

# STEADY-STATE MODELING OF EXTRUSION CAST FILM PROCESS, NECK-IN PHENOMENON AND RELATED EXPERIMENTAL RESEARCH: A REVIEW

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## ABSTRACT

This review paper provides the current state of knowledge of steady-state modeling of the extrusion cast film process used to produce flat polymer films, as well as related experimental research with a particular focus on the flow instability neck-in. All kinematic models used (i.e. 1, 1.5, 2 and 3-dimensional models) together with the utilized constitutive equations, boundary conditions, simplified assumptions and numerical methods are carefully summarized. The effect of draw ratio, Deborah number (i.e. melt relaxation time related to experimental time), film cooling, second to first normal stress difference ratio at the die exit, uniaxial extensional strain hardening and planar-to-uniaxial extensional viscosity ratio on the neck-in is discussed.

## 1 INTRODUCTION

Extrusion film casting is an industrially important process, which in practice has a solid place among polymer processing technologies. It can be classified as a continuous, high-speed manufacturing process, during which monolayer or co-extruded multilayer thin, highly oriented films are produced. A wide range of the plastic films and sheets produced by this technology are used in many different applications of daily and technical use: plastic bags, consumer packaging, magnetic tapes for storing audio video content, optical membranes for liquid crystal displays, flexible electronics, foils for capacitors and microporous membranes used primarily in separation processes (from microfiltration to reverse osmosis or as separators in lithium-ion batteries for mobile devices and electric vehicles [1–4]) or as a product for further processing by other technologies such as thermoforming and biaxial orientation [5, 6].

The growing demand for the quantity production and quality of manufactured films, together with the introduction of new materials, requires new approaches in production line. Of particular interest is to reach desirable properties of the produced films and to keep film thickness uniform and width as close as possible to the designed extrusion die width. In order to eliminate an expensive and time-consuming trial-and-error approach widely used in the plastics industry to optimize film casting process, one can use a computer modeling for the optimization of die design and process conditions for a given polymer system. This strategy can provide a better insight into the problem, broaden the knowledge on relationships between process/rheological variables and propose possible approaches to deal with them to optimize the process or provide a better understanding of basic underlying mechanics [6].

## 1.1 Film casting process description

The extrusion film casting is a technology, in which polymer pellets are conveyed, homogenized, compressed and melted in an extruder. Then, the polymer melt is pushed through the uniform slit die (center-fed T die or coat-hanger die) with typically about 1–2 mm gap size [5]. The thick sheet is then intensively stretched in the machine direction using a constant rotary whose circumferential velocity,  $v_x(X)$ , is higher than the average polymer melt velocity at the die exit,  $v_x(0)$ . This leads to the orientation of macromolecules and reduced film thickness, and due to a sufficiently high cooling rate, the final film dimensions are fixed. Intensity of the stretching is given by a draw ratio, which is defined as  $DR = v_x(X)/v_x(0)$ . Additionally, an increase in  $DR$ , cooling rate or stretching distance can cause temperature and/or stress induced crystallization, which can enhance final film properties. The process is visualized in Figure 1.

At the chill roll, several other technological devices can be used to provide a better contact line between the film and the chill roll and to increase the heat transfer rate, such as an air knife (a slit nozzle blows a jet of cooled air to film) or electrostatic pinning [5–7]. In the latter device, a high voltage wire is positioned parallel to the grounded chill roll that generates an electrostatic discharge exerting electrostatic force on the film to increase the film-chill roll contact. Another alternative with the similar result is a vacuum box, which provides a vacuum between the film and the chill roll [7–9]. In addition to cooling on the chill roll, the polymer film is naturally cooled to some extent, depending on the length of the drawing zone, by passing through the surrounding environment. This can be enhanced by introducing convection air or an inert gas source into this section or by passing the film through a fluid bath [10]. Additionally, the produced polymeric film can also be subjected to treatment (plasma treating, heating and biaxial orientation) depending on the desired properties and purpose of the final product. Polymer behavior and extensional conditions in the drawing zone have been shown to be key factors determining the final mechanical and optical properties of the film [6, 11].

To produce highly functional films with tailored properties, multiple layers of different polymer melts can be coextruded and stretched, i.e., the properties of the film are given by each individual layer. In this way, multilayer films with enhanced properties, such as oxygen and moisture impermeability, strength, chemical resistance or color can be produced [12]. An alternative continuous film production technology is called the extrusion film blowing process. In this process, the extruded tube is inflated by the internal pressure into a bubble shape having a thin wall thickness, which is simultaneously quenched and hauled off [13–17]. In contrast to this competing film production technology, films made by extrusion film casting have good transparency, uniformity of thickness, a smoother surface and are produced at a higher production rate [6].

According to the current industry practice, where a wide variety of films are produced with a requirement for use in heterogeneous applications, manufactures process a broad range of materials by using film casting technology. Frequently used polymeric materials include low-density polyethylene, LDPE; high-density polyethylene, HDPE; linear low-density polyethylene, LLDPE; polypropylene, PP; polyethylene terephthalate, PET; and polystyrene, PS. The extrusion film casting is also suitable for low viscosity polymers [18] and biodegradable polymers such as for example polylactide (PLA) or its blends with polybutylene succinate (PBS) or poly(3-hydroxybutyrate-co-3-hydroxyvalerate) (PBAT) [19, 20]. Since these films have a wide range of applications, there is a requirement to produce a wide range of sizes. The film width can typically range from 0.1 m to 10 m, thicknesses from 20  $\mu\text{m}$  to 2000  $\mu\text{m}$  [11] at production rates ranging from 70 to 200 m/min. Tolerable thickness variation is reported to be from 3 to 5 % [5]. The plastics industry, which focuses on the production of plastic foils, is currently undergoing a major change, due to the gradual transition from conventional commodity polymers to more advanced [6]. These include, for example, metallocene polymers with an easily modifiable structure which make it possible to

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significantly improve the final properties of the film. Structural polymers such as polyethylene terephthalate, polycarbonate, polyamide, polyphenylsulfide have become popular materials for producing films with high heat resistance. The line speed for the production of polymer films is gradually increasing for economic reasons and in some cases (e.g. polypropylene or polyethylene terephthalate) may reach up to 500 m/min [6].

## 2 FLOW INSTABILITIES

The presence of an air-polymer interface in the drawing zone makes it possible to develop various types of flow instabilities that severely limit the desired film quality and quantity. Their formation is influenced by processing conditions, heat transfer and rheology of the processed polymer. For example, if the draw ratio reaches some critical value (for the given process conditions, die design and polymer used), transient hydrodynamic instability, called draw resonance, begins to occur [21]. This instability causes oscillations of the film dimensions, although the volumetric flow supplied from the slit die and take-up speed are kept constant, see Figure 2. These periodic fluctuations in film width and thickness (measured in the center of the film) are offset by half-wavelength (i.e. maximum in width corresponds to the minimum thickness) and vice versa [22]. Extension of the drawing distance, increased cooling effects and the use of polymers with strong extensional strain hardening can stabilize the process and move the onset of draw resonance toward higher draw ratios [23].

