engineering: R: Reports* **39**, 1–28 (2002).
29. Hosoda, S. *et al.* Effect of the Sequence Length Distribution on the Lamellar Crystal Thickness and Thickness Distribution of Polyethylene: Perfectly Equisquential ADMET Polyethylene vs Ethylene/ α -Olefin Copolymer. *Macromolecules* **44**, 313–319 (2011).
30. Hung, J., Cole, A. P. & Waymouth, R. M. Control of Sequence Distribution of Ethylene Copolymers: Influence of Comonomer Sequence on the Melting Behavior of Ethylene Copolymers. *Macromolecules* **36**, 2454–2463 (2003).
31. Roland, C. M. & Boehm, G. G. A. Macromolecular diffusion and the autoadhesion of polybutadiene. *Macromolecules* **18**, 1310–1314 (1985).
32. Su, G. M., Best, K., Ranganathan, T., Emrick, T. & Crosby, A. J. Tailored Nanoparticles for Enhancing Polymer Adhesion. *Macromolecules* **44**, 5256–5261 (2011).
33. Léger, L. & Creton, C. Adhesion mechanisms at soft polymer interfaces. *Phil. Trans. R. Soc. A* **366**, 1425–1442 (2008).
34. Micic, P. & Bhattacharya, S. N. Rheology of LLDPE, LDPE and LLDPE/LDPE blends and its relevance to the film blowing process. *Polymer international* **49**, 1580–1589 (2000).

35. Ghijssels, A., Ente, J. & Raadsen, J. Melt Strength Behavior of PE and its Relation to Bubble Stability in Film Blowing. *International Polymer Processing* **5**, 284–286 (1990).
36. La Mantia, F. P. & Acierno, D. Influence of the molecular structure on the melt strength and extensibility of polyethylenes. *Polymer Engineering & Science* **25**, 279–283 (1985).
37. Brown, H. R. The Adhesion Between Polymers. *Annual Review of Materials Science* **21**, 463–489 (1991).
38. Qureshi, N. Z., Stepanov, E. V., Capaccio, G., Hiltner, A. & Baer, E. Self-adhesion of polyethylene in the melt. 1. Heterogeneous copolymers. *Macromolecules* **34**, 1358–1364 (2001).
39. Wool, R. P. Adhesion at polymer–polymer interfaces: a rigidity percolation approach. *Comptes Rendus Chimie* **9**, 25–44 (2006).
40. Wool, R. P. Self-healing materials: a review. *Soft Matter* **4**, 400–418 (2008).
41. Vega, J. F., Rastogi, S., Peters, G. W. M. & Meijer, H. E. H. Rheology and reptation of linear polymers. Ultrahigh molecular weight chain dynamics in the melt. *Journal of Rheology (1978-present)* **48**, 663–678 (2004).
42. Fetters, L. J., Lohse, D. J. & Colby, R. H. in *Physical Properties of Polymers Handbook*
43. Wang, W.-J., Yan, D., Zhu, S. & Hamielec, A. E. Kinetics of Long Chain Branching in Continuous Solution Polymerization of Ethylene Using Constrained Geometry Metallocene. *Macromolecules* **31**, 8677–8683 (1998).
44. Halle, R. W. & Davis, D. S. Heat sealing linear ethylene plastomers to ionomers or LLDPEs. *Tappi journal* **78**, 200–206 (1995).
45. Farley, J. M., Meka, P., Stehling, F. C., Trudell, B. C. & Kurtzman, M. B. United States Patent: 5530065 - Heat sealable films and articles made therefrom. (1996).
46. Shih, H. H., Wong, C. M., Wang, Y. C., Huang, C. J. & Wu, C. C. Hot tack of metallocene catalyzed polyethylene and low-density polyethylene blend. *Journal of Applied Polymer Science* **73**, 1769–1773 (1999).
47. Majumdar, A. & Kale, D. D. Properties of films made from ternary blends of metallocene and conventional polyolefins. *Journal of Applied Polymer Science* **81**, 53–57 (2001).
48. Von Seggern, J., Klotz, S. & Cantow, H. J. Reptation and constraint release in linear polymer melts: an experimental study. *Macromolecules* **24**, 3300–3303 (1991).
49. Pearson, D. S., Fetters, L. J., Graessley, W. W., Ver Strate, G. & von Meerwall, E. Viscosity and self-diffusion coefficient of hydrogenated polybutadiene. *Macromolecules* **27**, 711–719 (1994).
50. Qureshi, N. Z. *et al.* Self-adhesion of polyethylene in the melt. 2. Comparison of heterogeneous and homogeneous copolymers. *Macromolecules* **34**, 3007–3017 (2001).
51. Wu, D. T. & Fredrickson, G. H. Effect of Architecture in the Surface Segregation of Polymer Blends. *Macromolecules* **29**, 7919–7930 (1996).
52. Jones, R. A. & Richards, R. W. *Polymers at surfaces and interfaces*. (Cambridge University Press, 1999).

53. McEvoy, R. L. & Krause, S. Interfacial Interactions between Polyethylene and Polypropylene and Some Ethylene-Containing Copolymers. *Macromolecules* **29**, 4258–4266 (1996).
54. Scheffold, F. *et al.* Surface phase behavior in binary polymer mixtures. II. Surface enrichment from polyolefin blends. *The Journal of Chemical Physics* **104**, 8795–8806 (1996).
55. Brant, P., Karim, A., Douglas, J. F. & Bates, F. S. Surface Composition of Amorphous and Crystallizable Polyethylene Blends As Measured by Static SIMS. *Macromolecules* **29**, 5628–5634 (1996).
56. Magonov, S. & Godovsky, Y. Atomic force microscopy, Part 8: Visualization of granular nanostructure in crystalline polymers. *American Laboratory* **31**, 52–58 (1999).
57. Schuman, T. *et al.* Solid state structure and melting behavior of interdiffused polyethylenes in microlayers. *Polymer* **40**, 7373–7385 (1999).
58. Reiter, G. & Steiner, U. in *Application of Scattering Methods to the Dynamics of Polymer Systems* (Ewen, B., Fischer, E. W. & Fytas, G.) 93–96 (Steinkopff, 1993).
59. Schuman, T. *et al.* Interdiffusion of linear and branched polyethylene in microlayers studied via melting behavior. *Macromolecules* **31**, 4551–4561 (1998).

CHAPTER 6

**ARTICLE 3: INTERFACIAL SELF-ADHESION OF POLYETHYLENE
BLENDS: THE ROLE OF LONG CHAIN BRANCHING AND
EXTENSIONAL RHEOLOGY***

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6.1 Abstract

The interfacial self-adhesion strength of polyethylene binary blends in temperatures higher than T_m was investigated in relation to their components' structures and elongational rheological properties. Four binary blends were prepared from the combination of a conventional polyethylene; LDPE or Ziegler-Natta LLDPE, with a metallocene catalyzed ethylene α -olefin copolymer; linear (Linear-m) or long chain branched (LCB-m). Differential scanning calorimetry (DSC) melting curve of the blend films showed the gradual shift of T_m and partial cocrystallization for all blend compositions, which influenced the adhesion temperature. Interfacial bonding was carried out through intimate contact under slight pressure and heat for 0.5 s. Adhesion strength was measured by peeling immediately after sealing, while the adherents were still in molten state. Incorporation of a metallocene, of both kinds, into conventional PEs enhanced the self adhesion. The adhesion strength of the blends containing 60wt% metallocene resins was far lower than the superior adhesion strength of neat metallocene resins. This was due to the formation of a segregated layer of highly branched short chains of LDPE or LLDPE at the surface of films. Blending with Linear-m caused higher enhancement in adhesion strength than blending with LCB-m. This was attributed to the faster reptation of linear chains, hence superior diffusion across the interface. The larger increase of adhesion strength was observed for all compositions of LDPE blends compared to their LLDPE counterparts. This suggested that for LDPE blends, a fast diffusion was provided by both Linear-m and LCB-m. The upper and lower limits of the temperature range in which the film showed a plateau of its highest adhesion strength were determined for all compositions. The temperature window of plateau adhesion

* Submitted to Rheologica Acta, April 2014.

strength for LDPE blends was broader than their Zn-LLDPE counterparts. This work suggests that the final plateau temperature (T_{pf}) can be correlated with the area under stress-strain curve of extensional rheological measurements. Also, the higher melt toughness can result in a broader adhesion strength plateau.

6.2 Introduction

Self-adhesion in glassy or semicrystalline polymers is created by heating the materials above their glass transition or melting temperature to permit interdiffusion and is referred to as welding, sealing or crack healing. This area of melt adhesion strength is an ongoing topic in fundamental polymer research. So far, the majority of studies on self-adhesion of either semicrystalline or glassy polymers have been concentrated on the adhesion strength after cooling¹⁻⁴. However, in some applications, the strength of the joint is critical while it is still molten and under tension. For example in modern machinery of flexible packaging industry where the speed must be as high as possible, the heated joint should resist the stress caused by the weight of the product inside the packaged. The strength of an interface is estimated as the required force when separating the adherents, while they are still molten and hot. The melt strength of the adherent is critical as well as its ability for interdiffusion at the interface⁵.

In the case of semicrystalline polymers, the focus has been on measuring the interfacial adhesion after heated bonding at room temperature where crystallization occur⁶⁻¹³. One of the main reasons for limited fundamental studies on self adhesion in the melt state is that the measurement of interfacial adhesion between two polymer melts is challenging. Therefore, there is less information on interfacial adhesion of semicrystalline polymers at temperatures higher than the melting point.

The main distinction between adhesion measurement before and after cooling is not the joining mechanism but in peeling and fracture mechanics. Fracture toughness is controlled by the local yielding mechanisms and the rheological properties of the materials, and thus tends to differ significantly before and after cooling^{6,8,14}.

The use of single-site catalyst technology for the polymerization of olefins allows the efficient control of the molecular structure and led to the formation of polyethylene chains with narrow MWD and relatively homogeneous comonomer distributions among and along chains. This novel

technology has been particularly useful for more uniform α -olefin comonomer incorporation in ethylene chains than it is with conventional Ziegler-Natta catalysts. This technology offered also the possibility for the production of metallocene catalyzed ethylene olefins with controlled long chain branches ¹⁵.

In a previous work ¹⁶, the influence of molecular architecture such as molecular weight (M_w), molecular weight distribution (MWD), amount and distribution of long chain branch (LCB), and short chain branch (SCB) distribution among and along polyethylene chains on interfacial self-adhesion at temperatures higher than melting point was studied. However, because of economical concerns and difficulties in metallocene extrusion and film processing, they have been used in blends with conventional PEs. In addition, the improved toughness, clarity, mechanical properties and down gauging are the advantages of these blends. Metallocene resins have been the subject of extensive studies on thermal and rheological properties as well as miscibility and processability. Several reports investigated the miscibility/immiscibility of LDPE/m-LLDPE and Zn-LLDPE/m-LLDPE blends ¹⁷⁻²³. The main outcome of these studies was that the blend properties depend on M_w , MWD, blend composition, comonomer type and the content of each component. Due to the broad diversity of these parameters in PEs, a universal conclusion might not be accurate. However, the majority of these studies reported the miscibility in the melt state and partial miscibility with the probability of partial co-crystallization in solid state ¹⁷⁻²³.

To our knowledge, there is no study on interfacial adhesion of polyethylene blends at temperatures higher than the melting point. In this study, hot tack equipment was used in order to seal film specimens above the melting point and measure peeling force immediately after sealing. This technique provided the possibility to measure the adhesion strength in the first milliseconds of peeling. This creates a unique opportunity to study self adhesion at very short time, which is very limited in literature.

In this study, two metallocene ethylene α -olefin copolymers were selected. To investigate the role of sparse LCB in metallocene chains in the blends with conventional PEs, one contains a low level of LCB while its counterpart has no LCB. They were individually blended with Zn-LLDPE and LDPE at 20, 40 and 60wt%. Accordingly, by producing four types of blends we investigated the presence of LCB based on branch distribution and content in this work. The adhesion strength experiments were discussed with regard to the melting behaviour and melting temperature which

control adhesion temperature. The correlation between the adhesion strength of the blends and the extensional rheology was also explored at different temperatures.

Results of this study bring new insights about the behaviour of polyethylene blends in interfacial self-adhesion at molten stage and under tension. This is of particular interest for modern machinery of high speed flexible packaging.

6.3 Experiments

6.3.1 Materials and films preparation

Conventional LDPE and LLDPE resins and the commercial metallocene, linear and containing sparse LCB ethylene α -olefin copolymers, were supplied by ExxonMobil Company. Table 6-1 shows the main characteristics of these resins. Films from blends of conventional/metallocene polyethylene resins containing 20, 40, and 60wt% of metallocene resins were prepared using a cast film line. A 45mm Killion single screw extruder followed by 2 mm opening and 20 cm width slit cast die was used. Films of 50 μ m in thickness were prepared from the entire blend compositions. The temperature profile along the barrel (from hopper to die) was set at 160/180/190/200/200°C. The extrusion was carried out at 40 rpm and the distance between the die exit to the nip roll was 10 cm. An air knife and calendar cooling system were used to cool the films after the die exit. Extrusion speed and the collecting speed were kept the same for all resins in order to control draw ratio and have uniform final thickness.

Table 6-1 Main characteristics of the resins

Nomenclature	MFI ^a (190°C/2.16kg) (g/10min)	α -olefin comonomer ^a	SCB		Density ^a (g.cm ⁻³)	T _m ^a (°C)	M _w ^c (kg.mol ⁻¹)	PDI ^c
			content/ 1000C	atom ^b				
LDPE	2	-	-		0.923	110	160	8.756
Zn-LLDPE	2	butene	42.45		0.918	123	133	5.448
Linear-m	1.2	hexene	25.41		0.900	96	123	2.329
LCB-m	0.5	hexene	24.12		0.920	114	115	2.636

^a Provided by the manufacturer. ^b Obtained from high temperature ¹³C NMR. ^c Obtained from high temperature GPC.

6.3.2 Differential scanning calorimetry (DSC)

In order to determine melting temperature of the films of various blend compositions, differential scanning calorimetry (DSC) was used with the help of a TA Instruments Q1000 calorimeter. The samples were heated from room temperature to 200°C at a heating rate of 10°C/min. The melting temperatures of films were determined from the first heating ramp.

6.3.3 Extensional rheology

The melt extensional behaviour of the polymers was determined using the SER geometry of the ARES Rheometric Scientific rheometer. The data were recorded at a Hencky strain rate of $\dot{\epsilon}=0.1$, 1 and 10 s⁻¹ on samples having dimensions of 13mm×18mm×0.8mm at 150°C. In order to study the role of molecular structure on melt strength and strain hardening, the temperature was kept constant for all extensional rheometry experiments. The temperature of 150°C was selected to make sure all the resins could melt in the time scale of experiment. The melt toughness of resins was calculated from the area under stress-strain curve of extentional flow at $\dot{\epsilon}=10$ s⁻¹ at three temperatures of 125°C, 135°C, and 150°C.

6.3.4 Interfacial self-adhesion measurements

Adhesion experiments were performed using SL10 Lako-Tool equipment based on T-Peel test method. Heat bonding was made with two metallic jaws of 19.1mm×25.4mm covered by Teflon.

The film samples were cut in $2.54\text{cm} \times 33\text{cm}$ strips specimens parallel to machine direction. This method is consisted of two stages of bonding and peeling: in the first stage, two layers of film were located between the two jaws, then the jaws closed to provide the contact between the two sides of the film with a controlled pressure for 0.5 s. In the second stage, the jaws were removed and simultaneously two arms of the machine pulled the films to measure the adhesion strength, while it was still hot (also called hot tack). To avoid film distortion due to direct contact to heated jaw, the PE films were laminated to a layer of PET support film using a polyurethane based solventless adhesive. To obtain reliable results for adhesion in the molten state, the PE films must be strongly attached to the support film. The adhesion experiments were performed in a range of jaw temperatures from 80°C to 160°C at a constant pressure of 0.5 MPa and dwell time of 0.5 s. The values reported in adhesion graphs were averaged over at least five tests.

6.4 Results and discussion

6.4.1 Differential scanning calorimetry (DSC)

In the heat bonding of thermoplastic films, the primary requirement for chain diffusion across the interface is the capability of polymer chains to reptate. The chain reptation motion in semicrystalline polymers is possible when the chains are not stocked in crystals. It is known that the number of chains capable of diffusion is determined by the fraction of amorphous phase in LLDPE films ²⁴. The un-melted crystals however act like obstacles hindering chains' free diffusion across the interface, thus the crystal content of films is important. Therefore, in the self-adhesion of thermoplastic films, the temperature in which adhesion is done is determined by the melting temperature of the film. Especially, for blends of two thermoplastic materials because of the different melting points of components, the adhesion temperature might vary based on the melting behaviour of every component. The size of crystals and the possibility of co-crystallization during the cooling of polymer films after extrusion influence the melting behaviour of film in heat bonding. Therefore, thermal characterization was performed to investigate the melting behaviour of PE blend films.

Figure 6-1 illustrates DSC melting endotherm curves of the first heating run for films of the neat polymers and their corresponding blends. The neat LLDPE showed higher melting temperature (T_m) than Linear-m and LCB-m blends. (See Figure 6-1a and b) The T_m shifted to lower

temperatures as the amount of Linear-m and LCB-m increased in the blend. The fact that the melting peaks in LL/Linear-m blends were more similar to that of neat LLDPE than neat Linear-m indicated weak miscibility in crystalline phase. Also, the gradual decrease of T_m in blends represents a partial miscibility of metallocene chains in LL rich crystalline phase. Linear-m showed the lowest T_m , which was attributed to its high number of short chain branches and homogeneous distribution of these branches along and among its chains (See Table 6-1).

In LL/Linear-m blends, although the difference between the T_m of the two neat resins was large, a distinct peak in 40wt% and 60wt% Linear-m was not observed. Only the LLDPE corresponding melting peak was observed and became broader as it shifted to the lower temperatures. This broad endotherm was attributed to the distribution of lamellar thicknesses and might be due to cocrystallization of LLDPE and Linear-m.

In the DSC melting thermogram of certain types of LLDPE, the presence of two or more distinct peaks was reported^{25, 26}. This was explained by heterogeneity in short chain branch distribution in Ziegler-Natta LLDPE chains which was due to the presence of different chain fractions. A fraction rich in comparatively linear chains is capable of creating larger lamellar crystals, while another fraction concentrated in SCB, which do not contribute in crystalline phase, results in lower crystallinity and smaller lamellae²⁵⁻²⁷. The melting curve of LLDPE, in addition to its main melting peak, showed a distinct shoulder indicating another population of the crystals. (See Figure 6-1)

In the melting endotherm of the blends in Figure 6-1a, the fraction of LL consisted of linear chains with the high T_m which were only slightly affected by blending with Linear-m. By blending LL with linear-m the small shoulder beside the main melting peak of LL was broadened and shifted to lower temperatures. Latter showed that the possible co-crystallization occurred mostly among the highly branched fraction of LLDPE and Linear-m chains. Knowing that the comonomer branches do not contribute into the crystal structure, the sequence and content of short chain branches in ethylene α -olefin copolymers is determinant in subsequent miscibility/immiscibility in crystalline phase of their blends^{23, 28-31}.

As mentioned earlier, in heat bonding of semicrystalline polymers, the melting of crystals at bonding temperature controls adhesion strength. Therefore, the co-crystallization of blend components in the fast cooling of film production provides the opportunity for re-melting of both

crystal components at the same temperature. The influence of melting behaviour will be discussed further in adhesion strength results.

The difference between the T_m of LLDPE and LCB-m was very small. (Figure 6-1b) By incorporating the LCB-m into LLDPE, the T_m corresponding LLDPE shifted to lower temperatures and the melting peak broadens. Blending with LCB-m broadened the small shoulder of LLDPE melting peak and shifted it to lower temperatures. For 60wt%LCB-m the melting peaks of components merged together and show one single peak.

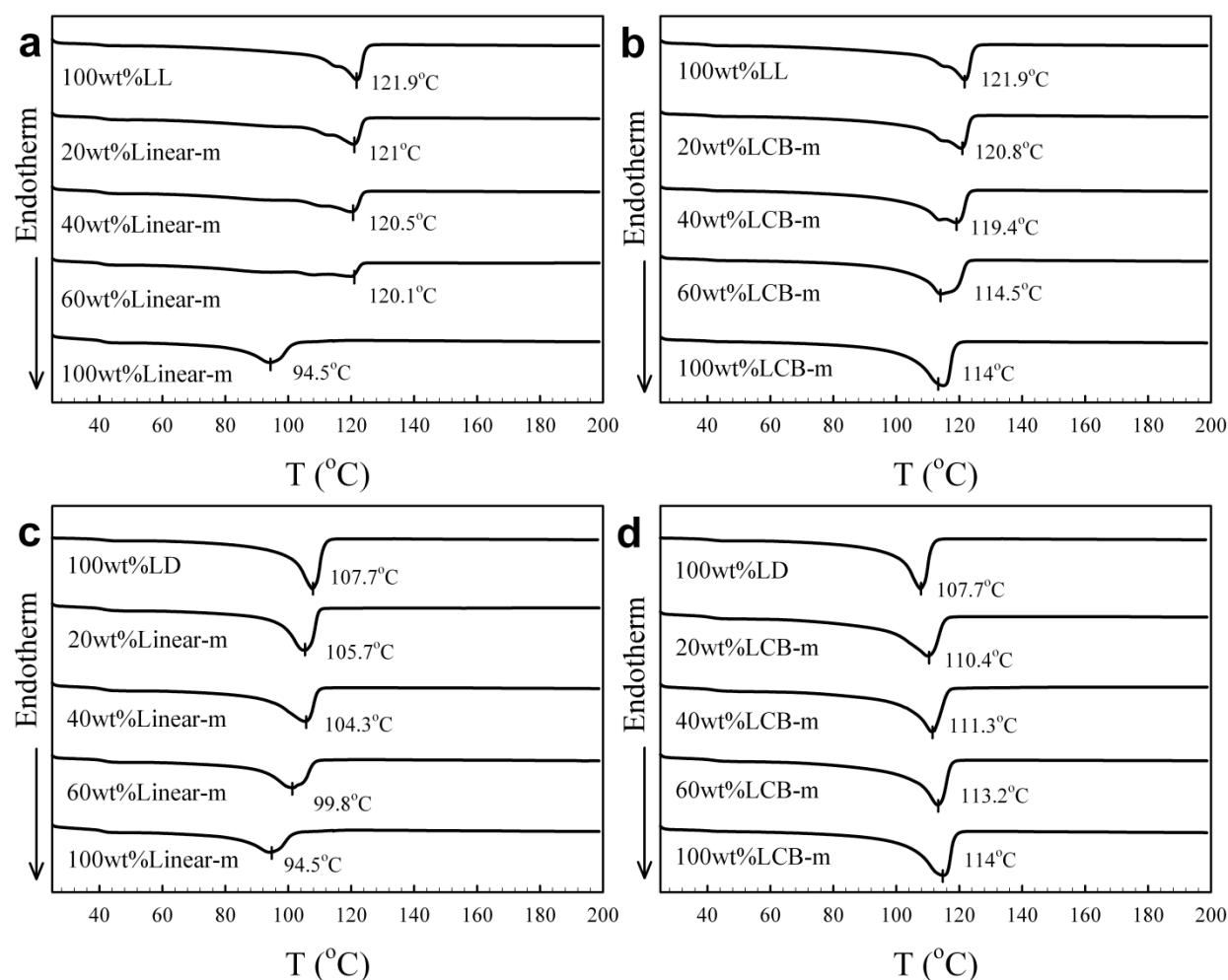


Figure 6-1 DSC melting curve of the first heating run for films of the neat polymers and their corresponding blends, (a) LL/Linear-m blends, (b) LL/LCB-m blends, (c) LD/Linear-m blends, (d) LD/LCB-m blends

The influence of the cooling rate which is very fast in film processing needs to be addressed. Fast cooling rates increase co-crystallization. Hill et al.²² studied the effect of cooling rate on co-crystallization and phase segregation in polyethylene blends. They reported that the phase separation increased with decreasing cooling rate. For high cooling rates, crystallization occurs at conditions far away from thermodynamic equilibrium. The morphology of blends after quenching is strongly affected by either phase separation or miscibility in the melt.

LDPE showed a higher T_m than Linear-m. (Figure 6-1c and d) As the amount of Linear-m increased in the blend, T_m shifted to lower values. Furthermore, all LD/Linear-m blends showed a single melting peak which was broadened by increasing Linear-m content. The single melting peak indicated that co-crystallization occurred between all LDPE and Linear-m chains. The T_m of LDPE was however lower than that of LCB-m and it increased gradually by incorporating the LCB-m. (Figure 6-1c)

The miscibility of LDPE blends with a variety of ethylene α -olefin copolymers in solid and melt states has been widely studied^{17,18,32-34}. Due to complexity of LDPE chain structure and branch distribution, there is high possibility of similar chain crystallize-ability with other types of polyethylene. The miscibility in the melt state and co-crystallization in solid state were reported in several studies^{32, 33}. However, for every binary blend the extent of co-crystallization and melt miscibility was reported to depend on density, SCB content, and composition.

6.4.2 Transient extensional rheology

The measurement of interfacial self-adhesion in this work was performed while either partially or completely molten. The melt strength and physical entanglement had a strong contribution to adhesive bond strength. In addition, the adhesive laminate provided a strong carrier web which transferred the stress completely to the adherents. Thus the transient extensional behaviour of the resins and blends could assist in understanding their self-adhesion performance.

The transient elongational viscosity of the resins and blends at different strain rates at 150°C is illustrated in Figure 6-2. The uniaxial extension flow is very sensitive to the molecular and microstructural characteristics of the resins and the resistance against extensional flow can be described by the extensional viscosity η_E^+ . The extensional viscosity of a polymer results from the resistance of chains and branches entanglements. The melt strength is controlled by the rate of

chain disentanglement in relation to the deformation rate and will be discussed further below. As expected, the linear-m and LLDPE extensional viscosity followed a linear viscoelastic behaviour over a large strain range and no strain hardening was seen in the rate range studied here. (See Figure 6-2a and b) Surprisingly, the LCB-m did not show strain hardening. The content of long chain branches in this LCB-m was determined as 0.3LCB/ 10^4 C atom with gel permeation chromatography. Thus, this low level of LCB did not develop the strain hardening behaviour.

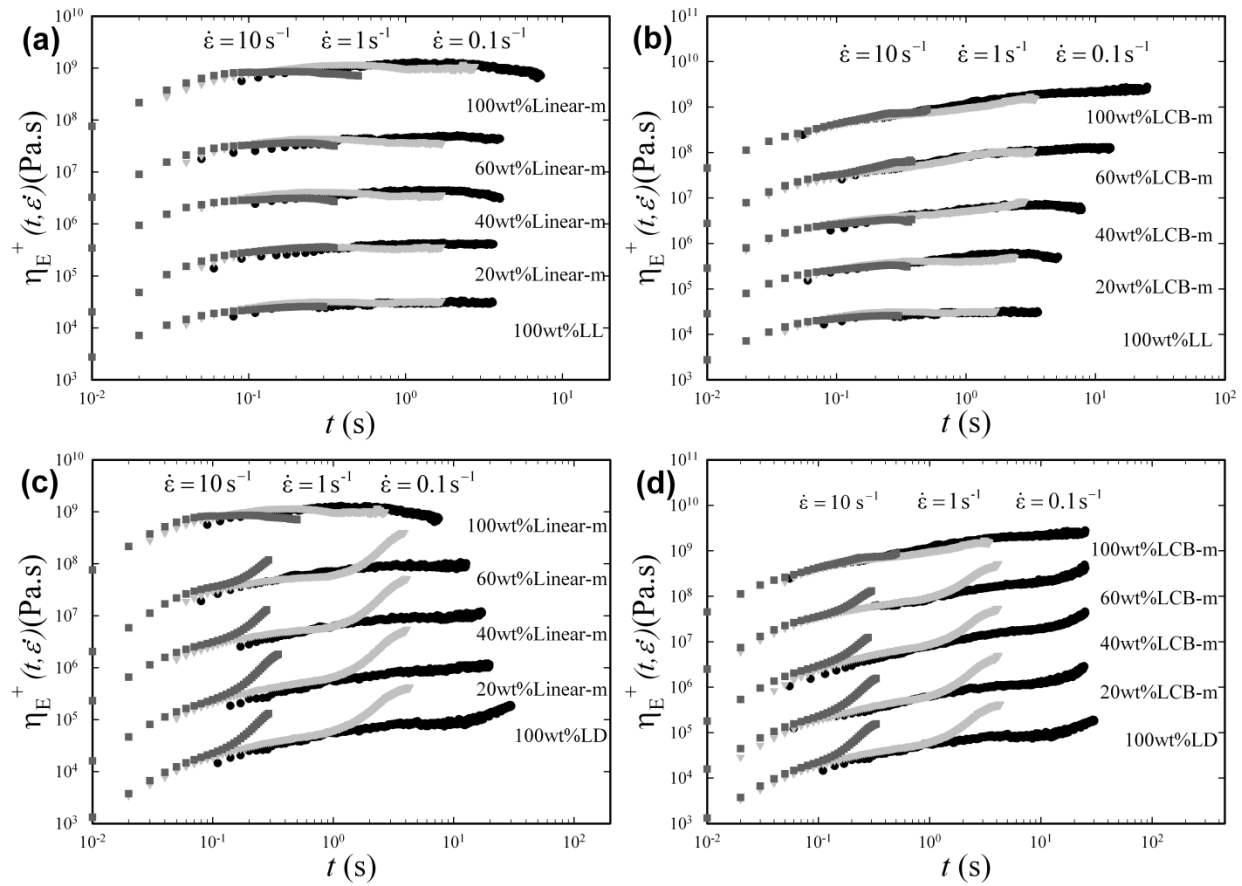


Figure 6-2 Transient elongational viscosities as a function of time at different strain rates, (a) LL/Linear-m blends, (b) LL/LCB-m blends, (c) LD/Linear-m blends, (d) LD/LCB-m blends, $T=150^{\circ}\text{C}$, the curves have been shifted by multiplication

For high extension rates, by increasing time and strain, η_E^+ rapidly increased above the linear viscoelastic limit. (Figure 6-2c and d) At low strain rates, η_E^+ of LDPE followed the linear viscoelastic behaviour. By blending, the decreasing branch content in the whole composition

caused a decrease in the nonlinear behaviour and shifted the maximum η_E^+ to lower values. These results are in agreement with previous reports in literature for linear and branched PEs^{35–37}.

6.4.3 Interfacial self-adhesion strength

The ultimate interfacial self-adhesion strength for all blend compositions is illustrated in Figure 6-3. As expected, the superior adhesion strength of metallocene resins observed. The mechanisms involved in the interfacial adhesion of conventional PEs compared to metallocene resins were discussed in a previous work¹⁶. Briefly, it is believed that the polymer chains at the surface of the film are important in strengthening the adhesion at interface^{6,38}. The thickness of the important surface layer for self adhesion is of the order of R_g of average molecular weight chains⁶. This layer is enriched by short chains in conventional PEs, which cause the formation of a weak adhesion^{39–41}. Because of the broad molecular weight and heterogeneous composition distribution in conventional PEs, a segregated layer of highly branched low molecular weight chains formed at the surface of conventional PE films. The thickness of this concentrated highly branched short chains layer was reported to be up to 100 nm in Zn-LLDPE^{42,43}. The segregated low molecular weight chains at the surface made weaker adhesion at the interface compared to high molecular weight chains. The presence of such layer was not observed in metallocene catalyzed resins because of their homogeneous composition distribution⁴².

Incorporation of metallocene resins to both LD and LL enhanced adhesion strength. By increasing metallocene content to 40wt% and 60wt%, the adhesion strength increased for all four types of blends. (See Figure 6-3) However, the amount of this enhancement varied among different types of blends. The % increase was more for LL/Linear-m blends, than for LL/LCB-m blends. Similarly, the % increase was more for the blends of LD/Linear-m than for the blends of LD/LCB-m. The increase in interfacial adhesion in both solid and melt state by blending metallocene LLDPE with conventional PEs has been reported previously^{42,44}. Referring to the segregated layer of short chains at the surface would explain the enhancement of adhesion strength by blending with metallocene resins. Blending metallocene with conventional PEs increased the number of long chains and reduced the ratio of short chains relative to the whole composition of the blends. Therefore, the thickness of the layer formed from the segregation of short chains decreased. However, even the adhesion strength of 60wt% metallocene in all four types of blends was quite far beneath its value for neat metallocene resins. This result indicated

that even low amount of heterogeneous component in the blend was capable of causing the formation of this layer ⁴². Consequently, the relative rate of diffusion of polymer chains across the interface would favor the more mobile low molecular weight chains and would not make strong adhesion.

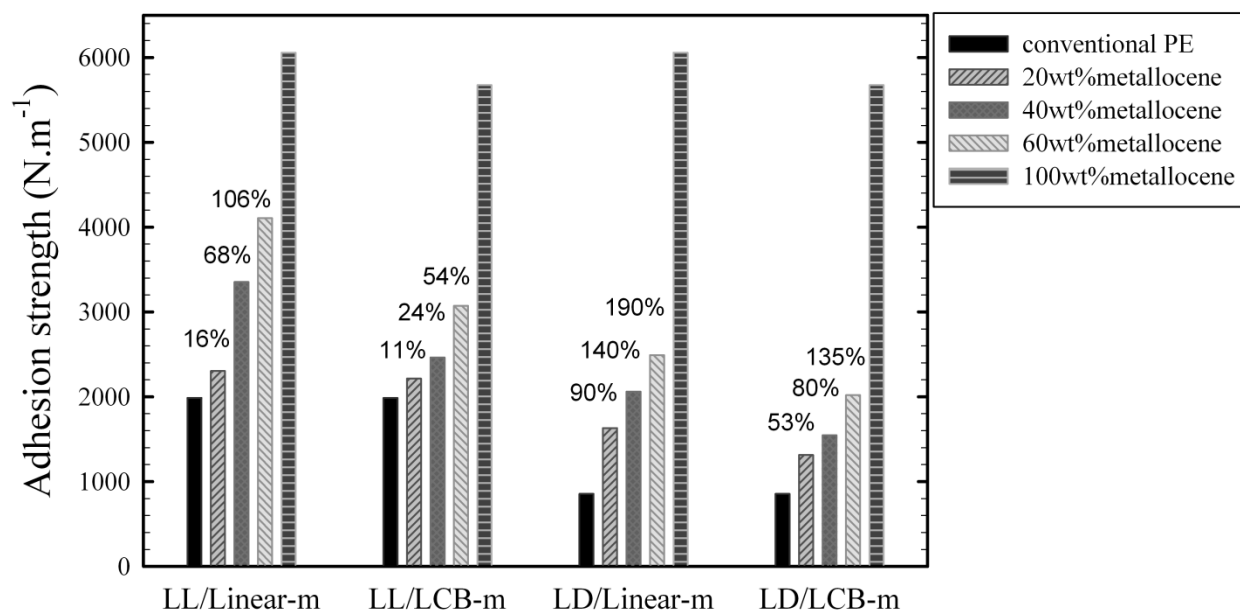


Figure 6-3 Maximum interfacial self-adhesion strength of LL/Linear-m, LL/LCB-m, LD/Linear-m, and LD/LCB-m; Labels on top of every column illustrates the %increase of maximum adhesion strength for every blend composition

In the case of LD blends, there was an additional source for adhesion strength enhancement which could explain the higher % increase of adhesion strength. Due to its highly branched chains, LDPE lack the fast interdiffusion. Linear chains of Linear-m resin provided faster interdiffusion and more entanglements on the other side of interface. The % increase in adhesion strength in both LD/Linear-m and LD/LCB-m were higher than their counterpart blends with LLDPE. (Figure 6-3) In addition, the blends of 20wt%, 40wt%, and 60wt% LD/Linear-m showed higher % increase than their counterparts of LD/LCB-m blend. It indicated that the faster diffusion of Linear-m compared to LCB-m caused larger improvement of interfacial adhesion.

The adhesion strength of all blend compositions in correlation with their extensional viscosity indicated that the melt strength and strength hardening were not the primary factors to determine

the adhesion strength. These results indicated that the interdiffusion is the main parameter controlling the maximum self-adhesion strength. However, the role of melt strength and extensional rheological behaviour of resins will be discussed further sections.

6.4.4 Temperature dependence of interfacial self-adhesion strength

As mentioned earlier, interfacial self-adhesion of semicrystalline polymers depends strongly on the temperature. Presence of the crystals at the interface vicinity hinders the chain motion and diffusion across the interface. Therefore, the adhesion experiments should be performed in a range of temperature for any semicrystalline polymer. Figure 6-4 illustrates the adhesion strength in a range temperature for every blend composition. At low temperatures, a weak adhesion was created. Few degrees increase in temperature improved the adhesion strength sharply. Afterward, adhesion strength remained almost constant for a temperature interval, called plateau adhesion strength. This plateau covered a temperature range which had a lower and an upper boundary, called plateau initiation temperature (T_{pi}) and final plateau temperature (T_{pf}), respectively. For a few degrees higher than T_{pf} , a sharp decrease in adhesion strength occurred. The upper boundary of the temperature plateau was bounded by the damage to the joint caused by heat and pressure; the lower temperature boundary was bounded by the amount of heat available at the interface to melt the two adherents and was affected by crystallinity and T_m . The lower limit was controlled by crystallinity and the upper limit was controlled by rheological properties, specifically melt strength of the polymer. In order to achieve the highest adhesion strength, the temperature must be high enough to melt the interface between the two polymer layers, yet low enough to prevent the distortion at the joint area. The temperature windows of plateau for all blend compositions are illustrated in Figure 6-5.