Film breakage is another feature that can be observed during increasing the draw ratio. In this case, the chains cannot be reorganized to relieve local stresses within the time frame imposed by the deformation, resulting in a cohesive failure between the polymer chains and disintegration of the film. This can be seen in polymers containing long chain branches or a high molecular weight portion processed at high line speeds and cooling rates, leading to good process stability but also to the development of high tensile stress [6].

Neck-in and edge-beading are flow phenomena which are the most common instabilities in the production of flat films because they occur and destabilize the flow at any processing conditions. These instabilities are described and reviewed below in greater detail.

## 2.1 Neck-in

Upon leaving the die, the extruded polymer in the form of a thick sheet exhibits swelling due to its viscoelastic nature. This relaxation of molecular stress is then influenced by the velocity rearrangement that occurs during the transition from a confined shear flow in the slit die to the downstream extension. When the polymer sheet is hauled off further downstream and stable processing conditions are met, its cross-sectional dimensions are monotonically reduced due to the external drawing force exerted on the sheet by the rotary winding drum. In addition to the desirable reduction in film thickness, the width of the film is reduced. This defect is called the neck-in and can be defined as the difference between the half-width of the film at the die exit and the final half-width of the solidified film (Figure 3). The neck-in is considered to be a typical instability occurring in extensional flows as explained by Larson in [24], even if it occurs under steady state extrusion conditions (i.e. that the stress and the velocity are not time dependent at the given point of the stretched film) because it can have serious consequences since it might lead to breakage of the film.

The role of extrudate swell on the film drawing was investigated by using viscoelastic constitutive equations such as the Leonov model [25] and the linear PTT model [26]. It was demonstrated that the negative value of the second normal stress difference causes swelling in the thickness direction much higher than in the width direction of the extrudate [26]. Even if the intensity of the extrudate swell rapidly decreases by increased take-up velocity, there might be “a certain amount of swelling persisting near the die exit” lowering the melt velocity at this region [26]. This can increase the melt orientation because the actual  $DR$  “expressed in terms of the velocity at the point of the film’s maximum thickness” is higher than the conventional  $DR$  based on the melt velocity at the die exit [26].

Based on experimental studies (Table 1) and theoretical analyses (Tables 2–5), the following material parameters and process variables have been identified to have a significant



impact on the neck-in phenomenon: molecular weight ( $M_w$ ), molecular weight distribution ( $MWD$ ), relaxation time ( $\lambda$ ), the ratio of the second and first normal stress difference at the die exit ( $-N_2/N_1$ ), long chain branching ( $LCB$ ), strain hardening in uniaxial extension ( $SH$  in  $\eta_{E,U}$ ), planar to uniaxial extensional viscosity ratio ( $\eta_{E,P}/\eta_{E,U}$ ), draw ratio,  $DR$ , take-up length,  $X$ , take-up rate,  $v_x(X)$ , melt speed at the die exit,  $v_x(0)$ , and temperature  $T$ . The role of each individual parameter on this phenomenon is summarized in Table 6. In order to clarify the reading of Table 6, let us provide here an example explaining its first line, which should be read as follows: In 1986, Dobroth and Erwin reported that the neck-in for LDPE increases if the draw ratio,  $DR$ , (adjusted via the average polymer melt velocity at the die exit,  $v_x(X)$ ) increases or if the take-up length,  $X$ , increases.

As can be seen, the reduction of the neck-in can be achieved by an increasing the polymer melt relaxation time  $\lambda$  (via broadening  $MWD$  and/or increasing  $M_w$  and/or decreasing  $T$ ), increasing the melt speed at the die exit  $v_x(0)$  (maintaining a constant  $DR$ ) or reducing the air-gap (distance between die and roll),  $X$ . All these three variables determine the elasticity of the melt, which can be evaluated in terms of the Deborah number defined as

$$De = \frac{\lambda v_x(0)}{X} \quad (1)$$

It is obvious, that if the Deborah number (i.e. melt elasticity) increases, the neck-in decreases (although the stability of the process in terms of the maximum attainable draw ratio,  $DR$ , at which the film breaks may be lowered [27–29]). Thus, it is appropriate to maintain the level of elasticity reasonably high to minimize neck-in, which can be achieved by increasing the relaxation time and/or the melt speed at the die exit or by reducing the air-gap (see Eq. 1). The effect of relaxation time on the neck-in phenomenon determined experimentally for two linear low-density polyethylenes, LLDPEs [30], and two linear polypropylenes, PPs [31–34], is provided in Figures 4–5. It is important to mention that a different definition of relaxation time

can be found in the reviewed literature. In the studies based on single-mode constitutive equations, the utilized Maxwell relaxation time [27], the shortest [35] or characteristic relaxation time (determined by the reciprocal frequency at the intersection of the storage modulus  $G'$  and the loss modulus  $G''$  curves [36] or by fitting the strain rate dependent steady uniaxial extensional viscosity data [37, 38]) are typically used to calculate  $De$ . In the case of multi-mode constitutive equations, the relaxation time for each mode [39–44] or an average relaxation time,  $\bar{\lambda}$ , is calculated to determine  $De$  by using the following expression [45–47]:

$$\bar{\lambda} = \frac{\sum_{j=1}^N G_j \lambda_j^2}{\sum_{j=1}^N G_j \lambda_j} \quad (2)$$

where  $\lambda_j$  and  $G_j$  is the relaxation time and the modulus, respectively, in the  $j^{\text{th}}$  relaxation mode. Also, in some experimental studies, the longest relaxation time ( $\lambda = \eta_0 J_E^0$ , where  $\eta_0$  is the zero-shear viscosity and  $J_E^0$  is the linear steady-state elastic compliance) [33] or the characteristic (reptation-mode) relaxation time representing the onset of shear-thinning [30], are used.

The role of  $DR$  on the neck-in is complex, depending whether the polymeric chains are linear or branched or if  $DR$  is changed via  $v_x(X)$  or  $v_x(0)$ . Current experimental studies showed that for linear polymers (i.e. for PP, PET, LLDPE, HDPE), an increase in  $DR$  (by an increase in  $v_x(X)$ ) always increases the neck-in but for branched polymers (such as LDPEs), interestingly, the trend can even become an opposite (see Table 6 and Figure 6). This unexpected trend was attributed to the strain-hardening behavior of LDPE in elongational flow [48]. The situation becomes also complex, if  $DR$  is increased by reduction in  $v_x(0)$  (keeping the  $v_x(X)$  constant). It was reported for branched LDPE and linear PET that an increase in  $DR$  (by decrease in  $v_x(0)$ ) reduces the neck-in, but for linear isotactic polypropylene (iPP) the trend was found to be

surprisingly opposite for given processing conditions. This unexpected trend for iPP was attributed to the increased crystallization rate, which caused quicker film solidification.

It was found that the introduction of strain hardening,  $SH$ , in uniaxial extensional viscosity,  $\eta_{E,U}$ , by incorporating long chain branches into polymer backbone chains, reduces the neck-in phenomenon. This trend is illustrated in Figure 7 for linear low-density polyethylene and highly branched low-density polyethylene. Seay and Baird [30] revealed that the addition of sparse long chain branching ( $LCB$ ) to polymer chains, i.e.  $SH$  in  $\eta_{E,U}$ , is more significant for film width conservation than broadening the molecular weight distribution ( $MWD$ ). In addition, they have found that increasing  $LCB$  of long and short chains reduced the neck-in at low and high draw ratios, respectively. This effect is sometimes used to improve final width for films made from polymers prone to the neck-in (such as HDPE, LLDPE) using coextrusion technology in which the surface/edge portion of the film is made of a material having a long-chain branching (such as LDPE) and a core from linear polymer [49–51].

Recent viscoelastic modeling of the extrusion film casting process, which followed the corresponding neck-in measurements, suggests that reduction of  $\eta_{E,P}/\eta_{E,U}$  or  $-N_2/N_1$  at the die exit (if  $De > 0.1$ ) can also reduce the neck-in phenomenon (see Table 6).

In order to understand the role of extensional rheology and the die exit stress state, it is necessary to discuss the mechanism and physical background of the neck-in in more detail. Ito et al. [52], performed an experimental study on metallocene-catalyzed linear low-density polyethylene, mLLDPE, aimed to visualize the flow in the air-gap region during a film casting operation. They used small aluminum particles and made streamline measurement by using the particle tracking method. These particles were placed across the film width at the die exit (one particle for each measurement) and their movement was monitored by a camera for different draw ratios ( $DR = 4.4, 7.5$  and  $12.2$ ). It has been found that streamlines at the film center are straight regardless the drawing intensity (i.e. there is planar extensional flow) whereas the

streamlines at the near-edge region were found to be curved with a tapered transversal spacing in the flow direction confirming presence of uniaxial extensional flow (see Figure 8). Moreover, an increase in the draw ratio caused an increase in the streamlines tapering, which lead to a more pronounced neck-in phenomenon. Therefore, if  $\eta_{E,U}$  increases due to  $SH$  in such a way that  $\eta_{E,P}/\eta_{E,U}$  is decreased (i.e. if the resistance against the uniaxial extensional flow becomes much higher than the resistance against the planar extensional flow), the polymer melt starts to prefer the planar extensional flow at the expense of the uniaxial elongation flow and thus, the neck-in is decreased. Similarly, the reduction in  $-N_2/N_1$  at the die exit physically means an induction of a planar prestretch inside the extrusion die (for example by using converging instead of a parallel flow channel), which (if remembered by the melts, i.e. if  $De > 0.1$ ) increases dominance of the planar extensional flow in the post die area and therefore, the neck-in is reduced.

In the industrial practice, it is useful to have a tool that can provide a reasonable evaluation of the neck-in for a particular polymeric material and processing conditions prior to film production itself, where its determination via a trial-and-error approach can be very expensive. It is therefore not surprising that considerable efforts have been made to relate the neck-in with the air-gap, extensional strain rate and relaxation time [53] (Eqs. 3–4), both planar and uniaxial extensional viscosities [54] (Eq. 5), and to the strain hardening, the ratio of planar to uniaxial extensional viscosity, the Deborah number and the die exit stress state [38] (Eq. 6). These simple analytical models are easy to use and have the advantage to gain a correlation between the particular model variables with the naked eye in order to identify key process and material parameters to optimize them for neck-in reduction. In more detail, Ito et al. [53] in 2003 developed a model based on the Dobroth-Erwin model [55], which assumes a planar extensional flow in the middle of the film and a uniaxial extensional flow at the edge. According to their

theory, the final film width is determined by the ratio of planar viscosities in the axial and transverse directions with respect to the flow. Proposed relationship for the neck-in,  $NI$ , considering that the polymer melt behaves as a Newtonian fluid, is following

$$NI = \sqrt{\frac{1}{2}}X \quad (3)$$

where the air-gap,  $X$ , is the only variable (i.e. draw ratio, deformation rate, relaxation time or viscosity are not included). The model was tested using experimental data for linear HDPE, short chain branched LLDPE and long chain branched LDPE melts expressed as the neck-in plotted against the air-gap at four different draw ratios. The model was shown to correctly predict the general trend between  $NI$  and  $X$ , i.e. an increase in  $X$  causes an increase in  $NI$ , and the predicted slope of the theoretical line was close to experimental data for HDPE and LLDPE at short air-gap values and the highest draw ratios at given processing conditions. On the other hand, the model tended to overpredict  $NI$  (especially for LDPE) without the ability to predict the effect of draw ratio on  $NI$ , as expected due to the absence of deformation rate and any rheological parameters in Eq. 3. If the upper convected Maxwell model is used, the expression for  $NI$  yields the following form

$$NI = \sqrt{\frac{1}{2}(1 - 2\dot{\epsilon}_p\lambda)}X \quad (4)$$

where  $\dot{\epsilon}_p$  denotes the extensional strain rate of the planar part defined as  $\dot{\epsilon}_p = d\epsilon_p(t)/dt$  ( $\epsilon_p$  is the Hencky strain of the planar deformation) and  $\lambda$  is the characteristic relaxation time.

Although the neck-in trend predictions are consistent with the observations with respect to the melt elasticity or the air-gap, the model unrealistically predicts neck-in decrease for the increased draw-down ratio, which was attributed to the used constitutive equation. Further works have moved further and tried to predict neck-in based on uniaxial and planar extensional viscosities. Shiromoto et al. [54] in 2010 developed a theoretical model based on the force

balance and film deformation in the post die area. The authors found that  $NI$  can be correlated with the planar to uniaxial extensional viscosity ratio rather than with the strain hardening in uniaxial extension or with the ratio of planar viscosities. These findings were transformed into the following expression for  $NI$

$$NI \cong X \left\langle \frac{\eta_{E,P}}{\eta_{E,U}} \right\rangle^{0.5} \quad (5)$$

where  $\eta_{E,P}$  and  $\eta_{E,U}$  denotes planar and uniaxial extensional viscosity. Eq. 5 was validated using relevant experimental data for three long chain branched LDPEs having different  $M_w$  and  $MWD$ . The ratio  $\eta_{E,P}/\eta_{E,U}$  was determined for all three samples using the multi-mode exponential type of Phan–Thien and Tanner constitutive model utilizing parameters identified on the experimental data obtained from rotational and capillary rheometers.  $\eta_{E,U}$  at low deformation rates and high extensional rates was determined with a Meissner-type rheometer (ARES-EVF, TA Instruments) and the Cogswell method [56], respectively. The key limitation of the Eq. 5 is the absence of variables allowing to evaluate the role of the Deborah number and the die exit stress state on  $NI$ . Barborik and Zatloukal [37, 38] continued the research of the neck-in phenomenon in the period 2017-2018 with respect to the ratio of the second and first normal stress difference at the die exit,  $-N_2/N_1$ , uniaxial extensional strain hardening,  $\eta_{E,U,max}/3\eta_0$ , melt elasticity (captured via the Deborah number,  $De$ ) and the ratio of planar-to-uniaxial extensional viscosity,  $\eta_{E,P}/\eta_{E,U}$ . Using an isothermal 1.5-dimensional (1.5D) membrane model and viscoelastic modified Leonov constitutive equation, the following expression for maximum attainable neck-in was proposed:

$$NI = \frac{X}{\sqrt[7.43]{\frac{\eta_{E,U,\max}}{3\eta_0}}} \left\{ \begin{aligned} &0.593 \left[ 1 - \exp(-1073.742 De^{2.113}) \right] \left( \sqrt{\frac{\eta_{E,P}}{\eta_{E,U}}} - 1 \right) + \\ &\left\{ \begin{aligned} &1 + 1.027 \arctan \left( 0.849 \frac{N_2}{N_1} \right) \\ &\arctan \left[ 0.514 \frac{\eta_{E,U,\max}}{3\eta_0} \tanh(3.953 De) \right] \end{aligned} \right\} \\ &1.027 \left[ 1 - \exp(-0.849 De^{0.514}) \right] \end{aligned} \right. \quad (6)$$