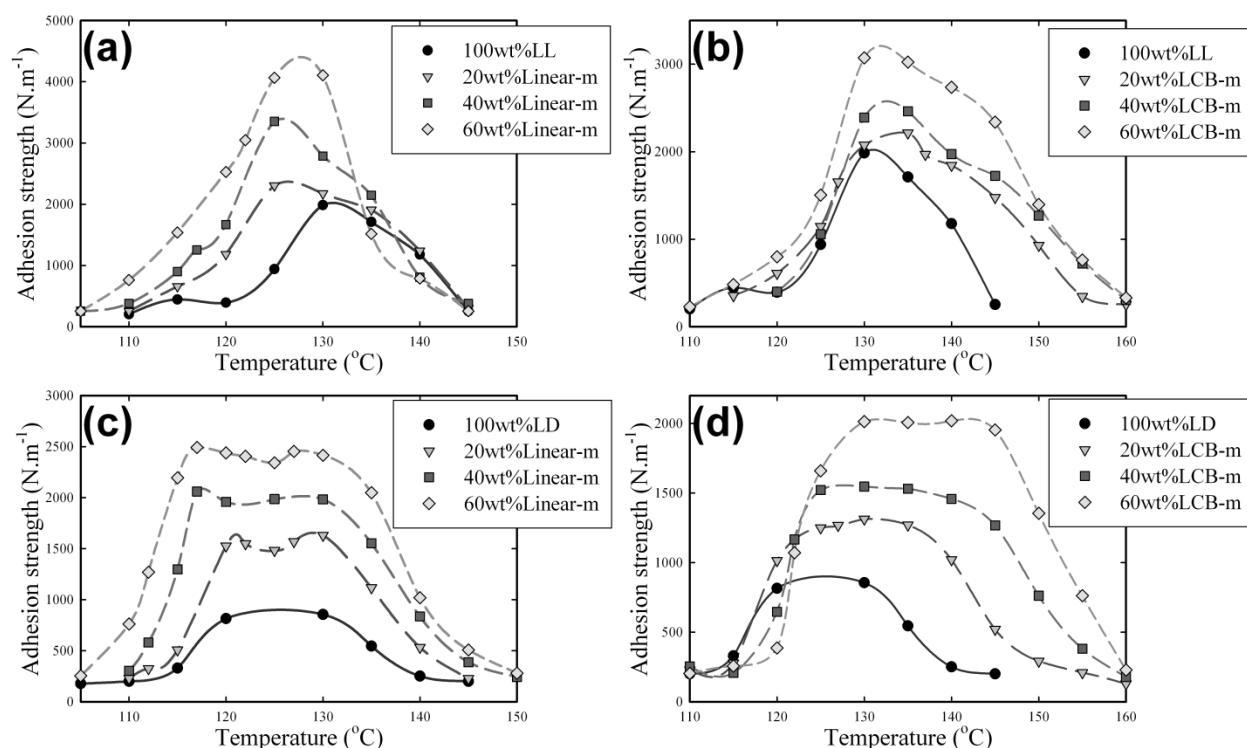


Figure 6-4 Interfacial self-adhesion strength of (a) LL/Linear-m, (b) LL/LCB-m, (c) LD/Linear-m, and (d) LD/LCB-m, as a function of temperature

It can be seen in Figure 6-4, the incorporation of metallocene caused either an increase or a decrease in T_{pi} . Regarding the thermal properties, the melting point and crystal content of LD/Linear-m and LL/Linear-m blends decreased when the metallocene content increased. (See Figure 6-1) Consequently, a decrease in T_{pi} was observed in Figure 6-4a, b, and c. In LD/LCB-m blends, increasing metallocene content increased T_{pi} because of higher T_m of LCB-m than LD. Therefore, the presence of LCB had no significant effect on T_{pi} and the amount and distribution of SCB were rather the controlling parameters. Smaller crystal sizes induced bonding at lower temperatures. The temperature dependence of interfacial adhesion and the role of both T_g in glassy and T_m in thermoplastic polymers has been reported^{44,45}. To our knowledge, there is no study discussing the temperature boundaries for a polymer's interfacial adhesion.

The sharp drop down of adhesion strength occurred in high temperatures a few degrees higher than the upper limit. This was the result of complete merging of the two film layers together. Consequently, the crack propagation in peeling did not follow the interface. Then a breakage at the edge of joint area occurred, followed by a cohesive failure in the vicinity of the support layer

^{8,14,45}. This means that increasing the temperature caused lower melt strength with the possibility of defects, which caused breakage at the edge. The low interface strength at the temperatures above a certain limit has been also reported before ⁴⁴.

Figure 6-5 illustrates the temperature window of plateau broadness for all blend compositions. For LL/Linear-m and LL/LCB-m blends as well as neat LLDPE, the temperature window was narrow. For all compositions of LD/Linear-m and LD/LCB-m blends as well as neat LDPE, the temperature window was very broad. Moreover, both metallocene resins showed a broad temperature window. As mentioned earlier, the lower limit of plateau was controlled by crystallinity. Accordingly, the T_{pf} was the origin of the difference between these two groups of blends. It is thus important to explore the influence of molecular architecture in ethylene olefin blends on the position of T_{pf} .

Peeling a joint at temperatures higher than melting point involve unsteady-state and time dependent deformations. The deformation of entangled network of chains is elastic while the pull-out of chains at the interface is a viscous process. If v designate the peel rate and is 33mm.s^{-1} and h is the film thickness= $50\text{ }\mu$, the strain rate could be estimated roughly from $\dot{\gamma} = \frac{v}{h}$. A rough estimation for strain rate would thus be 660 s^{-1} . So, peeling is equivalent to applying an extensional force on entangled chains at very high strain rates. It is therefore necessary to examine the dynamic characteristics of these different macromolecular structures of polyethylene.

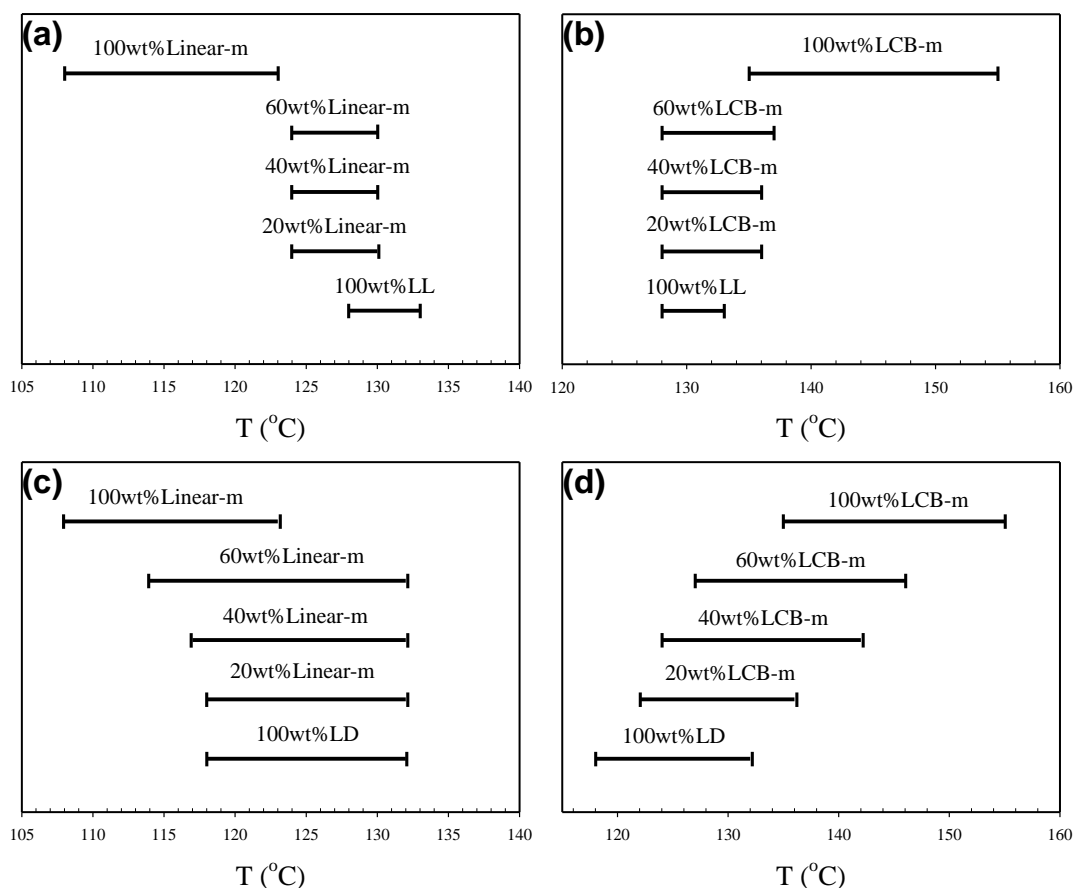


Figure 6-5 Adhesion strength plateau broadness for (a) LL/Linear-m blends, (b) LL/LCB-m blends, (c) LD/Linear-m blends, (d) LD/LCB-m blends

Polymer entanglements act as crosslink points at short processing time scales. On the other hand, the polymer chains can reptate along each other at longer timescales and open the entanglement ties. Then, the behaviour is more viscous. It is useful to have a rough estimation of time scales of chain motions versus time scale of peeling. The relaxation time τ , obtained from Maxwell relaxation time was used to obtain Deborah number (De) as a reduced parameter. The Maxwell relaxation time (τ) was obtained from the frequency of $G'(\omega)$ and $G''(\omega)$ cross-over point in frequency sweep shear measurements. This relaxation time has been used in several studies on adhesives behaviours and rheological characterization of polymers^{8,46-49}. De was defined as $De = \frac{\tau}{t}$ in which τ is the time scale of the material's response and t is the timescale of the measurement process. The time scale of peeling was 600 ms. Table 6-2 shows the relaxation time

and the De calculated for four neat resins in this study. The relaxation time τ is one or two order of magnitude smaller than timescale of peeling.

Creton et al.⁸ also calculated De in debonding of SBR elastomers as pressure sensitive adhesives. Analyzing the corresponding failure mechanism, they defined a critical value for Deborah number (De_{crit}) which was estimated ≈ 0.8 for the elastomers studied and the probe geometry in their work. They showed that for $De < De_{crit}$, SBR behaves as a viscoelastic fluid. The exact value of $De_{crit}=0.8$ might not be accurate to apply for the experiment and materials in this study. However, the very small values represented in Table 6-2 for De for all the resins indicate a viscous dominant regime, indicating an irreversibly consumed energy when fracture develops and propagate.

Among all resins, LD has the highest τ and consequently highest De . As long branches are incorporated to the chains and get involved in entanglements, the relaxation mechanism of chains changes from simple reptation to arm retraction^{50,51}. The relaxation time of the material increases strongly and it becomes elastic at larger timescales. Due to the presence of long chain branches, τ for LDPE is one order of magnitude higher than for linear resins.

Table 6-2 Relaxation time and De number of neat resins

Resin	τ (ms)	De
LDPE	199	0.332
Zn-LLDPE	10	0.016
Linear-m	15	0.025
LCB-m	25	0.042

It is necessary to compare the viscous dissipation of the resins, which provides a better understanding of the broadness of plateau adhesion strength. As mentioned earlier, the type of deformation in melt adhesion measurement resembles the uniaxial extensional flow. Therefore, the toughness calculated from the area under the stress-strain curve in extensional flow can be used to estimate the energy dissipated by melt before rupture. Although the high strain rate involved in peeling is practically unattainable, the data from lower strain rates provides a comparative estimation. The toughness of neat resins obtained from area under the extensional flow curve at $\dot{\epsilon}=10 \text{ s}^{-1}$ at three different temperature of 125°C, 135°C and 150°C are listed in

Table 6-3. These temperatures were selected with regard to the plateau temperature window of all four resins.

The highest toughness belongs to LDPE compared to the others. The toughness of LLDPE is significantly lower than all other resins. The amount of energy per volume that LLDPE can absorb before rupture is much lower than LDPE or metallocene resins. This could explain the very narrow plateau temperature window for LLDPE and its blends.

The presence of LCB in LDPE deteriorates the adhesion strength. As discussed in a previous work ¹⁶, this is due to the hindered diffusion of its chains. However, the melt toughness in blends containing LDPE remains higher for the range of adhesion temperatures. (Table 6-3) This high toughness in elongational flow causes LDPE to maintain its strength in a wider range of temperature compared to LLDPE.

The high toughness of metallocene resins was because of high strain at break. The adhesion strength of some other grades of metallocene catalyzed ethylene α -olefin was examined ¹⁶. Our findings showed that among linear metallocene resins, narrower plateau was observed for low molecular weight resins but broader plateau occurred for high molecular weight resins. Moreover, a wide plateau was observed for LCB containing metallocene resins with different LCB contents.

Table 6-3 Melt toughness obtained from area under σ - ϵ curve in extensional flow at $\dot{\epsilon}=10 \text{ s}^{-1}$

Resin	Toughness (kJ.m^{-3})		
	125°C	135°C	150°C
LDPE	55.1	46.0	19.1
Zn-LLDPE	13.2	12.4	7.6
Linear-m	45.5	36.0	16.6
LCB-m	40.0	34.0	15.4

Based on our observations of the failure mechanism, the T_{pf} is the boundary in which a transition occurs from interfacial failure and bulk deformation to edge breakage. This transition can occur depending on the characteristics of the material such as the length of the polymer chains, the number and the nature of entanglements, and the process conditions such as the applied time, pressure, and temperature. At temperatures higher than T_{pf} , the provided energy by the heated

bars overcomes the required energy for polymer chains to disentangle. Latter would allow the polymer chains to slip out of the bulk by applying lower peeling force.

High melt toughness in elongational flow indicates higher required energy to overcome the chain entanglements at a certain temperatures. Hence for a polymer with higher melt toughness to fail in a joint a higher temperature would be required which in result would broaden the adhesion plateau. The broad temperature plateau window in LD and its blends could be explained by higher melt toughness in extensional melt flow. Creton et al.¹⁴ also examined the adhesion energy at the soft interface of poly dimethylsiloxane elastomers in terms of energy dissipation mechanisms at molecular and macroscopic scales. They concluded that both large extensional bulk deformations and interfacial failure determine the adhesion strength.

6.5 Conclusions

Investigation of interfacial self adhesion for polyethylene blend films in melt state has provided novel information on the role of molecular architecture of components on adhesion performance. Four binary blends were prepared from the combination of a conventional polyethylene; LDPE or Ziegler-Natta LLDPE, with a metallocene catalyzed ethylene α -olefin copolymer; linear (Linear-m) or long chain branched (LCB-m). These films were examined in terms of interfacial self-adhesion at temperatures higher than T_m .

Melting behaviour and partial co-crystallization, was observed for all blend compositions. All compositions of LL/Linear-m and LD/Linear-m blends showed higher increase percentage in adhesion strength compared to their counterparts in LL/LCB-m and LD/LCB-m blends. Moreover, the increase percentage of adhesion strength for LDPE blends was higher than LLDPE blends. The chain interdiffusion has the primary role to build up the strength at the interface. Moreover, melt toughness has a crucial role on maintaining its highest strength for a wide range of temperatures. The presence of LCB in either low content in LCB-m or high content in LDPE hinder the chain motion compared to linear chains with approximately the same length. The broad adhesion strength plateau of LDPE as highly LCB containing resin illustrated the important role of high relaxation time of chains and high level of entanglements in the melt behaviour. Based on the correlations, melt toughness in extensional flow provides a basis to predict the

relative plateau broadness of a polymer. The broad adhesion strength plateau for metallocene resins was attributed to their high melt toughness.

6.6 Acknowledgement

Financial support from 3S Pack NSERC/Saputo/Prolamina industrial research chair is gratefully acknowledged.

6.7 References

1. Wool, R. P., Yuan, B.-L. & McGarel, O. J. Welding of polymer interfaces. *Polymer Engineering & Science* **29**, 1340–1367 (1989).
2. Jud, K., Kausch, H. H. & Williams, J. G. Fracture mechanics studies of crack healing and welding of polymers. *J Mater Sci* **16**, 204–210 (1981).
3. Prager, S. & Tirrell, M. The healing process at polymer–polymer interfaces. *The Journal of Chemical Physics* **75**, 5194–5198 (1981).
4. Kausch, H. H. & Tirrell, M. Polymer Interdiffusion. *Annual Review of Materials Science* **19**, 341–377 (1989).
5. Kim, K. D., Sperling, L. H., Klein, A. & Hammouda, B. Reptation Time, Temperature, and Cosurfactant Effects on the Molecular Interdiffusion Rate during Polystyrene Latex Film Formation. *Macromolecules* **27**, 6841–6850 (1994).
6. Wool, R. P. *Polymer Interfaces: Structure and Strength*. (Hanser-Gardner Publications, 1995).
7. Brown. The Adhesion Between Polymers. *Annual Review of Materials Science* **21**, 463–489 (1991).
8. Schach, R. & Creton, C. Adhesion at interfaces between highly entangled polymer melts. *Journal of Rheology* **52**, 749–767 (2008).
9. Boiko, Y. M., Guérin, G., Marikhin, V. A. & Prud'homme, R. E. Healing of interfaces of amorphous and semi-crystalline poly(ethylene terephthalate) in the vicinity of the glass transition temperature. *Polymer* **42**, 8695–8702 (2001).
10. Gent, A. N., Kim, E.-G. & Ye, P. Autohesion of crosslinked polyethylene. *Journal of Polymer Science Part B: Polymer Physics* **35**, 615–622 (1997).
11. Smith, G. D., Plummer, C. J. G., Bourban, P.-E. & Manson, J.-A. E. Non-isothermal fusion bonding of polypropylene. *Polymer* **42**, 6247–6257 (2001).
12. Xue, Y.-Q., Tervoort, T. A. & Lemstra, P. J. Welding behaviour of semicrystalline polymers. 1. The effect of nonequilibrium chain conformations on autoadhesion of UHMWPE. *Macromolecules* **31**, 3075–3080 (1998).