The predictions of the model were found to be in good agreement with the corresponding experimental data [38] (see Figure 9) capturing all the trends obtained numerically, i.e. that  $NI$  is reduced if

- $-N_2/N_1$  at the die exit decreases (i.e. for increased planar pre-stretch of the melt inside the extrusion die),
- $De$  increases,
- $\eta_{E,P}/\eta_{E,U}$  decreases,
- $\eta_{E,U,\max}/3\eta_0$  increases.

It has also been revealed that there is a threshold of about 0.1 for  $De$  above which the neck-in phenomenon starts to be strongly dependent on the  $-N_2/N_1$  ratio at the die exit. In other words, if  $De > 0.1$ , the flow history inside the die (i.e. the die design) starts to significantly affect the neck-in phenomenon. It is important to mention that Eq. 6 represents an analytical approximation of numerical solutions based on an isothermal (1.5D) membrane model utilizing the modified Leonov constitutive equation (single mode) for the processing conditions, in which the maximum attainable neck-in is achieved (i.e. for very high draw ratios only) were

$$0.011 \leq De \leq 0.253, \quad 0.825 \leq \frac{\eta_{E,P}}{\eta_{E,U}} \leq 1.910, \quad 2.047 \leq \frac{\eta_{E,U,\max}}{3\eta_0} \leq 10.096 \quad \text{and}$$

$0.017 \leq -\frac{N_2}{N_1} \leq 0.680$ . The basic form of Eq. 6 has been derived from the assumed linear

function between  $Nl$  and  $\sqrt{\eta_{E,P} / \eta_{E,U}}$ , in which its constants were allowed to vary with  $De$  according to the Avrami exponential functions. Detailed derivation of Eq. 6 is provided in [37, 38]. Validation of Eq. 6 was performed for different highly branched LDPEs. All rheological quantities appearing in Eq. 6 were predicted by the single-mode modified Leonov model whose parameters were identified from uniaxial extensional viscosity data only. This procedure seems to be reasonable at least for the given LDPEs and applied processing conditions. In order to experimentally evaluate  $\eta_{E,P}/\eta_{E,U}$ , one could use the Cogswell model and measured entrance pressure drops on capillary rheometer by using circular and rectangle capillaries [57, 58] whereas  $-N_2/N_1$  can be evaluated using Han's methods utilizing exit pressure drop measures by using a capillary rheometer equipped by a slit die [59–64]. The key advantage of Eq. 6 in contrary to Eqs. 3–5 is a consideration of uniaxial and planar extensional viscosities along with the Deborah number and die exit stress state (quantified via  $-N_2/N_1$ ). On the other hand, the model is only applicable to very high draw ratios and does not take into account the full relaxation spectrum, film cooling and crystallization, which can be considered as its key limitations.

## 2.2 Edge-beading

In addition to the neck-in phenomenon, an interrelated defect, referred to as the edge-beading or the dog-bone defect, is formed making the edge portions of the film substantially thicker than its central part (Figure 10). The size of these raised parts can be five times higher compared to the center and several centimeters wide. The predominant cause of the edge-beads formation is the edge-stress effect [55] arising due to the neck-in phenomenon and its intensity increases with the draw ratio (see Figure 11). The following equation was derived in [55] to evaluate the edge-beading:



$$\frac{h_f^{\text{edge}}}{h_f^{\text{center}}} = B = \sqrt{DR} \quad (7)$$

where  $B$  is the bead ratio and  $h_f^{\text{edge}}$  and  $h_f^{\text{center}}$  represents edge and center final thickness, respectively. This equation was derived by simply comparing the strains of the center (undergoing planar stretch) and edge (undergoing uniaxial stretch) elements between the roll and the die neglecting surface tension, extrudate swell and assuming melt incompressibility (i.e. without the need to use any constitutive equation). Equation 7 was successfully validated for LDPE for  $DR$  between 1 and 20 [55] (see Figure 12). It has also been shown (when comparing simulations based on the Newtonian and UCM models) that increasing the melt elasticity (by increasing the Deborah number) decreases the intensity of the edge-beading [28].

The raised edges are often trimmed with a slit razer, scrapped and optionally reprocessed to achieve a uniform film surface. Regardless of the large amount of waste material, the occurrence of edge-beads also causes air entrapment between the film and the chill roll, which in turn results in poorer film quality. In the manufacturing practice, a technological procedure of opening lateral parts of the extrusion slit die (i.e. the gap size at the edge is bigger than in the center) can be found in order to create thicker edges that would restrain the neck-in level in comparison to the situations when the edge-beads and the neck-in would evolved in the natural way [11].

## 3 MATHEMATICAL MODELING OF THE EXTRUSION FILM CASTING

## PROCESS

The drawing of polymer films or filaments has taken an enormous amount of attention and has been extensively studied both experimentally and theoretically in the past four decades because of its great importance in the polymer processing industry.

## 3.1 Flow kinematics

Individual research groups focused on experimental works [30, 48, 52, 55, 65–74] dealing with flow visualization, effects of temperature, crystallization, molecular weight distribution or long chain branching on kinetics of the film casting process (see Table 1). Theoretical research has not disappeared, and attention has been drawn to the development and use of numerical models (primarily considering steady-state conditions) of different dimensionality such as 1D [22, 25, 26, 45, 75–85], 1.5D [18, 27, 47, 53, 86–93, 29, 94–100, 35, 37, 38, 41–44], 2D [8, 12, 28, 36, 39, 40, 46, 54, 101–113] and full 3D [114–116] (see Tables 2–5) using different types of constitutive equations taking into account non-isothermal effects, crystallization, inertia, and gravity. The 1D model here is based on the assumption of an infinite film width and assumes the following velocity field

$$v_x = v_x(x) \quad (8)$$

$$v_y = 0 \quad (9)$$

$$v_z = -z \frac{\partial v_x}{\partial x} \quad (10)$$

that is, the flow deformation in the drawing region is planar [52, 55]. The 1.5D model is simply the 1D model with variable film width proposed in [27, 28]. This simplified model, which

retains the ability to cover the reduction in film width in the drawing length while reducing the dimensionality of the solved problem, is based on the assumption that all velocity components are an exclusive function of the drawing length position,  $x$ , and vary linearly with respect to its corresponding direction. In this case, a velocity field is assumed in form of

$$v_x = v_x(x) \quad (11)$$

$$v_y = yf(x) \quad (12)$$

$$v_z = -zg(x) \quad (13)$$

The 2D approximation developed in [112, 113] is based on the so-called membrane hypothesis considering that one dimension of the film is small compared to the others [112]. The film thickness is much smaller (several orders of magnitude) than the film width and the take-up length, so it can be assumed that the velocity component in the machine and the transversal direction are independent of thickness direction, i.e., uniform across the thickness. The reduced velocity field is given in the following form

$$v_x = v_x(x, y) \quad (14)$$

$$v_y = v_y(x, y) \quad (15)$$

$$v_z = -z \left( \frac{\partial v_x}{\partial x} + \frac{\partial v_y}{\partial y} \right) \quad (16)$$

The 3D model utilizes velocity components without any restriction, which are given bellow:

$$v_x = v_x(x, y, z) \quad (17)$$

$$v_y = v_y(x, y, z) \quad (18)$$

$$v_z = v_z(x, y, z) \quad (19)$$

### 3.2 Constitutive equations

Different types of constitutive equations were used to model film casting, as shown in Tables 2–5. They are introduced and briefly discussed in this chapter. For simplicity, they are usually provided in a single mode version. Note that in the multi-mode approach, a discontinuous relaxation spectrum is used and the stress tensor is expressed as  $\underline{\underline{\tau}} = \sum_{j=1}^N \underline{\underline{\tau}}_j$  where  $\underline{\underline{\tau}}_j$  represents the stress tensor in the  $j^{\text{th}}$  mode.

#### 3.2.1 Newtonian model

This constitutive equation describes the behavior of ideal Newtonian fluids by Eq. 20.

$$\underline{\underline{\tau}} = 2\eta_0 \underline{\underline{D}} \quad (20)$$

Here  $\underline{\underline{\tau}}$  is the extra stress tensor,  $\eta_0$  is the Newtonian shear viscosity (zero-shear-rate viscosity) and  $\underline{\underline{D}}$  is the deformation rate tensor defined as

$$\underline{\underline{D}} = \frac{1}{2} (\underline{\underline{L}}_v + \underline{\underline{L}}_v^T), \underline{\underline{L}}_v = \nabla v \quad (21)$$

where  $v$  represents the velocity field,  $T$  denotes the transpose of the tensor and  $\nabla$  is the gradient operator. The Newtonian model predicts constant steady shear viscosity ( $\eta_0$ ), steady uniaxial ( $3\eta_0$ ) and planar ( $4\eta_0$ ) extensional viscosities, which is correct also for polymer melts, but only at very low extensional rates, where they behave as Newtonian fluids. The key advantage of this model is its mathematical simplicity and utilization of only one adjustable parameter,  $\eta_0$ , which can be determined from simple shear flow measurements. On the other hand, the model does not have the ability to describe the elasticity and memory of fluids, the extensional strain thinning or the extensional strain hardening, typically occurring for polymer melts at medium and high extensional strain rates. It has been found that the model provides reasonable  $NI$  values

only at low  $DR$  values, it predicts essentially parabolic thickness profile across the film width as well as increased  $NI$  values with increased  $DR$  [105]. The convergence is almost guaranteed [89]. At high DRs, the Newtonian model predicts artificially high neck-in, and there are also discrepancies between experiments and temperature profile predictions, as shown for PET in [90] (see Figure 13). This constitutive equation was used in the following studies: 1.D (4 works) [80], [82], [22], [26]; 1.5D (10 works) [18], [100], [86], [87], [88], [27], [89], [90], [53], [97]; 2D (10 works) [112], [12], [28], [101], [102], [39], [104], [105], [113], [111]; 3D (1 work) [114].

### 3.2.2 Generalized Newtonian model

The generalized Newtonian model is simply the Newtonian model given by Eq. 20, in which  $\eta_0$  is replaced by a viscosity scalar function  $\eta(\Pi_{\underline{D}})$ , which is allowed to vary with the second invariant of deformation rate tensor  $\Pi_{\underline{D}}$  defined as  $2tr(\underline{\underline{D}}^2)$ . In the simple shear flow,  $\Pi_{\underline{D}} = \dot{\gamma}^2$ , uniaxial extensional flow,  $\Pi_{\underline{D}} = 3\dot{\epsilon}^2$ , and planar extensional flow,  $\Pi_{\underline{D}} = 4\dot{\epsilon}^2$ . Here  $\dot{\gamma}$  and  $\dot{\epsilon}$  represents shear and extensional strain rate, respectively.

*Power-law (or Ostwald–de Waele) model* [117–119]

$$\eta(\Pi_{\underline{D}}) = m \left( \sqrt{\Pi_{\underline{D}}} \right)^{n-1} \quad (22)$$

where  $m$  (the flow consistency index) and  $n$  (the flow behavior index, which is lower than 1 for polymer melts) are adjustable parameters. This model allows to model shear, uniaxial and planar extensional viscosities plotted as the function of deformation rates as a simple line in log-log scale with the slope equal to  $n-1$ . The key advantage of this model its mathematical simplicity and low number of used parameters. A key disadvantage of this model is over

prediction of shear and extensional viscosities at low deformation rates, the incapability to describe a smooth transition from Newtonian to non-Newtonian flow regime and the incapability to represent fluid elasticity, memory and extensional rheology. It was found that model predictions start to significantly deviate from the Newtonian solution when the power-law index  $n$  becomes less than 0.8 [113]. This model was utilized in the following studies within this review: 1.D (1 work) [81]; 1.5D (0); 2D (2 works) [113], [8]; 3D (0).

*Cross model* [120]

$$\eta(\underline{\Pi}_{\underline{D}}) = \eta_{\infty} + \frac{\eta_0 - \eta_{\infty}}{1 + \left(\lambda \sqrt{\underline{\Pi}_{\underline{D}}}\right)^a} \quad (23)$$

The model was used in: 1.D (0); 1.5D (3 works) [91], [93], [96]; 2D (1 work) [105]; 3D (0).

*Carreau model* [121]

$$\eta(\underline{\Pi}_{\underline{D}}) = \eta_{\infty} + \frac{\eta_0 - \eta_{\infty}}{\left[1 + \left(\lambda \sqrt{\underline{\Pi}_{\underline{D}}}\right)^2\right]^{\frac{1-n}{2}}} \quad (24)$$

The model was used in: 2D (2 works) [110], [111]; 3D (1 work) [115].

*Carreau-Yasuda model* [122]

$$\eta(\underline{\Pi}_{\underline{D}}) = \eta_{\infty} + \frac{\eta_0 - \eta_{\infty}}{\left[1 + \left(\lambda \sqrt{\underline{\Pi}_{\underline{D}}}\right)^a\right]^{\frac{1-n}{a}}} \quad (25)$$

The model was used in: 1.D (1 work) [79]; 1.5D (0); 2D (1 work) [40]; 3D (0).