13. Xue, Y.-Q., Tervoort, T. A., Rastogi, S. & Lemstra, J. Welding Behavior of Semicrystalline Polymers. 2. Effect of Cocrystallization on Autoadhesion. *Macromolecules* **33**, 7084–7087 (2000).
14. Léger, L. & Creton, C. Adhesion mechanisms at soft polymer interfaces. *Phil. Trans. R. Soc. A* **366**, 1425–1442 (2008).
15. Brydson, J. A. *Plastics Materials*. (Butterworth-Heinemann, 1999).
16. Najarzadeh, Z. & Ajji, A. Role of molecular architecture in interfacial self-adhesion of polyethylene films.
17. Delgadillo-Velázquez, O., Hatzikiriakos, S. G. & Sentmanat, M. Thermorheological properties of LLDPE/LDPE blends: Effects of production technology of LLDPE. *Journal of Polymer Science Part B: Polymer Physics* **46**, 1669–1683 (2008).
18. Hussein, I. A., Hameed, T., Abu Sharkh, B. F. & Mezghani, K. Miscibility of hexene-LLDPE and LDPE blends: Influence of branch content and composition distribution. *Polymer* **44**, 4665–4672 (2003).
19. Fang, Y., Carreau, P. J., Lafleur, P. G. & Ymmel, S. Properties of mLLDPE/LDPE blends in film blowing. *Polymer Engineering and Science* **45**, 343–353 (2005).
20. Fang, Y., Carreau, P. J. & Lafleur, P. G. Thermal and rheological properties of mLLDPE/LDPE blends. *Polymer Engineering & Science* **45**, 1254–1264 (2005).
21. Chen, F., Shanks, R. A. & Amarasinghe, G. Rheological and thermal properties of single-site polyethylene blends. *Journal of Applied Polymer Science* **95**, 1549–1557 (2005).
22. Hill, M. J. & Barham, P. J. Morphology maps of binary blends of copolymers produced using the metallocene catalyst process. *Polymer* **41**, 1621–1625 (2000).
23. Rana, D., Kim, H. L., Kwag, H. & Choe, S. Hybrid blends of similar ethylene 1-octene copolymers. *Polymer* **41**, 7067–7082 (2000).
24. Najarzadeh, Z. & Ajji, A. A novel approach toward the effect of seal process parameters on final seal strength and microstructure of LLDPE. *Journal of Adhesion Science and Technology* (2014).
25. Bensason, S. *et al.* Classification of homogeneous ethylene-octene copolymers based on comonomer content. *Journal of Polymer Science Part B: Polymer Physics* **34**, 1301–1315 (1996).
26. Bubeck, R. A. Structure–property relationships in metallocene polyethylenes. *Materials Science and Engineering: R: Reports* **39**, 1–28 (2002).
27. Hosoda, S. *et al.* Effect of the Sequence Length Distribution on the Lamellar Crystal Thickness and Thickness Distribution of Polyethylene: Perfectly Equisquential ADMET Polyethylene vs Ethylene/ α -Olefin Copolymer. *Macromolecules* **44**, 313–319 (2011).
28. Rana, D., Cho, K., Woo, T., Lee, B. H. & Choe, S. Blends of ethylene 1-octene copolymer synthesized by Ziegler–Natta and metallocene catalysts. I. Thermal and mechanical properties. *Journal of Applied Polymer Science* **74**, 1169–1177 (1999).

29. Rana, D. *et al.* Blends of ethylene 1-octene copolymer synthesized by Ziegler–Natta and metallocene catalysts. II. Rheology and morphological behaviors. *Journal of Applied Polymer Science* **76**, 1950–1964 (2000).
30. Kuwabara, K., Kaji, H., Horii, F., Bassett, D. C. & Olley, R. H. Solid-State ¹³C NMR Analyses of the Crystalline–Noncrystalline Structure for Metallocene-Catalyzed Linear Low-Density Polyethylene. *Macromolecules* **30**, 7516–7521 (1997).
31. Janicek, M., Cermak, R., Obadal, M., Piel, C. & Ponizil, P. Ethylene Copolymers with Crystallizable Side Chains. *Macromolecules* **44**, 6759–6766 (2011).
32. Hameed, T. & Hussein, I. A. Rheological study of the influence of MW and comonomer type on the miscibility of m-LLDPE and LDPE blends. *Polymer* **43**, 6911–6929 (2002).
33. Hussein, I. A. & Hameed, T. Influence of branching characteristics on thermal and mechanical properties of Ziegler-Natta and metallocene hexene linear low-density polyethylene blends with low-density polyethylene. *Journal of Applied Polymer Science* **97**, 2488–2498 (2005).
34. Robledo, N., Vega, J. F., Nieto, J. & Martínez-Salazar, J. The role of the interface in melt linear viscoelastic properties of LLDPE/LDPE blends: Effect of the molecular architecture of the matrix. *J. Appl. Polym. Sci.* **114**, 420–429 (2009).
35. Wagner, M. H., Kheirandish, S. & Yamaguchi, M. Quantitative analysis of melt elongational behavior of LLDPE/LDPE blends. *Rheol Acta* **44**, 198–218 (2004).
36. Kasehagen, L. J. & Macosko, C. W. Nonlinear shear and extensional rheology of long-chain randomly branched polybutadiene. *Journal of Rheology (1978-present)* **42**, 1303–1327 (1998).
37. Tabatabaei, S. H., Carreau, P. J. & Ajji, A. Rheological and thermal properties of blends of a long-chain branched polypropylene and different linear polypropylenes. *Chemical Engineering Science* **64**, 4719–4731 (2009).
38. Basin, V. E. Role of diffusion in the formation of adhesion bonds between polymers. *Polymer Mechanics* **3**, 561–563 (1967).
39. Kausch, H. H., Nguyen, T. Q. & Petrovska-Delacrétaz, D. Chain interdiffusion: macromolecules between Rouse and de Gennes. *Phys. Scr.* **1991**, 57 (1991).
40. Roland, C. M. & Boehm, G. G. A. Macromolecular diffusion and the autoadhesion of polybutadiene. *Macromolecules* **18**, 1310–1314 (1985).
41. Su, G. M., Best, K., Ranganathan, T., Emrick, T. & Crosby, A. J. Tailored Nanoparticles for Enhancing Polymer Adhesion. *Macromolecules* **44**, 5256–5261 (2011).
42. Qureshi, N. Z. *et al.* Self-adhesion of polyethylene in the melt. 2. Comparison of heterogeneous and homogeneous copolymers. *Macromolecules* **34**, 3007–3017 (2001).
43. Magonov, S. & Godovsky, Y. Atomic force microscopy, Part 8: Visualization of granular nanostructure in crystalline polymers. *American Laboratory* **31**, 52–58 (1999).
44. Shih, H. H., Wong, C. M., Wang, Y. C., Huang, C. J. & Wu, C. C. Hot tack of metallocene catalyzed polyethylene and low-density polyethylene blend. *Journal of Applied Polymer Science* **73**, 1769–1773 (1999).

45. Shih, H.-H. & Hamed, G. R. Peel adhesion and viscoelasticity of poly(ethylene-co-vinyl acetate)-based hot melt adhesives. I. The effect of tackifier compatibility. *Journal of Applied Polymer Science* **63**, 323–331 (1997).
46. O'Connor, A. E. & Macosko, C. W. Melt versus solvent coating: Structure and properties of block-copolymer-based pressure-sensitive adhesives. *J. Appl. Polym. Sci.* **86**, 3355–3367 (2002).
47. Ganapathy, R. & Sood, A. K. Tuning Rheochaos by Temperature in Wormlike Micelles. *Langmuir* **22**, 11016–11021 (2006).
48. Lin, G.-G., Shih, H.-H., Chai, P.-C. & Hsu, S.-J. Influence of side-chain structures on the viscoelasticity and elongation viscosity of polyethylene melts. *Polym Eng Sci* **42**, 2213–2221 (2002).
49. Gahleitner, M. Melt rheology of polyolefins. *Progress in Polymer Science* **26**, 895–944 (2001).
50. McLeish, T. C. B. & Larson, R. G. Molecular constitutive equations for a class of branched polymers: The pom-pom polymer. *J. Rheol.* **42**, 81 (1998).
51. Chen, X., Costeux, C. & Larson, R. G. Characterization and prediction of long-chain branching in commercial polyethylenes by a combination of rheology and modeling methods. *Journal of Rheology (1978-present)* **54**, 1185–1205 (2010).

CHAPTER 7

GENERAL DISCUSSION

Heat sealing is the process of joining thermoplastic materials and is the most popular method for bonding two polymer films in package closure. Heat sealing is an immensely complicated type of symmetric interfacial adhesion of polymer surfaces. The process is non-isothermal and generally occurs in a very short time. The strength of heat seal joint is critical in two conditions: before cooling while the joint is still molten and after cooling and re-crystallization of the joint area. The final package integrity depends on the heat seal performance. The quality of a heat seal is assessed based on the integrity of the joint, the behaviour of seal strength at various times and temperatures, broadness of the temperature range, seal initiation temperature, seal failure mode, and seal uniformity. The factors, which determine the quality of a heat seal, can be classified as heat seal process parameters and film parameters. Process parameters are applied temperature, dwell time, and pressure. These parameters should be optimized to maximize seal performance while minimizing costs. Film parameters are generally the material characteristics, film thickness, crystallinity, orientation, etc. Polyethylene is the most used polymer in the seal layer. Thus, in this work different types of polyethylene resin were used to study the role of molecular architecture on seal performance.

The concurrent role of temperature, time and pressure should be considered when the effect of process parameters on final seal strength is analyzed. Furthermore, optimization of each of these parameters should be done with respect to the other two, which has not been done in previous studies. Therefore, in the first part of this study, the seal experiments were conducted for LLDPE in order to optimize the sealing parameters and to perceive the effect of concurrent change of “temperature - dwell time” and “temperature - pressure” on the seal strength. Mapping of process safety zone was introduced for seal strength in a range of heat seal process variables.

It is practically important to predict seal initiation temperature (T_{si}) and plateau initiation temperature (T_{pi}) of sealant by knowing its melting point. The T_{si} and T_{pi} are correlated with the initial amorphous fraction of the film. In the melting distribution curve of LLDPE, the interval corresponding $T_{si} - T_{pi}$ temperature window is shown for every dwell time. We argue here that

the $T_{pi} - T_{mf}$ relationship is strongly influenced by the dwell time of the heat seal process which, to our knowledge, was not considered by the previous studies.

The time dependence of seal strength before complete sealing of the interface is studied in regard to adhesion models based on reptation theory. We found that the seal strength has a linear correlation with the square root of dwell time. This is supported by the prediction of reptation motion theory and adhesion models. Moreover, seal strength increases faster over time at higher temperature.

Analysis of the AFM and SEM images reveal topography and morphology of surfaces after peeling of seal from microscopic point of view. The AFM 3D height micrographs at different sealing temperatures along with the heat sealing curve showed the morphology of fractured surface after debonding. Our findings indicated that the crack propagation through the sealed interface leaved numerous fractured craze fibrils on the peeled surfaces. The origin of these fibrils was attributed to the bridges formed during sealing by the interdiffusion of chains across the interface and the entanglements made by them. These entanglements act as stitches attaching two film layers. The density and length of these fibrils depends on the depth of diffusion and the number of chains available for motion at every defined sealing temperature. Our findings revealed that, by increasing the temperature the fibrils became thicker and larger.

The roughness analysis on film surfaces after peeling could quantitatively characterize the topography of peeled surfaces. The RMS and R_a roughness parameters illustrate the much rougher surfaces after breakage of the strong bonds. Then, increasing the temperature causes rougher surfaces. Moreover, the roughness parameters are in the same range in the case of the peeled surfaces after sealing at T_{pi} for every dwell time. Although T_{pi} is different at each dwell time, the roughness parameters are similar for various sealing temperatures and dwell times. This indicates that the diffusion had reached to a certain point at T_{pi} , regardless of the process conditions.

The fibrillar surface morphologies observed by AFM are also confirmed by the SEM micrographs. The fibrillar morphology is originated from interlamellar tie molecules which have formed tight stretched links between crystals across the interface. As it was also reported in literature, the local yielding and microvoiding pull the material into highly oriented craze fibrils.

In the second phase of this work we focused on the role of molecular architecture of sealant material in heat sealing. A series of polyethylene polymers with variety of chain structures were

chosen. In order to avoid complications due to crystallization in seal area after cooling, the hot tack strength has to be examined. Furthermore, the importance of hot tack in flexible packaging modern machinery as well as the lack of knowledge in open literature about controlling mechanisms of hot tack emphasises the need for a fundamental investigation. In current work, the adhesion strength terminology was used as the substitute for industrial term of hot tack, because the adhesion strength terminology has been used in fundamental studies on interfacial adhesion in literature. In this part, the level of SCB and LCB in resins was quantitatively determined using NMR and GPC techniques.

For the three linear metallocene catalyzed LLDPE containing equal amount of SCB, increasing M_w augments the adhesion strength. Furthermore, a linear dependence of adhesion strength to M_w and a similar linear relationship for elongational viscosity was observed. These results revealed that the adhesion strength at temperatures higher than the melting point is closely related to the melt strength of the resins. However, the fast diffusion of linear chains across the interface is a prerequisite to create entanglements across the interface.

The influence of LCB should be examined in two categories of conventional and single site catalyzed polyethylene resins. In conventional polyethylene resins, highly long branched chains cause LDPE to have very low self-adhesion strength. In metallocene catalyzed resins, the presence of a very low amount of long chain branches leads to lower adhesion strength compared to their linear counterparts. We argued that even low level of LCB hinder the reptation motion and diffusion. As the main conclusion of this part, the interfacial self-adhesion strength is primarily the outcome of chain interdiffusion across the interface and then the melt strength of polymer.

The ultimate adhesion strength of linear Ziegler-Natta and metallocene catalyzed LLDPE are evaluated. Our findings indicate that the heterogeneity in composition distribution of conventional polyethylene result in much lower self-adhesion strength compared to their homogeneous counterparts. This was explained to occur because of high concentration of highly branched low molecular weight chains in the surface of heterogeneous films. The segregated layer of short chains at the surface, which is doubled upon sealing, is less effective in strengthening the interface than long chains. Therefore, the composition distribution of polymer chains including SCB distribution is the influential parameter.

Due to cost and processability concerns, metallocene catalyzed ethylene –olefins are being generally used in the form of blends with conventional polyethylene resins. Therefore, the performance of variety of chain structures in the blends should be explored. Long chain branching is a key factor in both categories of metallocene catalyzed and conventional polyethylene resins. In the third part of the our work, four binary blends were prepared from one component of conventional polyethylene either LDPE or Ziegler-Natta LLDPE, and another component of metallocene catalyzed ethylene α -olefin copolymer either linear (Linear-m) or long chain branch containing (LCB-m). In order to determine the adhesion temperature range, the melting behaviour of films should primarily be examined. The melting thermographs of the first DSC cycle of the blend films indicate cocrystallization and hence partial miscibility for all blend compositions. It also showed the gradual shift of T_m toward the melting point of second component by blending.

Our findings showed that the incorporation of metallocene of both kinds into conventional PEs enhance the self adhesion. However, the adhesion strength of the blends containing 60wt% metallocene resins is far weaker than the adhesion strength of neat metallocene resins. This is due to the formation of a segregated layer of highly branched short chains of LDPE or LLDPE at the surface of the films (as discussed in the second paper). In addition, blending linear-m with both LDPE and LLDPE leads to higher improvement in adhesion strength than blending with LCB-m. This is attributed to the superior diffusion of Linear-m across the interface, because of its linear chains. All compositions of LDPE blends showed higher %increase for adhesion strength compared to their LLDPE counterparts. This finding suggests that for LDPE blends, a fast diffusion was provided by both Linear-m and LCB-m. The temperature ranges in which the films show a plateau of their highest adhesion strength were determined. For all LDPE/metallocene blends composition, the broadness of temperature window was broader than for the blends of LLDPE/metallocene. This indicates an inherent difference in melt behaviour of these two resins in extensional deformation. In order to understand this behaviour, the melt toughness obtained from extensional rheology was examined. Our findings indicated that the higher relaxation time of LDPE chains and higher level of entanglements in the melt results in higher melt toughness compared to LLDPE.

CHAPTER 8

CONCLUSIONS AND RECOMMENDATIONS

8.1 Conclusions

In this dissertation, the controlling parameters of heat sealing were explored. In the first phase of this work we studied the behaviour and effects of process parameters of heat seal quality for monolayer films of LLDPE as the most widely used polymer in seal layer applications. We investigated the seal strength of 50 μ monolayer LLDPE films supported by BOPP tape, in a range of 100-135°C for temperature, 0.1-3 s for dwell time and 0.1-3 for N.mm⁻² pressure. Increasing dwell time enhanced seal strength however, the rate of this enhancement varied for every interval of dwell time. We suggested that dwell time and temperature affect seal strength through the same mechanism of crystal melting and interdiffusion. However, pressure affected seal strength through wetting as a very different microscopic scale mechanism.

We also introduced the 3D mapping of process safety zone for the effect of concurrent change of “temperature - dwell time” and “temperature - pressure” on the seal strength. The temperature plateau broadness was studied in the range of dwell time and pressure. We showed that the time and temperature dependence of seal strength is a consequence of the amount of amorphous phase available at the interface of films. We investigated the time dependence of seal strengthening and confirmed our observations with the welding models. We showed that the seal strength has a linear correlation with the square root of sealing time. The slope of this linear correlation increased with temperature.

The topography and morphology of peeled surfaces which were sealed at different heat seal conditions were investigated. In the debonding failure mode, the crack propagates through the sealed interface and it leaves numerous highly oriented fractured craze fibrils on the peeled surfaces. The fractured fibrils were visible in AFM and SEM peeled surfaces of samples sealed at low temperatures. In specimens sealed at higher temperature (120°C) the fibrillar morphology developed into thicker and larger fibrils. They thus appear more like a network. This was explained by the better wetting and interdiffusion as a result of increasing temperature.

We also utilized the comparative RMS and R_a roughness parameters as representatives of the level of film surfaces' contact area and molecular interdiffusion. Increasing the seal temperature resulted in higher RMS and R_a values indicating much rougher surfaces. Thus, when strong adhesive bonds were broken, the resulting fracture surfaces were often extremely rough. It indicated the improved diffusion. The roughness parameters were similar for samples sealed at the T_{pi} for every dwell time, although T_{pi} is different for every dwell time. This finding indicated that the diffusion reached to a certain point at T_{pi} regardless of the process conditions.

In the second phase we fundamentally studied the role of molecular architecture on adhesion strength of variety of neat polyethylene resins. By increasing M_w , adhesion strength improved for linear metallocene ethylene α -olefins containing similar amount of SCB. For the metallocene ethylene α -olefins the presence of sparsely LCB resulted in lower adhesion strength compare to the linear chains. The presence of LCB hindered the reptation motion and diffusion. Highly branched chains of low density polyethylene (LDPE) yielded to a very low self-adhesion. Therefore, our findings indicated that the increase of LCB content results in decrease of interfacial self-adhesion strength.

The adhesion strength of Ziegler-Natta LLDPE was drastically lower than and metallocene catalyzed LLDPE. This was explained to be attributed to the homogeneous composition distribution of metallocene catalysed resins versus heterogeneous composition distribution of Zigler-Natta catalyzed resins. The enrichment of highly branched low molecular weight chains at the surfaces of the films in polymers with heterogeneous composition distribution was explained to cause this phenomenon. The segregated layer of short chains in the surface, which is doubled upon sealing, is less effective in strengthening the interface than the long chains.

The role of different molecular structures of polyethylene was studied through the designated blend compositions of conventional and metallocene catalyzed polyethylene resins. Four binary blends were prepared: LLDPE/Linear metallocene, LLDPE/LCB contain metallocene, LDPE/Linear metallocene, LDPE/LCB contain metallocene. Incorporation of metallocene resins to both LD and LL enhanced the adhesion strength. By increasing metallocene content to 40wt% and 60wt% the adhesion strength increased for all four types of the blends. However, the % increase of adhesion strength varied among different types of blends. All compositions of LL/Linear-m and LD/Linear-m blends showed higher % increase in adhesion strength compared

to their counterparts in LL/LCB-m and LL/LCB-m blends. These results indicated that the Linear-m was more effective than LCB-m in term of enhancement in adhesion strength. Moreover, adhesion strength improvement for all of the compositions of LDPE was higher than their LLDPE counterparts. This suggested that both Linear-m and LCB-m provided a faster interdiffusion in LDPE blends.