The above viscosity models use the following parameters:  $\eta_0$  (zero-shear-rate viscosity),  $\eta_{\infty}$  (infinite-shear-rate viscosity),  $\lambda$  (relaxation time),  $a$  (characterizes the sharpness of the transition from Newtonian to non-Newtonian flow regime) and  $n$  (characterizes the slope between viscosity and deformation rate in a log-log scale) are model parameters. Utilization of low number of parameters, mathematical simplicity, capability to represent steady shear viscosity of polymer melts in a wide range of shear rates and correct predictions of steady

extensional viscosities at low extensional rates (i.e. equal to  $3\eta_0$  and  $4\eta_0$  for uniaxial and planar extensional viscosities, respectively) represent the advantages of these models. The main disadvantages are the inability to represent fluid memory and extensional rheology for branched polymers.

The use of generalized Newtonian models in film casting modeling has made it possible to capture some very important trends observed experimentally, namely that  $NI$  intensity and the edge bead increase with  $DR$  or that an increase in planar to uniaxial extensional viscosity ratio increases  $NI$ , in agreement with the viscoelastic PTT model [40], [110].

### 3.2.3 Upper-Convected Maxwell (UCM) model

One of the simplest model allowing to represent some basic viscoelastic features of polymer melts is called the Upper-Convected Maxwell model, which is given by the Eq. 26.

$$\underline{\underline{\tau}} + \lambda \overset{\nabla}{\underline{\underline{\tau}}} = 2\eta_0 \underline{\underline{D}} \quad (26)$$

As can be seen, the key difference between the Newtonian and the Upper-Convected Maxwell

model is the elastic term  $\lambda \overset{\nabla}{\underline{\underline{\tau}}}$ , which consists of the relaxation time,  $\lambda$ , and the upper-convected

time derivative of the stress tensor,  $\overset{\nabla}{\underline{\underline{\tau}}}$ , defined as

$$\overset{\nabla}{\underline{\underline{\tau}}} = \frac{\partial \underline{\underline{\tau}}}{\partial t} + (\mathbf{v} \cdot \nabla) \underline{\underline{\tau}} - \underline{\underline{L}}_v^T \underline{\underline{\tau}} - \underline{\underline{\tau}} \underline{\underline{L}}_v \quad (27)$$

The key advantage of this model is its mathematical simplicity, low number of used parameters ( $\lambda$ ,  $\eta_0$ , or alternatively  $\lambda$  and the elastic modulus  $G$  where  $\eta_0 = \lambda G$ ), it takes into account the melt memory and the first normal stress difference,  $N_1$ , is predicted to be nonzero. Disadvantages: The model predicts unrealistically strong extensional strain hardening without the ability to predict the extensional strain thinning and it yields an infinite steady uniaxial and

planar extensional viscosities at  $\dot{\epsilon} = \frac{1}{2\lambda}$ , as can be deduced from Eqs. 28–29 resulting from this

model for steady uniaxial,  $\eta_{E,U}$ , and planar,  $\eta_{E,P}$ , extensional viscosities:

$$\eta_{E,U} = \frac{3\eta_0}{(1-2\dot{\epsilon}\lambda)(1+\dot{\epsilon}\lambda)} \quad (28)$$

$$\eta_{E,P} = \frac{4\eta_0}{(1-2\dot{\epsilon}\lambda)(1+2\dot{\epsilon}\lambda)} \quad (29)$$

The model also unrealistically predicts a constant steady shear viscosity (equal to  $\eta_0$ ) and the second normal stress difference,  $N_2$ , equal to zero.

The use of the UCM model made it possible to reveal the qualitative role of melt elasticity in the film casting process. It was found that the edge bead defect was more pronounced in the Newtonian case than in the viscoelastic case (see Figure 14) and that increasing the melt elasticity (i.e. the Deborah number) reduces  $NI$  [28], which is in good agreement with the experimental results. This constitutive equation was utilized in the following studies: 1.D (2 works) [85], [83]; 1.5D (8 works) [27], [89], [53], [94], [99], [29], [98], [44]; 2D (4 works) [113], [28], [8], [106]; 3D (0).

### 3.2.4 Generalized Upper-Convected Maxwell model

The generalized Upper-Convected Maxwell model is simply the UCM model, in which the relaxation time and the shear viscosity are allowed to vary with the second invariant of the deformation rate tensor:

$$\underline{\underline{\tau}} + \lambda \left( \Pi_{\underline{\underline{D}}} \right) \overset{\nabla}{\underline{\underline{\tau}}} = 2\eta \left( \Pi_{\underline{\underline{D}}} \right) \underline{\underline{D}} \quad (30)$$



For the film casting modeling, the shear viscosity,  $\eta(\Pi_{\underline{D}})$ , was chosen as the Carreau function (Eq. 24 with  $\eta_{\infty} = 0$ ) and the relaxation time,  $\lambda(\Pi_{\underline{D}})$ , in the form of the Eq. 31.

$$\lambda(\Pi_{\underline{D}}) = \frac{\lambda_0}{\left[1 + \left(\lambda_t \sqrt{\Pi_{\underline{D}}}\right)^2\right]^{n'}} \quad (31)$$

In this equation,  $\lambda_0$ ,  $\lambda_t$  and  $n'$  are adjustable parameters. The model can represent steady shear viscosity,  $N_1$  and extensional rheology more realistically than the UCM model but it still shares the key disadvantages of the original model, i.e.,  $N_2$  is predicted to be zero and steady uniaxial and planar extensional viscosities becomes infinite, if the extensional strain rate becomes equal to  $\frac{1}{2\lambda(\Pi_{\underline{D}})}$ . This model was used in the film casting modeling (1.D (1 work in total) [83]) to understand the role of the power-law exponent  $n$  and the characteristic relaxation time on the velocity profile and the relationship between  $DR$  and tensile force.

### 3.2.5 Giesekus model

The Giesekus model was proposed in 1966 from the simple dumbbell theory for dilute solutions considering anisotropic drag [119, 123–125]. The model is given as follows:

$$\underline{\underline{\tau}} = \underline{\underline{\tau}}_p + \underline{\underline{\tau}}_s \quad (32)$$

$$\underline{\underline{\tau}}_s = 2\eta_{\infty}\underline{\underline{D}} \quad (33)$$

$$\underline{\underline{\tau}}_p + \lambda_1 \frac{\alpha}{\eta_p} \underline{\underline{\tau}}_p^2 + \lambda_1 \overset{\nabla}{\underline{\underline{\tau}}}_p = 2\eta_p \underline{\underline{D}} \quad (34)$$

$$\eta_0 = \eta_p + \eta_{\infty} \quad (35)$$

$$\eta_0 = \lambda_1 G \quad (36)$$

$$\eta_{\infty} = \eta_0 \frac{\lambda_2}{\lambda_1} \quad (37)$$

where  $\underline{\underline{\tau}}$  is the extra-stress tensor,  $\underline{\underline{\tau}}_p$  and  $\underline{\underline{\tau}}_s$  is the polymer and the solvent contribution to the stress tensor,  $\eta_{\infty}$  is the solvent viscosity,  $\eta_p$  is the polymer viscosity,  $D$  is the deformation rate tensor,  $\lambda_l$  is the relaxation time, the symbol  $\nabla$  represents the upper-convected time derivative,  $\eta_0$  is the zero-shear-rate viscosity,  $G$  is the modulus and  $\alpha$  is the parameter characterizing anisotropic hydrodynamic drag. The minimum and maximum anisotropies correspond to  $\alpha = 0$  and  $\alpha = 1$ , respectively [125], but as shown by Bird [119], the model behaves realistically only if  $\alpha \leq 0.5$ . This model can represent a steady shear viscosity of polymer melts in very wide shear rate range and it correctly predicts non-zero values of  $N_1$  as well as negative value of  $N_2$ . On the other hand, its behavior in an extensional flow is not realistic. The key disadvantage of this model is the inability to predict the decrease in extensional viscosity, if the extensional strain rate increases. The model has been found to provide reasonable predictions for the film neck-in, the centerline velocity profile and the temperature drop in the air-gap and it also predicts an increase in film neck-in and the temperature drop in the air-gap as the air-gap length is increased [107].