The broad adhesion strength plateau of LDPE as highly LCB containing resin illustrated the important role of high relaxation time of chains and high level of entanglements in melt behaviour. Based on the correlations, melt toughness in extensional flow could provide a basis for prediction of relative plateau broadness of a polymer. The broad adhesion strength plateau was also observed for metallocene resins, which was attributed to their high melt toughness.

8.2 Original contributions

- The concurrent effects of heat seal temperature-dwell time and temperature-pressure on seal were illustrated in 3D maps, a methodology that is easily applicable to other materials. Our results indicated that the time and temperature dependence of seal strength is a consequence of the fraction of amorphous phase available at the interface of films. We found that the seal strength has a linear correlation with the square root of sealing time, particularly at short sealing times, which complements literature results for long welding models.
- A fundamental study was performed on the effect of different molecular architectures on hot tack performance of resins from the polyethylene family. To our best knowledge, this has not been addressed in the literature. Our findings indicated that by increasing M_w , the hot tack (self-adhesion strength) improved for linear metallocene ethylene α -olefins. Also, the presence of even low amount of LCB results in a decrease of interfacial self-adhesion strength.
- The lower adhesion strength of conventional PEs than metallocene catalyzed PEs was explained to be attributed to the homogeneous composition distribution of metallocene catalysed resins versus heterogeneous composition distribution of conventional PEs.
- Our findings indicated that the Linear-metallocene resins was more effective than LCB-metallocene in term of enhancement in adhesion strength. The broadness of adhesion

strength plateau depended on the melt toughness obtained from extensional rheological properties and which could provide a basis for prediction of relative plateau broadness of a polymer.

8.3 Recommendations

In the previous section, we summarized what has been accomplished in the current work concerning the optimization of process parameters and material characteristics for heat seal process. The following unexplored aspects are recommended for the continuation of this work and future research:

1. To optimize seal process parameters in hot tack performance of polyethylene resins. Since the trend of dwell time, pressure and temperature dependency of hot tack might differ from seal strength, it is important to optimize hot tack based on these process parameters
2. To model heat transfer from heated bars to the film interfaces. It is important to estimate the required time for heat transfer across the film thickness. It is necessary to consider the phase change of material from solid to melt in heat sealing. In modeling of heat transfer, the consumption of energy for melting the crystals and the change in heat capacity of material from solid to melt should be considered.
3. To study the effect of molecular architecture on the hot tack after variety of delay times. The seal strength is measured after complete cooling while hot tack is measured instantly after heat sealing. However, in some industrial heat sealing processes the product fell into the package after a certain delay time which is normally a fraction of a second. This delay time enables partial crystallization which varies for different polymers.
4. To design metallocene and conventional polyethylene resins in a multilayer seal layer. In a design that the metallocene resin covers a thin layer of a few micron at the surface of seal layer. In order to avoid the influence of the segregated layer of highly branched low molecular weight chains of heterogeneous conventional polyethylene resins. To investigate the performance of LCB or linear metallocene at the surface of seal layer.
5. To improve the sustainability by reducing the thickness of seal layer. The possible alternatives could be neat metallocene resins or using blends of different types of metallocene together with different compositions.

BIBLIOGRAPHIE

- Basin, V.E., 1967. Role of diffusion in the formation of adhesion bonds between polymers. *Polym. Mech.* 3, 561–563. doi:10.1007/BF00859242
- Bastien, L.J., Gillespie, J.W., 1991. A non-isothermal healing model for strength and toughness of fusion bonded joints of amorphous thermoplastics. *Polym. Eng. Sci.* 31, 1720–1730. doi:10.1002/pen.760312406
- Benkoski, J.J., Fredrickson, G.H., Kramer, E.J., 2002. Model for the fracture energy of glassy polymer–polymer interfaces. *J. Polym. Sci. Part B Polym. Phys.* 40, 2377–2386. doi:10.1002/polb.10288
- Boiko, Y.M., Guérin, G., Marikhin, V.A., Prud'homme, R.E., 2001. Healing of interfaces of amorphous and semi-crystalline poly(ethylene terephthalate) in the vicinity of the glass transition temperature. *Polymer* 42, 8695–8702. doi:10.1016/S0032-3861(01)00406-2
- Boiko, Y.M., Lyngaae, J., 2005. Fracture energy–fracture stress relationship for weak polymer–polymer interfaces. *Polymer* 46, 6016–6024. doi:10.1016/j.polymer.2005.05.064
- Boiko, Y.M., Prud'homme, R.E., 1998. Morphology of fractured polymer surfaces self-bonded below the glass transition temperature. *Mech. Compos. Mater.* 34, 473–482. doi:10.1007/BF02254711
- Boiko, Y.M., Prud'homme, R.E., 1999. Interdiffusion and adhesion at the interface of a polystyrene-poly(2,6-dimethyl-1,4-phenylene oxide) blend below the glass transition temperature. *Mech. Compos. Mater.* 35, 441–446. doi:10.1007/BF02329331
- Brown, 1991a. The Adhesion Between Polymers. *Annu. Rev. Mater. Sci.* 21, 463–489. doi:10.1146/annurev.ms.21.080191.002335
- Brown, 1991b. A molecular interpretation of the toughness of glassy polymers. *Macromolecules* 24, 2752–2756. doi:10.1021/ma00010a018
- Brown, H.R., 1989. Effect of a diblock copolymer on the adhesion between incompatible polymers. *Macromolecules* 22, 2859–2860. doi:10.1021/ma00196a059
- Brown, H.R., 1990. Mixed-mode effects on the toughness of polymer interfaces. *J. Mater. Sci.* 25, 2791–2794. doi:10.1007/BF00584881
- Brown, H.R., 2001. Relation between the Width of an Interface between Two Polymers and Its Toughness. *Macromolecules* 34, 3720–3724. doi:10.1021/ma991821v
- Brydson, J.A., 1999. *Plastics Materials*. Butterworth-Heinemann.
- Cherry, B.W., 1981. *Polymer surfaces*. Cambridge University Press, Cambridge [Eng.]; New York.
- Chum, P.S., Kruper, W.J., Guest, M.J., 2000. Materials Properties Derived from INSITE Metallocene Catalysts. *Adv. Mater.* 12, 1759–1767. doi:10.1002/1521-4095(200012)12:23<1759::AID-ADMA1759>3.0.CO;2-7
- Coles, R., McDowell, D., Kirwan, M.J., 2003. *Food packaging technology*. CRC Press.

- Colls, N., Torres, A., Méndez, F., 2006. Material Selector for LLDPE/LDPE Blends for FFS Applications. *J. Plast. Film Sheeting* 22, 19–27. doi:10.1177/8756087906062563
- Creton, C., Kramer, E.J., Hui, C.Y., Brown, H.R., 1992. Failure mechanisms of polymer interfaces reinforced with block copolymers. *Macromolecules* 25, 3075–3088. doi:10.1021/ma00038a010
- De Gennes, P.G., 1971. Reptation of a Polymer Chain in the Presence of Fixed Obstacles. *J. Chem. Phys.* 55, 572–579. doi:10.1063/1.1675789
- De Gennes, P.-G., 1979. *Scaling Concepts in Polymer Physics*. Cornell University Press, Ithaca, NY.
- De Gennes, P.-G., 1983. Entangled polymers. *Phys. Today* 36, 33–39. doi:10.1063/1.2915700
- Doi, M., Edwards, S.F., 1978a. Dynamics of rod-like macromolecules in concentrated solution. Part 1. *J. Chem. Soc. Faraday Trans. 2 Mol. Chem. Phys.* 74, 560–570. doi:10.1039/F29787400560
- Doi, M., Edwards, S.F., 1978b. Dynamics of concentrated polymer systems. Part 2.—Molecular motion under flow. *J. Chem. Soc. Faraday Trans. 2 Mol. Chem. Phys.* 74, 1802–1817. doi:10.1039/F29787401802
- Doi, M., Edwards, S.F., 1978c. Dynamics of concentrated polymer systems. Part 3.—The constitutive equation. *J. Chem. Soc. Faraday Trans. 2 Mol. Chem. Phys.* 74, 1818–1832. doi:10.1039/F29787401818
- Doi, M., Edwards, S.F., 1979. Dynamics of concentrated polymer systems. Part 4.—Rheological properties. *J. Chem. Soc. Faraday Trans. 2 Mol. Chem. Phys.* 75, 38–54. doi:10.1039/F29797500038
- Donovan, K.M., Kong, D.-C., Liu, L., Sexton, D.F., Su, T.-K., 1999. United States Patent: 5888648 - Multi-layer hermetically sealable film and method of making same. 5888648.
- F02 Committee, 1998. ASTM 1921-98 Test Methods for Hot Seal Strength (Hot Tack) of Thermoplastic Polymers and Blends Comprising the Sealing Surfaces of Flexible Webs. ASTM International.
- F02 Committee, 2000. ASTM F88 Test Method for Seal Strength of Flexible Barrier Materials. ASTM International.
- Farley, J.M., Meka, P., Stehling, F.C., Trudell, B.C., Kurtzman, M.B., 1996. United States Patent: 5530065 - Heat sealable films and articles made therefrom. 5530065.
- Feldman, D., 1996. *Synthetic Polymers: Technology, Properties, Applications*. Springer.
- Foster, K.L., Wool, R.P., 1991. Strength of polystyrene-poly(methyl methacrylate) interfaces. *Macromolecules* 24, 1397–1403. doi:10.1021/ma00006a028
- Fowler, M.E., Barlow, J.W., Paul, D.R., 1987. Kinetics of adhesion development at PMMA-SAN interfaces. *Polymer* 28, 2145–2150. doi:10.1016/0032-3861(87)90056-5
- Frederix, C., Beauchene, P., Seguela, R., Lefebvre, J.M., 2013. Kinetics of the non-isothermal fusion-welding of unlike ethylene copolymers over a wide crystallinity range. *Polymer* 54, 2755–2763. doi:10.1016/j.polymer.2013.03.038

- Gardon, J.L., 1963a. Peel adhesion. I. Some phenomenological aspects of the test. *J. Appl. Polym. Sci.* 7, 625–641. doi:10.1002/app.1963.070070219
- Gardon, J.L., 1963b. Peel adhesion. II. A theoretical analysis. *J. Appl. Polym. Sci.* 7, 643–665. doi:10.1002/app.1963.070070220
- Gent, A.N., Kim, E.-G., Ye, P., 1997. Autohesion of crosslinked polyethylene. *J. Polym. Sci. Part B Polym. Phys.* 35, 615–622. doi:10.1002/(SICI)1099-0488(199703)35:4<615::AID-POLB9>3.0.CO;2-O
- Halle, R.W., 1997. Use of Ethylene Plastomers to Improve Polyethylene Films, in: 1997 Polymers, Laminations & Coatings Conference. Presented at the TAPPI Polymers, Laminations & Coatings Conference.
- Halle, R.W., 2003. Plastomer-mVLDPE Blends for High Performance Heat Sealing Applications, in: TAPPI 2003 Conference. Presented at the TAPPI Conference, Orlando, Florida, pp. 7–45.
- Halle, R.W., Davis, D.S., 1995. Heat sealing linear ethylene plastomers to ionomers or LLDPEs. *Tappi J.* 78, 200–206.
- Hamed, G.R., Shieh, C.-H., 1983. Relationship between the cohesive strength and the tack of elastomers. *J. Polym. Sci. Polym. Phys. Ed.* 21, 1415–1425. doi:10.1002/pol.1983.180210812
- Hanlon, J.F., Kelsey, R.J., Forcinio, H.E., 1998. Handbook of package engineering. CRC.
- Hashimoto, Y., Ishiaku, U. s., Leong, Y. w., Hamada, H., Tsujii, T., 2006. Effect of heat-sealing temperature on the failure criteria of oriented polypropylene/cast polypropylene heat seal. *Polym. Eng. Sci.* 46, 205–214. doi:10.1002/pen.20452
- Hassan, A., 2007. Effect of bar sealing parameter on OPP/MCPP heat seal strength. *Express Polym. Lett.* 1, 773–779.
- Hosoda, S., 1988. Structural Distribution of Linear Low-Density Polyethylenes. *Polym. J.* 20, 383–397. doi:10.1295/polymj.20.383
- Jordens, K., Wilkes, G.L., Janzen, J., Rohlfing, D.C., Welch, M.B., 2000. The influence of molecular weight and thermal history on the thermal, rheological, and mechanical properties of metallocene-catalyzed linear polyethylenes. *Polymer* 41, 7175–7192. doi:10.1016/S0032-3861(00)00073-2
- Kausch, H.H., Tirrell, M., 1989. Polymer Interdiffusion. *Annu. Rev. Mater. Sci.* 19, 341–377. doi:10.1146/annurev.ms.19.080189.002013
- Khare, A.R., Westphal, S.P., Ling, M.T.K., Qin, C., Woo, L., 2000. Thermal and dynamic mechanical analysis on metallocene ULDPE/PP blends to optimize impact properties. *Thermochim. Acta* 357–358, 155–160. doi:10.1016/S0040-6031(00)00384-1
- Kim, K.D., Sperling, L.H., Klein, A., Hammouda, B., 1994. Reptation Time, Temperature, and Cosurfactant Effects on the Molecular Interdiffusion Rate during Polystyrene Latex Film Formation. *Macromolecules* 27, 6841–6850. doi:10.1021/ma00101a024
- Klein, J., 1990. The Interdiffusion of Polymers. *Science* 250, 640–646. doi:10.1126/science.250.4981.640

- Kline, D.B., Wool, R.P., 1988. Polymer welding relations investigated by a lap shear joint method. *Polym. Eng. Sci.* 28, 52–57. doi:10.1002/pen.760280109
- Kramer, E.J., 1983. Microscopic and molecular fundamentals of crazing, in: Kausch, H.H. (Ed.), *Crazing in Polymers*, *Advances in Polymer Science*. Springer Berlin Heidelberg, pp. 1–56.
- Kramer, E.J., Berger, L.L., 1990. Fundamental processes of craze growth and fracture, in: Kausch, H.-H. (Ed.), *Crazing in Polymers Vol. 2*, *Advances in Polymer Science*. Springer Berlin Heidelberg, pp. 1–68.
- Kunz, K., Stamm, M., 1996. Initial Stages of Interdiffusion of PMMA across an Interface. *Macromolecules* 29, 2548–2554. doi:10.1021/ma950187s
- Lee, L.H., 1991. *Fundamentals of Adhesion*. Springer.
- Malsen, J. van, Tenpierik, M.J., Looman, R.H.J., Cauberg, J.J.M., 2008. Heat Seal Strength of Barrier Films Used in Vacuum Insulation Panels At Room Temperature and At -130°C. *J. Plast. Film Sheeting* 24, 35–52. doi:10.1177/8756087908089092
- Manaure, A.C., Morales, R.A., Sánchez, J.J., Müller, A.J., 1997. Rheological and calorimetric evidences of the fractionated crystallization of iPP dispersed in ethylene/ α -olefin copolymers. *J. Appl. Polym. Sci.* 66, 2481–2493. doi:10.1002/(SICI)1097-4628(19971226)66:13<2481::AID-APP11>3.0.CO;2-0
- Manaure, A.C., Müller, A.J., 2000. Nucleation and crystallization of blends of poly(propylene) and ethylene/ α -olefin copolymers. *Macromol. Chem. Phys.* 201, 958–972. doi:10.1002/1521-3935(20000601)201:9<958::AID-MACP958>3.0.CO;2-0
- Meka, P., Stehling, F.C., 1994. Heat sealing of semicrystalline polymer films. I. Calculation and measurement of interfacial temperatures: Effect of process variables on seal properties. *J. Appl. Polym. Sci.* 51, 89–103.
- Mesnil, P., Arnauts, J., Halle, R.W., Rohse, N., 2000. Seal Through Contamination Performance of Metallocene Plastomers. Presented at the TAPPI POLYMERS LAMINATIONS AND COATINGS CONFERENCE, pp. 669–686.
- Miller, P., Buckley, D.J., Kramer, E.J., 1991. Microstructure and origin of cross-tie fibrils in crazes. *J. Mater. Sci.* 26, 4445–4454. doi:10.1007/BF00543666
- Mirabella, F.M., Ford, E.A., 1987. Characterization of linear low-density polyethylene: Cross-fractionation according to copolymer composition and molecular weight. *J. Polym. Sci. Part B Polym. Phys.* 25, 777–790. doi:10.1002/polb.1987.090250407
- Miyata, K., Toshiyuki, H., 2011. EFFECT OF LLDPE CONTENT ON HEAT SEAL PROPERTIES FOR HDPE/LLDPE FILM, in: ANTEC 2011 *Plastics: Annual Technical Conference Proceedings*. Presented at the ANTEC 2011.
- Morris, B.A., 2002. Predicting the Heat Seal Performance of Ionomer Films. *J. Plast. Film Sheeting* 18, 157–167. doi:10.1177/8756087902018003002
- Mueller, C., Capaccio, G., Hiltner, A., Baer, E., 1998. Heat sealing of LLDPE: relationships to melting and interdiffusion. *J. Appl. Polym. Sci.* 70, 2021–2030.

- Nase, M., Langer, B., Grellmann, W., 2009. Influence of Processing Conditions On the Peel Behavior of Polyethylene/ Polybutene-1 Peel Systems. *J. Plast. Film Sheeting* 25, 61–80. doi:10.1177/8756087909343139
- Nitta, K., Suzuki, K., Tanaka, A., 2000. Comparison of tensile properties in the pre-yield region of metallocene-catalyzed and Ziegler-Natta-catalyzed linear polyethylenes. *J. Mater. Sci.* 35, 2719–2727. doi:10.1023/A:1004710127185
- Nitta, K., Tanaka, A., 2001. Dynamic mechanical properties of metallocene catalyzed linear polyethylenes. *Polymer* 42, 1219–1226. doi:10.1016/S0032-3861(00)00418-3
- Poisson, Hervais, V., Lacrampe, M.F., Krawczak, P., 2006a. Optimization of PE/Binder/PA extrusion blow molded films. I. Heat sealing ability improvement using PE/EVA blends. *J. Appl. Polym. Sci.* 99, 974–985.
- Poisson, Hervais, V., Lacrampe, M.F., Krawczak, P., 2006b. Optimization of PE/binder/PA extrusion blow molded films. II. Adhesion properties improvement using binder/EVA blends. *J. Appl. Polym. Sci.* 101, 118–127.
- Prager, S., Tirrell, M., 1981. The healing process at polymer–polymer interfaces. *J. Chem. Phys.* 75, 5194–5198. doi:10.1063/1.441871
- Razavi-Nouri, M., Hay, J.N., 2001. Thermal and dynamic mechanical properties of metallocene polyethylene. *Polymer* 42, 8621–8627. doi:10.1016/S0032-3861(01)00377-9
- Russell, T.P., Deline, V.R., Dozier, W.D., Felcher, G.P., Agrawal, G., Wool, R.P., Mays, J.W., 1993. Direct observation of reptation at polymer interfaces. *Nature* 365, 235–237. doi:10.1038/365235a0
- Schnell, R., Stamm, M., Creton, C., 1998. Direct Correlation between Interfacial Width and Adhesion in Glassy Polymers. *Macromolecules* 31, 2284–2292. doi:10.1021/ma971020x
- Schnell, R., Stamm, M., Creton, C., 1999. Mechanical Properties of Homopolymer Interfaces: Transition from Simple Pullout To Crazing with Increasing Interfacial Width. *Macromolecules* 32, 3420–3425. doi:10.1021/ma980860o
- Schweizer, K.S., 1989. Mode-coupling theory of the dynamics of polymer liquids: Qualitative predictions for flexible chain and ring melts. *J. Chem. Phys.* 91, 5822–5839. doi:10.1063/1.457534
- Selke, S.E.M., Culter, J.D., Hernandez, R.J., 2004. *Plastics Packaging: Properties, Processing, Applications, And Regulations*, 2nd ed. Hanser Gardner Publications.
- Shanks, R.A., Li, J., Yu, L., 2000. Polypropylene–polyethylene blend morphology controlled by time–temperature–miscibility. *Polymer* 41, 2133–2139. doi:10.1016/S0032-3861(99)00399-7
- Shih, H.H., Wong, C.M., Wang, Y.C., Huang, C.J., Wu, C.C., 1999. Hot tack of metallocene catalyzed polyethylene and low-density polyethylene blend. *J. Appl. Polym. Sci.* 73, 1769–1773.
- Smith, G.D., Plummer, C.J.G., Bourban, P.-E., Manson, J.-A.E., 2001. Non-isothermal fusion bonding of polypropylene. *Polymer* 42, 6247–6257. doi:10.1016/S0032-3861(01)00060-X

- Stamm, M., Hüttenbach, S., Reiter, G., Springer, T., 1991. Initial Stages of Polymer Interdiffusion Studied by Neutron Reflectometry. *EPL Europhys. Lett.* 14, 451. doi:10.1209/0295-5075/14/5/011
- Stehling, F.C., Meka, P., 1994. Heat sealing of semicrystalline polymer films. II. Effect of melting distribution on heat-sealing behavior of polyolefins. *J. Appl. Polym. Sci.* 51, 105–119.
- Stokes, V.K., 1989. Joining methods for plastics and plastic composites: an overview. *Polym. Eng. Sci.* 29, 1310–1324.
- Tanrattanakul, V., Udomkichdecha, W., 2001. Development of novel elastomeric blends containing natural rubber and ultra-low-density polyethylene. *J. Appl. Polym. Sci.* 82, 650–660. doi:10.1002/app.1893
- Tetsuya, T., Hashimoto, Y., Ishiaku, U.S., Mizoguchi, M., Leong, Y.W., Hamada, H., 2006. Effect of heat-sealing temperature on the properties of OPP/CPP heat seals. Part II. Crystallinity and thermomechanical properties. *J. Appl. Polym. Sci.* 99, 513–519.
- Tetsuya, T., Ishiaku, U.S., Mizoguchi, M., Hamada, H., 2005. The effect of heat sealing temperature on the properties of OPP/CPP heat seal. I. Mechanical properties. *J. Appl. Polym. Sci.* 97, 753–760.
- Theller, H.W., 1989. Heatsealability of Flexible Web Materials in Hot-Bar Sealing Applications. *J. Plast. Film Sheeting* 5, 66–93. doi:10.1177/875608798900500107
- Troughton, M.J., 2008. *Handbook of Plastics Joining: A Practical Guide*. Cambridge University Press.
- Van Loon, A.J., Tierens, M.D., Permentier, D.J., 2009. United States Patent: 7588706 - Multilayer films with improved properties. 7588706.
- Vega, J.F., Muñoz-Escalona, A., Santamaría, A., Muñoz, M.E., Lafuente, P., 1996. Comparison of the Rheological Properties of Metallocene-Catalyzed and Conventional High-Density Polyethylenes. *Macromolecules* 29, 960–965. doi:10.1021/ma9504633
- Voyutskii, S.S., Kamenskii, A.N., Fodiman, N.M., 1966. Direct proofs of self- and mutual diffusion in the formation of adhesion bonds between polymers. *Polym. Mech.* 2, 279–283. doi:10.1007/BF00860301
- Wagner, J.R., 2009. *Multilayer Flexible Packaging: Technology and Applications for the Food, Personal Care, and Over-the-Counter Pharmaceutical Industries*. William Andrew.
- Wang, W.-J., Ye, Z., Fan, H., Li, B.-G., Zhu, S., 2004. Dynamic mechanical and rheological properties of metallocene-catalyzed long-chain-branched ethylene/propylene copolymers. *Polymer* 45, 5497–5504. doi:10.1016/j.polymer.2004.05.053
- Wild, L., Ryle, T.R., Knobloch, D.C., Peat, I.R., 1982. Determination of branching distributions in polyethylene and ethylene copolymers. *J. Polym. Sci. Polym. Phys. Ed.* 20, 441–455. doi:10.1002/pol.1982.180200307
- Wood-Adams, P.M., 1998. The effect of long chain branching on the rheological behavior of polyethylenes synthesized using constrained geometry and metallocene catalysts (Ph.D.Thesis). McGill University, Canada.

- Wool, R.P., 1995. *Polymer Interfaces: Structure and Strength*. Hanser-Gardner Publications.
- Wool, R.P., 2006. Adhesion at polymer–polymer interfaces: a rigidity percolation approach. *Comptes Rendus Chim.* 9, 25–44. doi:10.1016/j.crci.2005.04.008
- Wool, R.P., 2008. Self-healing materials: a review. *Soft Matter* 4, 400–418. doi:10.1039/b711716g
- Wool, R.P., O'Connor, K.M., 1981. A theory crack healing in polymers. *J. Appl. Phys.* 52, 5953–5963. doi:10.1063/1.328526
- Wool, R.P., Yuan, B.-L., McGarel, O.J., 1989. Welding of polymer interfaces. *Polym. Eng. Sci.* 29, 1340–1367. doi:10.1002/pen.760291906
- Xu, D.-B., Hui, C.-Y., Kramer, E.J., Creton, C., 1991. A micromechanical model of crack growth along polymer interfaces. *Mech. Mater.* 11, 257–268. doi:10.1016/0167-6636(91)90007-M
- Xue, Y.-Q., Tervoort, T.A., Lemstra, P.J., 1998. Welding behavior of semicrystalline polymers. 1. The effect of nonequilibrium chain conformations on autoadhesion of UHMWPE. *Macromolecules* 31, 3075–3080.
- Xue, Y.-Q., Tervoort, T.A., Rastogi, S., Lemstra, J., 2000. Welding Behavior of Semicrystalline Polymers. 2. Effect of CocrySTALLIZATION on Autoadhesion. *Macromolecules* 33, 7084–7087. doi:10.1021/ma000754y
- Yoo, J.N., Sperling, L.H., Glinka, C.J., Klein, A., 1991. Characterization of film formation from polystyrene latex particles via SANS. 2. High molecular weight. *Macromolecules* 24, 2868–2876. doi:10.1021/ma00010a036
- Zhang, M.Q., Rong, M.Z., 2012. Theoretical consideration and modeling of self-healing polymers. *J. Polym. Sci. Part B Polym. Phys.* 50, 229–241. doi:10.1002/polb.22387
- Zhao, W., Zhao, X., Rafailovich, M.H., Sokolov, J., Composto, R.J., Smith, S.D., Russell, T.P., Dozier, W.D., Mansfield, T., Satkowski, M., 1993. Segregation of chain ends to polymer melt surfaces and interfaces. *Macromolecules* 26, 561–562. doi:10.1021/ma00055a026

APPENDIX A

Article 4: Sealability and Seal Characteristics of PE/EVA and PLA/PCL Blends*

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1. Abstract

Seal strength behaviour of low density polyethylene and ethylene vinyl acetate copolymer (PE/EVA) blends as well as that of blends of a seal grade PLA with aliphatic polyester (PCL) was studied. Polyethylene is commonly used for seal application in packaging multilayer structures and amorphous PLA is considered to be its counterpart for compostable and/or biodegradable ones. Incorporation of EVA in polyethylene improves its sealability in terms of a decrease in seal initiation temperature and broadness of sealability plateau. This was interpreted as due to the formation of finer crystals, a decrease in the melting point and presence of vinyl acetate polar group. These were supported by results obtained from differential scanning calorimetry (DSC) and Scanning electron microscopy (SEM). For the PLA/PCL system, the dispersed phase was stretched into elongated ellipsoidal domains. This type of morphology affected the mechanical and seal properties of the blends. As a result of blending, both hot-tack initiation temperature and strength as well as seal initiation temperature were enhanced. The enhancement in these seal properties was significant when the concentration of the dispersed phase exceeded 20wt% in the blend. Hot-tack strength of up to twice of pure PLA was achieved through blending. This was attributed to the lower glass transition temperature of PCL, resulting in enhanced mobility of PLA chains and also the high aspect ratio of the dispersed phase. The maximum obtained hot-tack strength (1200 g/25mm) at 40% dispersed content compared advantageously to commercially available polyolefin based sealant resins. The seal and hot-tack initiation temperatures were shifted to lower temperatures by as much as 30 °C, which can allow faster and more energy efficient sealing process.

* Published in International Polymer Processing, 2014, Vol. 29, No. 1, pp. 95-102.

Keywords: Seal strength, hot tack, EVA, PLA/PCL blend, flexible packaging;

2. Introduction

Heat sealing is an important operation in packaging. Modern vertical and horizontal form fill seal (VFFS and HFFS) machines have a need for operating at higher speeds. This requirement and the fact that the final package integrity is ultimately dependent on the results of the sealing process requires that the polymers used for the seal layer have superior hot-tack strength, low seal initiation temperature and wider sealing temperatures range. Although resin suppliers provide a wide range of resins for the seal layer, with a variety of performances and physical properties, materials optimization is still a challenge. This challenge is even greater when the production line aims to use biodegradable materials which still cannot provide as superior properties as commercially available resins. Achieving sufficient adhesion upon sealing two semicrystalline polymer films in the fully or partially molten state is mainly ascribed to chain interdiffusion across the interface, amorphous fraction of material, and polymer chain functional groups.

Coextrusion of low density polyethylene (LDPE) and ethylene vinyl acetate copolymer (EVA) has been used in industry to tailor a sealant layer. However, studies on correlation of physical properties and final seal performance of LDPE/EVA blends in open literature are limited. Studies on PE/EVA blends in a multilayer structure showed the effect of incorporating a low content vinyl acetate EVA (9%) in seal layer and tie layer. (Poisson et al. 2006b; Poisson et al. 2006a; Zhang et al. 2009) In this work we investigate the effect of EVA incorporation in monolayer PE sealant for several compositions produced by the film blowing process. And the consequences of this blending on microstructure and physical properties of the films and their correlation to the final seal properties are established.

On the other hand, many efforts and energy are used to overcome the problems caused by plastic wastes, particularly those produced every year by the packaging industry. Increasing costs for the removal, incineration and landfill of municipal wastes urge the packaging industry to shift their productions to more sustainable packages. (Brogly, Nardin, and Schultz 1997) In this study, we also consider the applicability of biodegradable resins for multilayer packaging, and in particular for seal layer.

Some bio-based resins such as Polylactides (PLA) and Polyhydroxyalkanoate (PHA) have good moisture barrier but low flexibility, while others such as Polyvinylalcohol (PVA, PVOH) have significantly higher oxygen barrier, but are highly sensitive to humidity. (Mittal and Pizzi 2009) However, all these resin have a limited capability to be heat sealed at an acceptable dwell time and low enough temperature not to deteriorate other properties of the package. This hermetic seal property is a guaranty of the efficiency of barrier layers against oxygen permeation, odor loss and water vapor transmission. Among bio-based and compostable and/or biodegradable polymers that can be used as a hermetic seal layer are Polybutylene succinate (PBS) and amorphous PLA grade. PBS lacks seal strength and PLA has low flexibility and high seal initiation temperature. (Ichikawa and Mizukoshi 2011) In this study, another focus will be on the improvement of PLA seal properties through blending with another more flexible biodegradable and/or compostable resin such as polycaprolactone (PCL). This will be sought while maintaining reasonable mechanical properties and allowing the package to seal at temperatures as low as 60-90°C.

3. Materials and Methods

For LDPE/EVA blends, low density polyethylene (LDPE, MFI of 2.3) and ethylene vinyl acetate (EVA containing 18%VA, MFI of 8) were obtained from Dow Chemicals and Dupont respectively. The materials studied include pure EVA and blends with weight content of 60%, 40% and 20% of EVA and are designated as EVA100, EVA60, EVA40 and EVA20 respectively. For PLA/PCL blends, a seal grade PLA was obtained from Natureworks (grade is 4060D) and was a copolymer of D,L lactic acid, with D-monomer content of higher than 10%. An aliphatic polyester (polycaprolactone, PCL from Union Carbide with $M_w=80000$ g/mole) has been used to improve both flexibility and seal initiation temperature of the PLA sealant resin.

Melt blending of these materials needs a good understanding of melt behaviour and temperature dependency of melt properties. A technique usually used for this purpose is rheological characterization. Rheological properties can also be used in prediction of developed morphology in a specific stress field. Rheological characterization of the resins in our study was carried out using parallel-plate geometry on a constant strain rheometer (MCR 301 Anton Paar, Austria). The experiments were performed in the dynamic mode at 180°C for PE/EVA blends and 190 °C for PLA/PCL blends under a nitrogen atmosphere and in the frequency range of 0.1 to 100 Hz.

All LDPE/EVA samples have been melt blended directly into blown films using a 45 mm Killion single-screw extruder equipped with a helical blown film die. All the films have been drawn to a thickness of 40 μ m while circulating cooling air rest at room temperature.

For PLA/PCL blends, all samples have been melt blended directly into cast films using a twin-screw extruder (Leistritz model ZSE 18 HP co-rotating) equipped with a cast film die. All the films have been drawn to a thickness of 50 μ m for further analysis. The temperature profile of the extruder along the barrel was set at 170/180/190/190 and the die at a temperature of 190°C. Different blend compositions have been used to prepare the samples. The weight content of PCL in the blends was 40%, 20%, and 0%. The number following the sample name is the content of the dispersed phase (e.g. PLCL80-20 has 20% PCL in the blend) and in case of pure resin the samples have been coded as PLA or PCL.

As one of the main affecting factors in determination and control of seal properties is material's crystallinity before sealing; this content has to be determined precisely. Differential scanning calorimeter (DSC) has a capability to analyze size and content of crystals in different polymeric material both in granule form and as processed films. The TA Instruments Q1000 was used in our study to achieve this goal. The operating window of this instrument is between 2°C/min to 50°C/min. The heating rate for all the samples was set at 10°C/min and the temperature range adjusted from -20°C to 140°C. The melting peaks for polyethylene and EVA are found at around 110°C and 80°C respectively, and for PCL melting point is around 60°C. No melting point was observed for amorphous PLA.

Seal strength experiments were performed using SL100 Lako-Tool hot-tack machine based on the ASTM F88 and F2029 experimental methods. Heat seals were made with two metallic jaws (19.1mm \times 25.4mm) covered by Teflon on film strips in 1 inch width \times 13 inches length dimension. Biaxially oriented polypropylene tape was used to provide the support for PE/EVA sealant. For PLA/PCL samples, the support film was a semi-crystalline PLA with melting point of 168°C thermally laminated on the films. All the sealing experiments were performed in a range of jaw temperatures from 50°C to 140°C at a constant pressure of 0.5 (N/mm²) and dwell time of 0.5 (s).

Seal property, mechanical and thermal properties of any type of blend, will depend strongly on its morphology, in addition to the individual components. Scanning electron microscopy (SEM) is a

technique, which will allow the precise determination of blends morphology. For LDPE/EVA blends, the films were observed in the cross section. Prior to observation, the films were fractured under liquid nitrogen and the EVA phase extracted from the blends using xylene at 50°C for 5 hours. EVA component of the blend dissolves in the solvent and will be seen as black holes in the SEM images, while polyethylene remains unaffected. The etched surfaces, after proper drying, were gold sputtered and observed under a HITACHI S-4700 SEM. For PLA and PCL blends, film samples were molded in epoxy and then cryo-microtomed using a diamond knife and observed under the same SEM instrument.

4. Results and discussion for PE/EVA system

4.1. Differential Scanning Calorimetry (DSC) results

The results of thermal analysis using DSC are presented in Figure A- 1. It can be easily observed that an increase in the amount of EVA in the blend decreased crystallinity and broadened the range of crystal sizes.

Presence of vinyl acetate groups at crystal growth front can prevent polyethylene chains crystal growth in PE/EVA blends, which would reduce the lamellar thickness and crystal perfection. It also decreases the enthalpy of melting and shifts the melting point to lower temperatures. By increasing the EVA content, another melting peak appears at lower temperatures due to EVA chains crystal formation, which contain sequences of ethylene groups capable of forming fine lamellas. (Brogly, Nardin, and Schultz 1997)

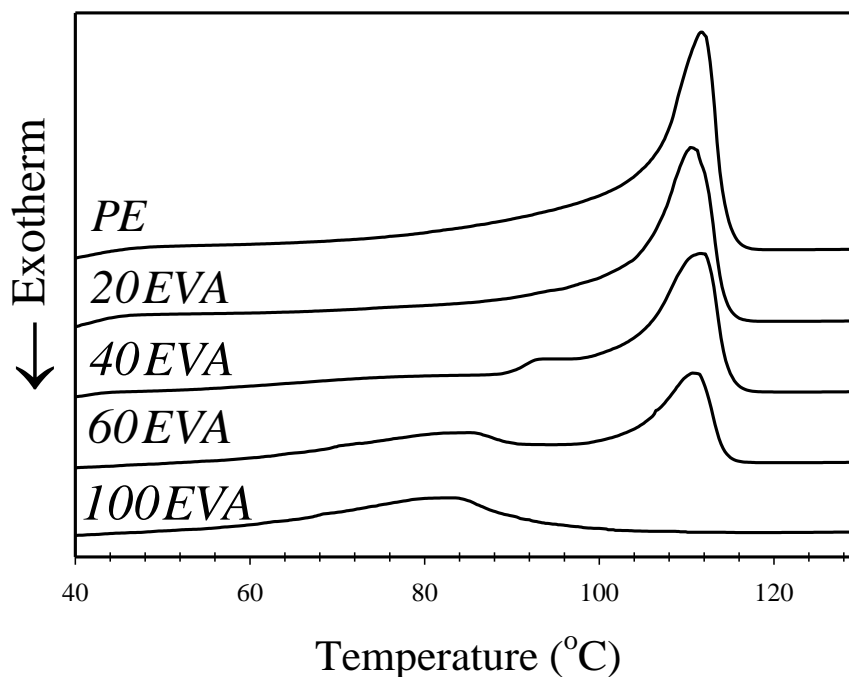


Figure A- 1 DSC thermographs of melting of films

4.2.Sealability results

In the heat sealing results of the blends, it is important to evaluate the evolution of the seal initiation temperature (T_{si}), plateau initiation temperatures (T_{pi}), sealability temperature window and plateau seal strengths for the various blends. The results are presented in Figure A- 2 and it can be seen that polyethylene shows $T_{si}=108^{\circ}\text{C}$ and $T_{pi}=115^{\circ}\text{C}$, while for neat EVA $T_{si}=74^{\circ}\text{C}$ and $T_{pi}=80^{\circ}\text{C}$ were observed. The results for the blends of these two resins are located in between. (Figure A- 2)

Incorporation of EVA in polyethylene causes a decrease in T_{si} and T_{pi} at all concentration. But from the results, one can observe two types of seal behaviour for the blends. Blends with lower concentration than 40% follow similar trend in sealing as pure polyethylene. For 60 % EVA content or more, the blends show a similar behaviour to pure EVA. The large difference in seal strength and seal initiation temperature between 40% and 60% EVA blends might be due to the dominance of EVA in 60% film. Vinyl acetate group polarity creates attraction force between the two films surfaces.