This constitutive equation has been used in the following studies: 1.D (1 work) [79]; 1.5D (0) ; 2D (3 works) [113], [107], [111]; 3D (0).

### 3.2.6 Modified Giesekus model

The original Giesekus model is not able to realistically represent extensional flows because the polymeric chains are assumed to be infinitely extensible. In order to overcome this model drawback, Wiest [126] modified the original Giesekus model [124] by incorporating the finite extensibility of polymer molecules by using the Peterlin approximation. The set of equations

remains the same as in the case of the original Giesekus model except for Eq. 34, which must be replaced by the following expression

$$\left[ Z - \lambda_1 \frac{D \ln(Z)}{Dt} \right] \tau_{=p} + \frac{\alpha \lambda_1 Z}{\eta_p} \tau_{=p}^2 + \lambda_1 \tau_{=p}^\nabla = 2\eta_p \left[ \underline{\underline{D}} + \frac{1}{2} \frac{D \ln(Z)}{Dt} \underline{\underline{\delta}} \right] \quad (38)$$

where  $\underline{\underline{\delta}}$  is the Kronecker delta and  $Z$  is the function defined as

$$Z = \frac{1}{b_c} \left[ b_c + 3 + \frac{\text{tr}(\tau_{=p})}{G} \right] \quad (39)$$

The term  $D()/Dt$  represents substantial time derivative, which is defined as

$$\frac{D(\ )}{Dt} = \frac{\partial(\ )}{\partial t} + \mathbf{v} \cdot \nabla(\ ) \quad (40)$$

In this model  $b_c$  represents the chain extensibility parameter. Note that if  $b_c \rightarrow \infty$ ,  $Z \rightarrow 1$ , the modified model is reduced to the original Giesekus model whereas for  $\alpha = 0$ , the ‘‘FENE-P’’ dumbbell constitutive equation is recovered [126]. This model shares the same advantages with the original Giesekus model with the additional ability to describe extensional strain hardening as well as extensional strain thinning for steady uniaxial as well as planar extensional viscosities. In addition, the model is derived from kinetic theory, which allows to relate model parameters with molecular characteristics. On the other hand, the model predicts that both uniaxial and planar extensional viscosities are very similar at high deformation rates [84], which might not be realistic, as indicated in [127].

The model was used to investigate the role of the uniaxial extensional strain hardening on velocity and stress profiles and film tension in single layer as well as the multilayer film casting process by using 1.D flow kinematics [84], [75], [76].

### 3.2.7 eXtended Pom-Pom model

The eXtended Pom-Pom (XPP) model was proposed by Verbeeten et al. [128]. This model represents an approximation of the original Pom-Pom model proposed by McLeish and Larson [129], which is based on the Doi-Edwards reptation tube theory and uses the Giesekus anisotropy parameter  $\alpha$ . The model considers a simplified H topology of branched molecules and the relaxation time is expressed as a tensor to separate the stretch and orientation. The model is given by the following set of equations:

$$\underline{\underline{\underline{\tau}}} + \underline{\underline{\underline{\lambda}}}(\underline{\underline{\underline{\tau}}})^{-1} \underline{\underline{\underline{\tau}}} = 2G \underline{\underline{\underline{D}}} \quad (41)$$

where the relaxation time tensor is defined as

$$\underline{\underline{\underline{\lambda}}}(\underline{\underline{\underline{\tau}}})^{-1} = \frac{1}{\lambda_{ob}} \left\{ \frac{\alpha}{G} \underline{\underline{\underline{\tau}}} + f(\underline{\underline{\underline{\tau}}})^{-1} \underline{\underline{\underline{\delta}}} + G \left[ f(\underline{\underline{\underline{\tau}}})^{-1} - 1 \right] \underline{\underline{\underline{\tau}}}^{-1} \right\} \quad (42)$$

Extra function:

$$f(\underline{\underline{\underline{\tau}}})^{-1} = \frac{2\lambda_{ob}}{\lambda_s} \left( 1 - \frac{1}{\Lambda} \right) + \frac{1}{\Lambda^2} \left( 1 - \frac{\alpha \text{tr}(\underline{\underline{\underline{\tau}}})^2}{3G^2} \right) \quad (43)$$

Backbone stretch and stretch relaxation time:

$$\Lambda = \sqrt{1 + \frac{\text{tr}(\underline{\underline{\underline{\tau}}})}{3G}}, \quad \lambda_s = \lambda_{os} e^{-v(\Lambda-1)}, \quad v = \frac{2}{q} \quad (44)$$

where  $q$  (number of arms),  $\lambda_{os}$  (stretch relaxation time) are adjustable parameters, which are allowed to vary with the orientation relaxation time,  $\lambda_{ob}$ . Note that the Maxwell parameters are  $G$  and  $\lambda_{ob} = \lambda$ . The model has an excellent capability to describe the shear and extensional rheology for long-chain branched polymers such as LDPE, which is widely used in the film casting technology. The model also predicts non-zero values of  $N_1$  and  $N_2$  as it should be. Additionally, the model parameters are directly related to the molecular characteristics because the model is derived from molecular arguments. On the other hand, the model is not suitable

for linear polymers due to the assumed structural topology and uses a very high number of adjustable parameters, which makes it difficult to identify them from the measured experimental data. The model (similarly to the original Pom-Pom model) also predicts that steady uniaxial and planar extensional viscosities becomes practically identical at high extensional strain rates, which might not be realistic for some LDPEs, as indicated in [127] (see Figure 15). The XPP model correctly predicts that increasing  $DR$  and the air-gap reduce  $N_I$ . According to the authors, the agreement between the experimental data (LDPE [41], [42] and long chain branched PP [43]) and the simulation results was qualitative rather than quantitative in terms of necking behavior (see Figure 16 as the example for LDPE; here the 8 mode XPP model, in which all model parameters were allowed to vary with the relaxation mode, was used; step shear and step uniaxial extensional experiments were used to validate the XPP model). This constitutive equation was utilized in the following studies: 1.D (0); 1.5D (3 works) [41], [42], [43]; 2D (0); 3D (0).