As represented in Figure A- 1, blending PE and EVA caused the formation of finer crystals which led to lower melting point for the blends. This would result in a lower seal initiation temperature and higher mobility of the chains in a set temperature in the blends with higher content of EVA. (Mittal and Pizzi 2009; Petrie 2000)

The work of Stehling and Meka showed that there is a direct relationship between the seal strength of a film and its yield strength. (Meka and Stehling 1994; Stehling and Meka 1994) By blending polyethylene with EVA, the yield strength decreases gradually. On the other hand, the higher mobility of EVA chains incorporated in blended films would result in higher diffusion and higher surface adhesion after sealing, which tend to increase the seal strength of blended samples. These two phenomena are acting in opposite direction. In the blends with EVA contents of 20% and 40%, where PE forms the continuous phase, the effect of polarity and inter-diffusion is more pronounced. In blends with 60% and higher content of EVA the continuous phase will change from PE to EVA and the yield strength of the blends decreases significantly. The latter overcome the effect of higher polarity and higher inter-diffusion of blends. This would result in a sharp decrease in the seal strength for the sample with 60% EVA and a plateau in seal strength of the samples afterward. The results we obtained in Figure A- 2 are in agreement with the trend they have explained which others also confirmed. (Halle 2003; Mueller et al. 1998)

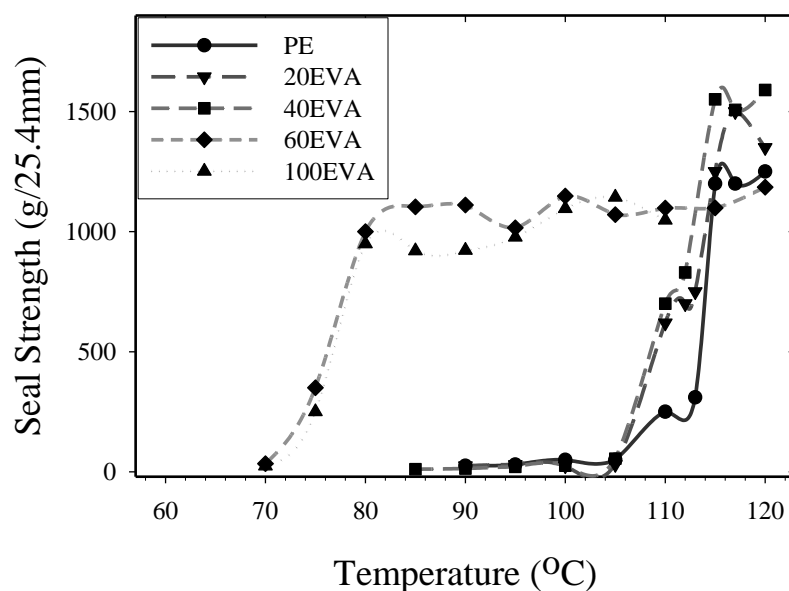


Figure A- 2 Seal Strength Temperature dependence of PE/EVA blends films

In failure mode analysis, three temperature zones were observed for each film, representing three different failure modes. As illustrated in Figure A- 3, peeling failure zone is the region in which the seal area peels apart at the film interface because of insufficient chain mobility. This lower chain mobility can be due to low sealing temperature difference to the melting point and would not allow the chains at the surface of the films to interdiffuse. In low EVA content blends (20%EVA and 40%EVA), this failure mode could be observed at 105°C to 112°C. These temperatures are very close to the melting point of polyethylene, which is the dominant phase. For 60%EVA blend, the peeling failure zone is close to the melting point of EVA.

Plateau zone, in which delamination and elongation failure occur, happen at the range of temperatures capable of melting crystals and providing enough interdiffusion and entanglements such that the seal area is strong. This plateau is usually expected at temperatures above the melting point of the dominant phase in the blend.

In 60%EVA blend this plateau is broad. Broadness of the plateau will provide us with the wider processing window, which facilitates industrial process optimization. In low content blend compositions (20%EVA and 40%EVA), the plateau is narrower and the operating range is more limited.

Breakage zone is where high temperature causes merging of the two films surfaces in a way that makes adhesive debonding impossible. This type of failure at very high temperature might cause pin holes and defects in other layers as well as the seal edges.

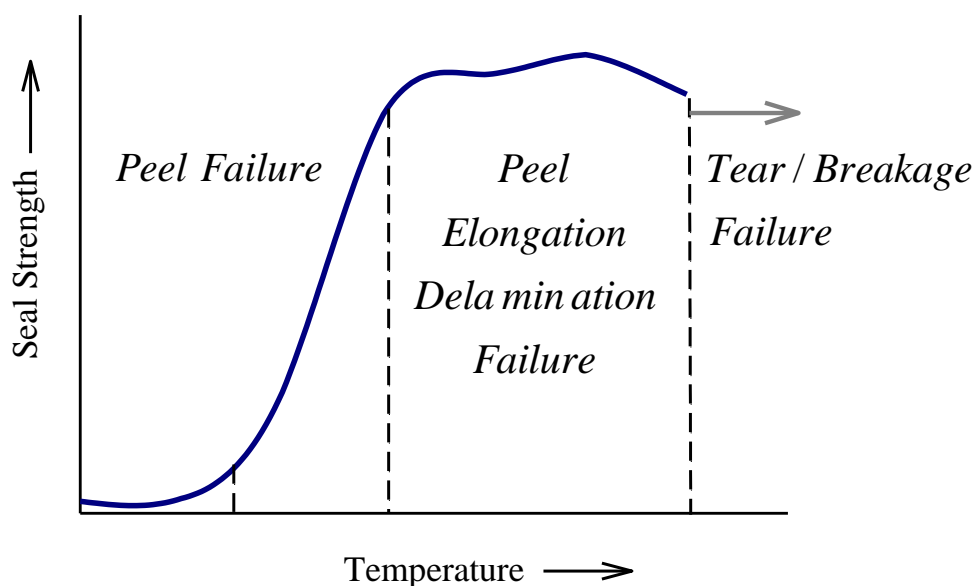


Figure A- 3 Failure modes observed in monolayer blown PE/EVA blend films

The term “hot-tack” represents strength of the seal while it is still molten and hot. It should be measured in the first few milliseconds after sealing. In the vertical form-fill and seal process, there are two seals, one in the bottom of the package and the other in the top. The package is usually filled right after the bottom seal is done, so while it is hot the seal has to withstand the product weight. To have an acceptable performance in industrial form-fill seal process, seal layer has to bear the weight of product immediately after seal formation. So the seal layer must be capable of making a seal strong enough to tolerate force and stress created by filling. Hot-tack plateau broadness indicates the range of temperatures in which heat sealing can be done to attain the maximum hot-tack achievable by the material. Since the seal jaws temperature in industrial process of form fill sealing shows usually significant fluctuations, the broadness of this plateau is crucial.

Figure A- 4 shows the hot-tack curves of the blends and clearly indicates the superior performance of high EVA content blends. In fact, these blends show lower hot-tack initiation temperature and higher level of hot-tack. Both 60%EVA and 100%EVA systems show their maximum hot-tack at 82°C. At temperatures below that, the seal peeled apart and the measured force is the strength of seal adhesion failure at the interface. It is to be noted that the final plateau temperature in 60%EVA (100°C) is much higher than for 100%EVA sample (90°C). The hot-tack

behaviour of the blends could be divided into EVA like and LDPE like behaviour, which is the same trend observed for seal curves of these resins. (Figure A- 2 and Figure A- 4)

The measured force drops at high temperatures. This is attributed to film breakage while the sealed part remains intact. At these high temperatures, the seal was strong enough not to peel apart, so the stress concentration on the edge of seal and thermal weakening of film cause film breakage in that area. Poor LDPE hot-tack strength represented in Figure A- 4 is well known in literature and industrial applications. EVA incorporation enhanced it to more than two times, although for 20%EVA and 40%EVA compositions the hot-tack strength is still lower than desirable hot-tack strength (above 600 (g/25.4mm)). This is probably due to the dominance of LDPE adhesion mechanism and low EVA amount in the blends.

As hot-tack and seal results are significantly controlled by the surface properties and blend composition, for deeper understanding of this behaviour, a morphological study of the blends is necessary. SEM micrographs are presented in Figure A- 5. The dominance of EVA on the surfaces of films having high EVA content could be the main reason that at their seal performance is enhanced significantly. The detailed analysis of the blend structure is given in the morphology section below.

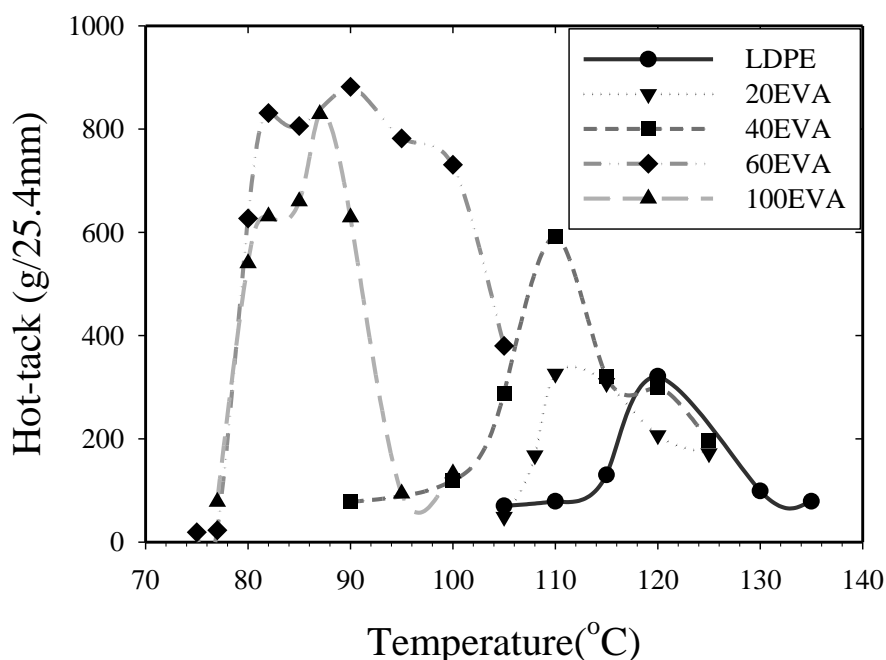


Figure A- 4 Hot-tack Temperature dependence of PE/EVA blend films

4.3. Blend morphology

Morphology analysis can provide us with a useful tool to explain the difference in the seal results of the two blend groups. The SEM images of etched cross section of 40%EVA and 60%EVA are shown in Figure A- 5 along with that of LDPE. The darker domains indicate the positions of the extracted EVA phase in both 60% and in 40%EVA. The extracted areas at the surface of the films indicate the presence of EVA at the surface. These domains were larger for 60%EVA compared to 40%EVA. For 60%EVA content, these domains cover most of the films surface.

Binary melt blends of several kinds of polyethylenes and EVAs with various vinyl acetate contents were extensively investigated and known to be partially compatible, leading to matrix disperse morphology at low contents and co-continuous morphology at high content of disperse phase. (Jin 2010) In most of the applications, small amounts of polyethylene grafted maleic anhydride were used as compatibilizer. (Moly et al. 2006) It should be mentioned here that it was not possible to add the PE-g-MA compatibilizer because of FDA restrictions concerning contact with food.

Finally, because of the large difference in viscosity of the resins, as illustrated in Figure A- 6, the morphology developed in the film blowing die could not be of the droplet-matrix type but rather co-continuous structure as reported in literature. (Chattopadhyay, Chaki, and Bhowmick 2001; Peón et al. 2003) The complex viscosity (η^*) of LDPE and EVA resins in Figure A- 6 shows a large difference in their viscosities at shear rates in the range of the film blowing process. The low molecular weight (low viscosity or high MFI) resin will go to the surface of the film in this case. It has also been reported that, in incompatible blends, chains with higher surface free energy or having polar groups have a preference to be located at the film surface. (Jalbert et al. 1993; H. Lee and Archer 2001; Chen and Gardella 1994; McNally 2005a; McNally 2005b) This higher concentration of EVA at the film surfaces causes superior hot-tack and seal strength results for high EVA content films. Polarity of vinyl acetate group in EVA, in addition to lower crystallinity, introduces the observed enhanced adhesion at the interface.

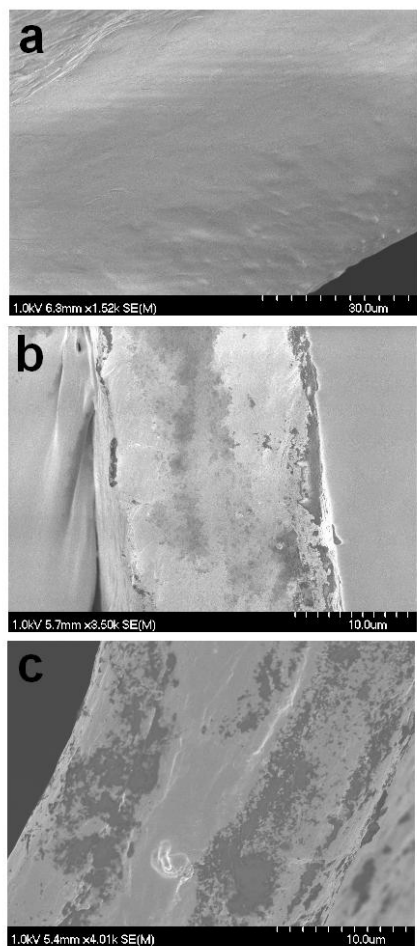


Figure A- 5 SEM images of film cross section after etching with xylene, (a) LDPE, (b) 40%EVA, (c) 60%EVA

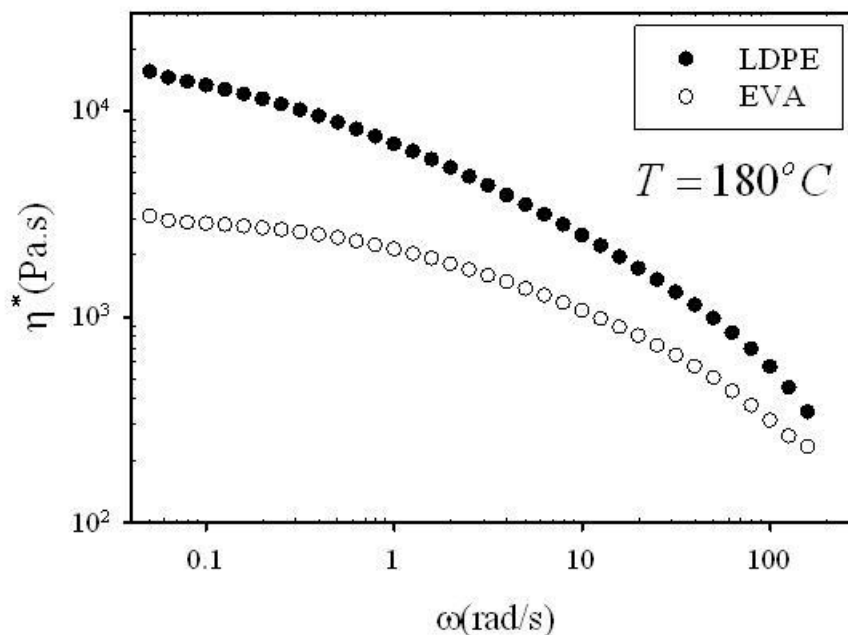


Figure A- 6 Frequency sweep of LDPE and EVA at 180°C

5. Results and discussion for PLA/PCL system

5.1. Viscoelastic Behaviour and Blends Morphology

The melt shear viscosity of PLA, PCL and blended samples was measured at 190°C and is illustrated in Figure A- 7. As discussed above, the rheological behaviour of the melts plays a determining role in the development of morphology in polymer blends. This morphology will affect later the mechanical and seal characteristics (particularly seal strength and hot tack) of the final films. In a blend of PLA and PCL, both resins are aliphatic polyesters which will favor interaction among them. (Sarazin, Roy, and Favis 2004) Knowing that the viscosity ratio plays a determining role in the formation of droplet disperse morphology or elongated disperse phase, depending on the type of flow during film processing using a cast die. (Kamal et al. 1995; S. Y. Lee and Kim 1997) As shown in Figure A- 7, the viscosity ratio in the shear rate range of the cast film process used (which can be estimated from the flow rate and die geometry to around 70/sec) will be around 2. In this study, this ratio coupled with favorable interaction of the components will most probably yield to the formation of a laminar morphology for all the compositions studied. More details will be given in the morphology section below.

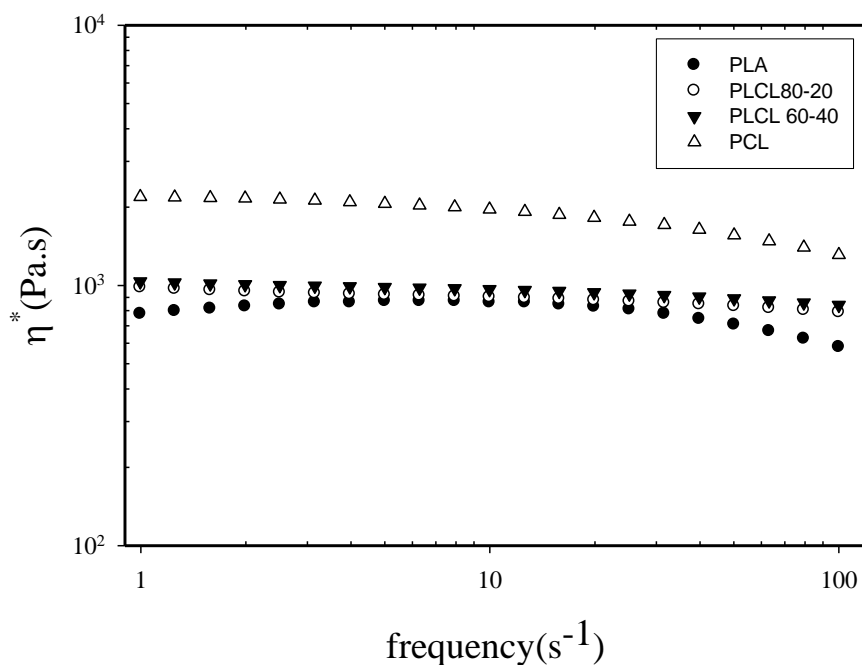


Figure A- 7 Frequency sweep of PLA, PCL and blends of 20% and 40% PCL weight content 190°C

5.2.Blend Morphology

As mentioned above, the presence of similar bonds in PLA and PCL would allow a good interaction between the dispersed phase and the matrix, which would fulfill the first criteria of the formation of elongated dispersed phase. (Kamal et al. 1995) In term of the viscosity of blends, the second criteria, as shown in Figure A- 7, both PLA and PCL have the same order of magnitude of viscosity at the processing temperature of 190°C, which will facilitate stress transfer from matrix to the disperse phase and increase the chances of laminar morphology development. Figure A- 8 shows the developed morphology in the blends cast film containing 20% and 40% of dispersed phase. The laminar morphology can be clearly observed.

Development of laminar morphology will increase the aspect ratio of the dispersed phased in the matrix. This increased area on the surface of the films will provide higher chance of PCL particles participating in the sealing and causing more interdiffusion at lower temperature. A significant change in the seal initiation temperature can thus be expected. This prediction is in agreement with the results of hot-tack and seal strength that will be discussed below.

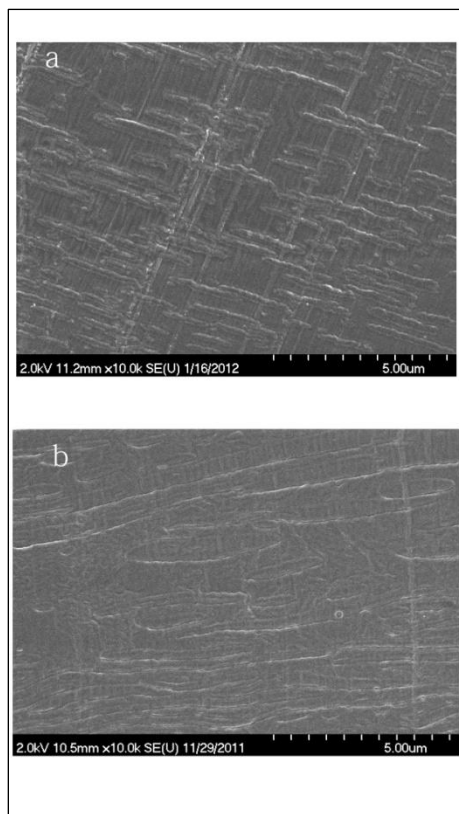


Figure A- 8 SEM images of film cross section after cryomicrotoming with diamond knife, (a) PLCL80-20, (b) PLCL60-40

5.3. Thermal analysis results

The DSC results are shown in Figure A- 9. The endothermic peak at around 60°C is associated with the melting point of PCL resin. The normalized enthalpy of melting of the blends obtained from the DSC curves show that, by increasing the PCL content over 60%, the crystallinity of the samples increase (results not shown), which indicates again some compatibility between PLA and PCL at higher concentration of PCL. Laredo et.al also showed some partial compatibility between PLA and PCL. (Newman et al. 2009) The experimental results we obtained are in agreement with their reported result on the miscibility of these two polymers.

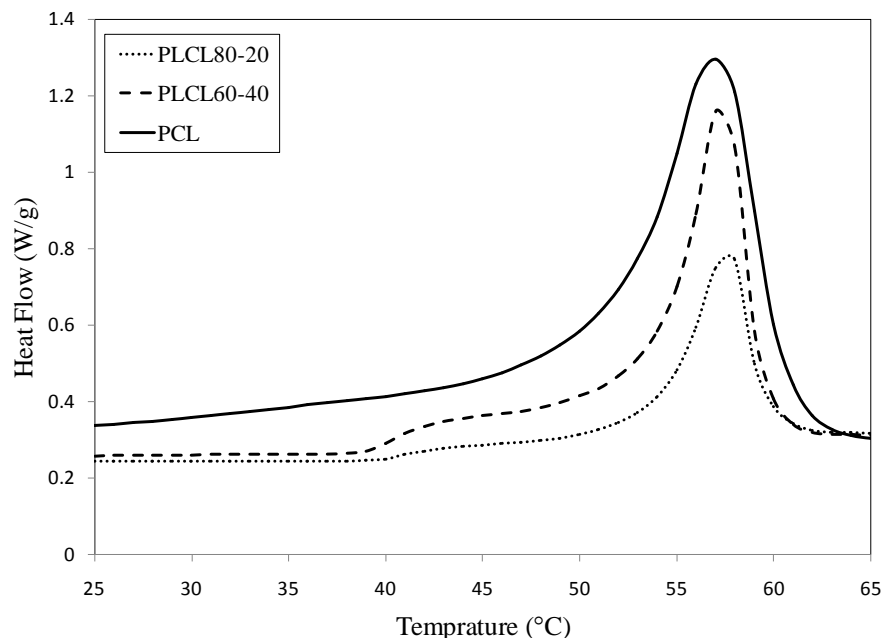


Figure A- 9 DSC thermographs of pure PCL and prepared blends

5.4.Sealability results

As mentioned above, the seal behaviour of film results from a combination of factors: seal initiation temperature, plateau seal strength, and hot-tack maximum peak strength. In order to be able to address the seal behaviour of any sealant film, all these characteristics need to be determined. (Zhang et al. 2009; Mittal and Pizzi 2009; Petrie 2000) The hot-tack results for PLA/PCL blends are presented in Figure A- 10. From the curves, it can be observed that the maximum peak of hot-tack strength for pure PLA is around 700 (g/24.5mm) and by incorporation of PCL, it increases gradually up to 1200 (g/24.5mm). The higher the hot-tack strength of the blend the higher the load we can put in the package before it fails. Expected hot-tack strength of the seal process determines the maximum hot-tack strength needed from the seal layer. Having a seal layer with significantly higher hot-tack strength will allow its use for packaging of heavier products.

Hot-tack initiation temperature shows a sharp decrease of 15°C by addition of 20% PCL and an additional 10°C for 40% PCL as illustrated in Figure A- 10. The lower melting point of PCL and, at the same time, its linear molecular chain structure are some of the reasons that cause higher chain mobility and higher rate of diffusion in the blends. Figure A- 10 and Figure A- 11 show the

decrease in hot-tack and seal initiation temperature as well as the increase in hot-tack strength by the use of PCL in the blend. The above mentioned higher mobility, in combination with the high aspect ratio of the dispersed PCL droplets in the blends, also allowed more flexible films as well as lower initiation temperature and higher hot-tack plateau. Higher flexibility of the films provides safer sealing, which means a lower chance for crack initiation and propagation in the PLA seal layer. This will be a significant advantage and, as previously explained, help maintain the performance of sealed package for prolonged period of time with fewer defects after manipulation and transportation.

This strong seal in PLA and PCL resins can be attributed to the combination of the low melting point and higher mobility of polymeric long chains. These, together with the similar viscosity of the two resins, resulted in higher aspect ratio of the dispersed phase and hence the higher diffusion of long chain molecules in the interface of the films. In addition, the polar chemical interaction coming from the nature of polyesters' heteroatomic bonds will give rise to very strong intermolecular interactions between polymer chains. This polar bond helps the seal to be stronger in the melt and solid phases. As shown in Figure A- 10 and Figure A- 11, higher hot tack and seal strength at lower temperatures is advantageous because in the processing line the temperature of sealing jaws can be adjusted to lower temperatures.

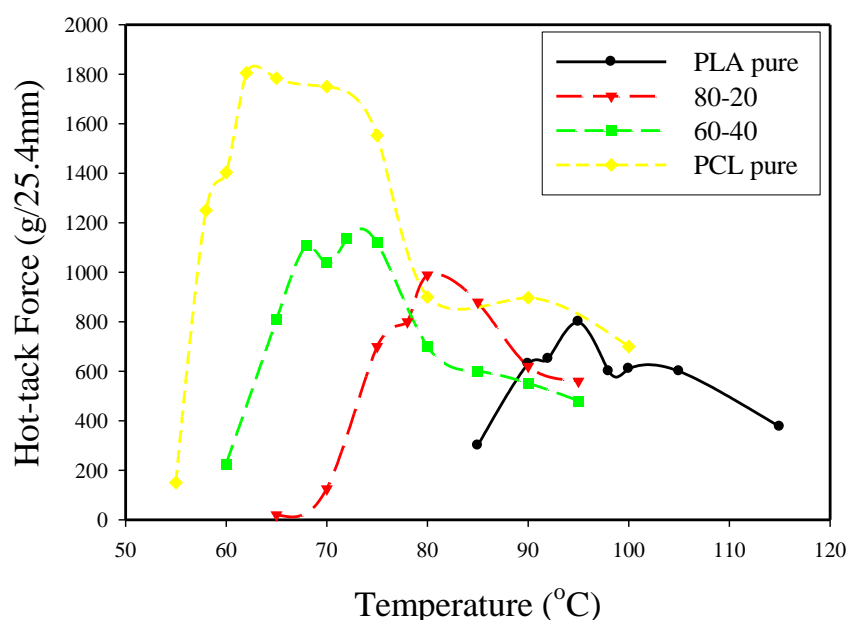


Figure A- 10 Hot-tack temperature dependence of PLA/PCL blend films

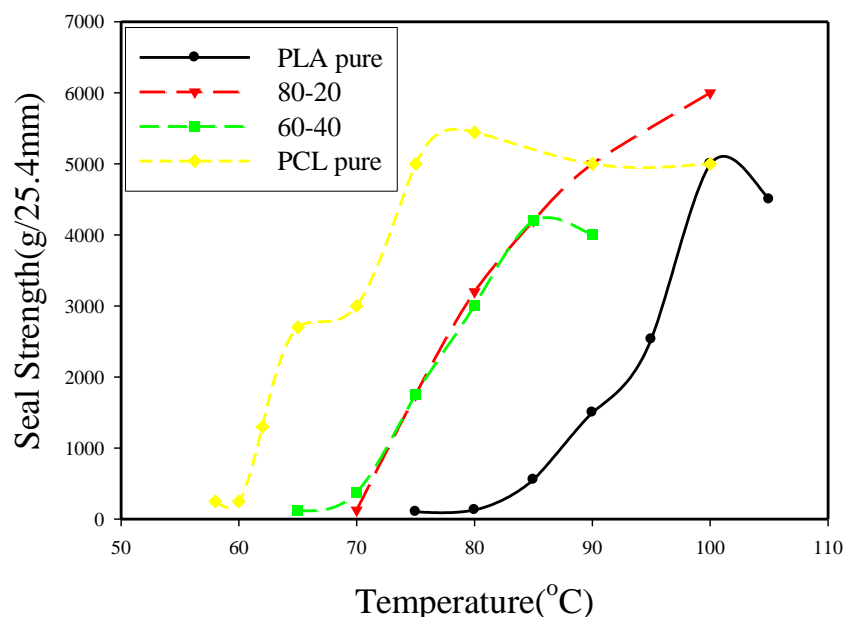


Figure A- 11 Seal strength temperature dependence of PE/EVA blend films

6. Conclusion

The results obtained from seal experiments on PE/EVA blend blown films clearly showed that their sealing performance can approach and even surpass the performance of pure EVA film. Reduction in crystallinity and melting temperature observed through DSC results as well as presence of vinyl acetate were the main reasons.

The failure mode analysis revealed that, at low temperatures, the seal fails by peeling. For intermediate temperatures, the failure occurs by delamination/elongation and at high temperatures by tearing. The presence of EVA reach phase at the surface of the films, observed in SEM micrographs, is also considered as an additional factor, which enhances the performance of EVA incorporation in polyethylene.

For PLA/PCL system, additions of PCL as a seal modifier polyester through melt extrusion to PLA significantly improved seal properties. This behaviour was related to the laminar morphology and low melting point of the dispersed phase, which cause higher chain mobility and higher interdiffusion. The polarity of the polyesters' heterogeneous bonds can also affect the seal properties of the blends. A laminar morphology was observed in the films and led to a significantly higher aspect ratio of the dispersed PCL particles, both in bulk and at the surface of

the film. This so called laminar morphology together with the higher mobility caused by incorporation of low melting point PCL resin, as shown in hot-tack and seal experiment results, have significantly improved the strength and initial hot-tack temperature as well as initial seal temperature of the blend samples. Incorporation of PCL in PLA has also led to toughening of PLA. This higher flexibility as discussed will provide safer and more reliable sealing process with fewer expected defects within and after sealing process and through transportation.

7. References:

Brogly, M., M. Nardin, and J. Schultz. 1997. "Effect of Vinylacetate Content on Crystallinity and Second-order Transitions in Ethylene—vinylacetate Copolymers." *Journal of Applied Polymer Science* 64 (10) (June 6): 1903–1912.

Chattopadhyay, S., T. K. Chaki, and Anil K. Bhowmick. 2001. "Structural Characterization of Electron-Beam Crosslinked Thermoplastic Elastomeric Films from Blends of Polyethylene and Ethylene-Vinyl Acetate Copolymers." *Journal of Applied Polymer Science* 81 (8): 1936–1950. doi:10.1002/app.1626.

Chen, Xin, and Joseph A. Jr. Gardella. 1994. "Surface Modification of Polymers by Blending Siloxane Block Copolymers." *Macromolecules* 27 (12) (June 1): 3363–3369. doi:10.1021/ma00090a034.

Halle, Richard W. 2003. "Plastomer-mVLDPE Blends for High Performance Heat Sealing Applications." In *TAPPI 2003 Conference*, 7–45. Orlando, Florida.

Ichikawa, Yasushi, and Tatsuya Mizukoshi. 2011. "Bionolle (Polybutylenesuccinate)." In *Synthetic Biodegradable Polymers*, edited by Bernhard Rieger, Andreas Künkel, Geoffrey W. Coates, Robert Reichardt, Eckhard Dinjus, and Thomas A. Zevaco, Chapter 8:285–313. Berlin, Heidelberg: Springer Berlin Heidelberg.

Jalbert, Claire, Jeffrey T. Koberstein, Iskender Yilgor, Paula Gallagher, and Val Krukonis. 1993. "Molecular Weight Dependence and End-Group Effects on the Surface Tension of Poly(dimethylsiloxane)." *Macromolecules* 26 (12) (June 1): 3069–3074. doi:10.1021/ma00064a012.

- Jin, J. 2010. "Non-Isothermal Crystallization Kinetics of Partially Miscible Ethylene-Vinyl Acetate Copolymer/low Density Polyethylene Blends." *eXPRESS Polymer Letters* 4 (3) (February 15): 141–152. doi:10.3144/expresspolymlett.2010.19.
- Kamal, M. R., H. Garmabi, S. Hozhabr, and L. Arghyris. 1995. "The Development of Laminar Morphology during Extrusion of Polymer Blends." *Polymer Engineering & Science* 35 (1): 41–51.
- Lee, H., and L. Archer. 2001. "Surface Modification of Polymer via Surface-Induced Migration of Copolymer Additives." In *ANTEC 2001 Conference Proceedings*, 1472–1476.
- Lee, Sang Young, and Sung Chul Kim. 1997. "Laminar Morphology Development and Oxygen Permeability of LDPE/EVOH Blends." *Polymer Engineering & Science* 37 (2): 463–475. doi:10.1002/pen.11690.
- McNally, G. M. 2005a. "The Effect of Polymer Properties on the Mechanical Behavior and Morphological Characteristics of Cast Polyethylene Film for Stretch and Cling Film Applications." *Journal of Plastic Film and Sheeting* 21 (1) (January 1): 39–54. doi:10.1177/8756087905052804.
- McNally, G.M. 2005b. "The Effect of Vinyl Acetate Content and Polyisobutylene Concentration on the Properties of Metallocene Polyethylene-Ethyl Vinyl Acetate Coextruded Film for Stretch and Cling Film Applications." *Journal of Plastic Film and Sheeting* 21 (1) (January 1): 69–83. doi:10.1177/8756087905053791.
- Meka, P., and F. C Stehling. 1994. "Heat Sealing of Semicrystalline Polymer Films. I. Calculation and Measurement of Interfacial Temperatures: Effect of Process Variables on Seal Properties." *Journal of Applied Polymer Science* 51 (1): 89–103.
- Mittal, K.L., and A. Pizzi. 2009. *Handbook of Sealant Technology*. 1st ed. Boca Raton: CRC Press.
- Moly, K. A., S. S. Bhagawan, G. Groeninckx, and S. Thomas. 2006. "Correlation between the Morphology and Dynamic Mechanical Properties of Ethylene Vinyl Acetate/linear Low-Density Polyethylene Blends: Effects of the Blend Ratio and Compatibilization." *Journal of Applied Polymer Science* 100 (6): 4526–4538.

- Mueller, C., G. Capaccio, A. Hiltner, and E. Baer. 1998. "Heat Sealing of LLDPE: Relationships to Melting and Interdiffusion." *Journal of Applied Polymer Science* 70 (10): 2021–2030.
- Newman, Dinorah, Estrella Laredo, Alfredo Bello, Angélica Grillo, José Luis Feijoo, and Alejandro J. Müller. 2009. "Molecular Mobilities in Biodegradable Poly(dl-lactide)/Poly(ϵ -Caprolactone) Blends." *Macromolecules* 42 (14) (July 28): 5219–5225. doi:10.1021/ma9007303.
- Peón, J, J.F Vega, B Del Amo, and J Martínez-Salazar. 2003. "Phase Morphology and Melt Viscoelastic Properties in Blends of Ethylene/vinyl Acetate Copolymer and Metallocene-Catalysed Linear Polyethylene." *Polymer* 44 (10) (May): 2911–2918.
- Petrie, Edward M. 2000. *Handbook of Adhesives and Sealants*. London: McGraw-Hill Professional.
- Poisson, C., V. Hervais, M. F Lacrampe, and P. Krawczak. 2006a. "Optimization of PE/binder/PA Extrusion Blow Molded Films. II. Adhesion Properties Improvement Using binder/EVA Blends." *Journal of Applied Polymer Science* 101 (1) (July 5): 118–127.
- Poisson, V. Hervais, M. F Lacrampe, and P. Krawczak. 2006b. "Optimization of PE/Binder/PA Extrusion Blow Molded Films. I. Heat Sealing Ability Improvement Using PE/EVA Blends." *Journal of Applied Polymer Science* 99 (3) (February 5): 974–985.
- Sarazin, Pierre, Xavier Roy, and Basil D. Favis. 2004. "Controlled Preparation and Properties of Porous Poly(l-Lactide) Obtained from a Co-Continuous Blend of Two Biodegradable Polymers." *Biomaterials* 25 (28) (December): 5965–5978.
- Stehling, F. C, and P. Meka. 1994. "Heat Sealing of Semicrystalline Polymer Films. II. Effect of Melting Distribution on Heat-Sealing Behavior of Polyolefins." *Journal of Applied Polymer Science* 51 (1): 105–119.
- Zhang, Jinguo, Evangelos Manias, George Polizos, Jin-Young Huh, Amos Ophir, Ponusa Songtipya, and Maria Jimenez-Gasco. 2009. "Tailored Polyethylene Nanocomposite Sealants: Broad-Range Peelable Heat-Seals Through Designed Filler/Polymer Interfaces." *Journal of Adhesion Science and Technology* 23 (5): 709–737.