### 3.2.8 Rolie–Poly Stretch (RP-S) model

Rolie–Poly (ROuse LLinear Entangled POLYmer) stretch model is a tube-based model proposed by Likhtman and Graham [130], which takes into account the convective constraint release, reptation and chain retraction. The model is given as

$$\underline{\underline{\tau}}^{\nabla} = -\frac{1}{\lambda_d}(\underline{\underline{\tau}} - \underline{\underline{\delta}}) - \frac{2(1 - \sqrt{3/\text{tr}\underline{\underline{\tau}}})}{\lambda_r} \left( \underline{\underline{\tau}} + \beta_c \left( \frac{\text{tr}\underline{\underline{\tau}}}{3} \right)^{\delta_0} (\underline{\underline{\tau}} - \underline{\underline{\delta}}) \right) \quad (45)$$

where  $\lambda_d$  and  $\lambda_r$  are the reptation and the Rouse relaxation times, respectively;  $\beta_c$  is the convective constraint release coefficient,  $\delta_0$  is a fitting scalar parameter. Being  $\beta_c$  equal to zero,  $\delta_0$  parameter does not have to be specified. This model has showed a good capability to describe the transient shear and extensional rheology of linear film casting resins (namely LLDPE and HDPE), it is mathematically simple and gives a non-zero value of  $N_I$ . On the other

hand, the model is not able to describe the rheological behavior of branched polymers and unrealistically predicts  $N_2 = 0$  [131]. This constitutive equation was utilized in the following theoretical studies summarized in this review: 1.D (0); 1.5D (3 works) [41], [42], [43]; 2D (0); 3D (0). The model predicted an increase in  $NI$  with increased  $DR$  and the air-gap. According to the authors, the agreement between the experimental data (HDPE [41]; LLDPE [41], [42]; and linear PP [43]) and the simulation results was qualitative rather than quantitative with respect to  $NI$  (see Figure 17 as an example for LLDPE; here the 8 mode RP-S model, in which  $\lambda_d$  and  $\lambda_r$  were allowed to vary with the relaxation mode, was used;  $\beta = 0$  and  $\delta_0 = -0.5$ ; step shear and step uniaxial extensional experiments were used to validate the RP-S model).

### 3.2.9 Modified Leonov model

The modified Leonov model is based on heuristic thermodynamic arguments resulting from the theory of rubber elasticity [132–137]. In this constitutive equation, a fading memory of the melt is determined by an irreversible dissipation process driven by the dissipation term,  $b$ . This model relates the stress and elastic strain stored in the polymer melt as:

$$\underline{\underline{\tau}} = 2 \left( \underline{\underline{c}} \frac{\partial W}{\partial \underline{\underline{I}}_{\underline{\underline{c}}}} - \underline{\underline{c}}^{-1} \frac{\partial W}{\partial \underline{\underline{II}}_{\underline{\underline{c}}}} \right) \quad (46)$$

where  $W$  is the elastic potential, which depends on the invariants  $\underline{\underline{I}}_{\underline{\underline{c}}}$  and  $\underline{\underline{II}}_{\underline{\underline{c}}}$  of the recoverable Finger tensor  $\underline{\underline{c}}$ ,

$$W = \frac{3G}{2(n_L + 1)} \left\{ (1 - \beta) \left[ \left( \frac{\underline{\underline{I}}_{\underline{\underline{c}}}}{3} \right)^{n_L + 1} - 1 \right] + \beta \left[ \left( \frac{\underline{\underline{II}}_{\underline{\underline{c}}}}{3} \right)^{n_L + 1} - 1 \right] \right\} \quad (47)$$

where  $G$  denotes the linear Hookean elastic modulus,  $\beta$  and  $n_L$  are numerical parameters.

Leonov assumed that the dissipative process acts to produce an irreversible rate of strain  $\underline{\underline{e}}_p$

$$\underline{\underline{e}}_p = \underline{\underline{b}} \left[ \underline{\underline{c}} - \frac{\underline{\underline{I}}_c}{3} \underline{\underline{\delta}} \right] - \underline{\underline{b}} \left[ \underline{\underline{c}}^{-1} - \frac{\underline{\underline{\Pi}}_c}{3} \underline{\underline{\delta}} \right] \quad (48)$$

which spontaneously reduces the rate of elastic strain accumulation. Here,  $\underline{\underline{\delta}}$  is the unit tensor and  $\underline{\underline{b}}$  stands for the dissipation function defined by Eq. 50. This elastic strain  $\underline{\underline{c}}$  is related to the deformation rate tensor  $\underline{\underline{D}}$  as follows

$$\overset{\circ}{\underline{\underline{c}}} - \underline{\underline{c}} \cdot \underline{\underline{D}} - \underline{\underline{D}} \cdot \underline{\underline{c}} + 2\underline{\underline{c}} \cdot \underline{\underline{e}}_p = 0 \quad (49)$$

where  $\overset{\circ}{\underline{\underline{c}}}$  is the Jaumann (corotational) time derivative of the recoverable Finger strain tensor. The dissipation function  $\underline{\underline{b}}$  proposed in [64] is given as

$$\underline{\underline{b}}(\underline{\underline{I}}_c) = \frac{1}{4\lambda} \left\{ \exp \left[ -\xi \sqrt{\underline{\underline{I}}_c - 3} \right] + \frac{\sinh \left[ \nu (\underline{\underline{I}}_c - 3) \right]}{\nu (\underline{\underline{I}}_c - 3) + 1} \right\} \quad (50)$$

where  $\xi$  and  $\nu$  are adjustable parameters of the model.

The model has a very good capability to describe shear viscosity,  $N_1$ ,  $N_2$ , uniaxial and planar extensional viscosities for linear as well as branched polymers [57, 58, 62–64, 138]. The model also offers an independent control of uniaxial and planar extensional viscosities, which was used for systematic investigation of the role of planar to uniaxial extensional viscosity ratio on the film casting process for different LDPEs. On the other hand, the interpretation of molecular meaning of the used model parameters is limited because the model is derived from thermodynamics rather than molecular arguments. Note that the original Leonov model is recovered if  $n_L = \beta = \xi = \nu = 0$ .

The model demonstrated the ability to describe film casting experimental data for linear (PP) as well as branched (LDPE) polymers even by using a single mode (i.e. utilizing a single pair of  $\lambda$  and  $G$ ) only [35, 37, 38] (see Figure 18; In the case of PP, the lowest relaxation time typical for polyolefins was chosen and  $G$  was calculated from the known  $\eta_0$ . The nonlinear model

parameters were adjusted as the typical values for linear polymers, i.e.  $\xi = 0$ ,  $n_L = 0$ ,  $\nu = 0.5$ ,  $\beta = 0.5$ . In the case of LDPE, the model parameters, namely  $\lambda$ ,  $G$ ,  $\xi$ ,  $\nu$ ,  $\beta$  were identified using deformation rate dependent ‘steady state’ uniaxial extensional viscosity experimental data and known  $\eta_0$ . The parameter  $n_L$  was kept equal to zero). The modified Leonov model was used in the following studies: 1D (0); 1.5D (3 works) [37], [38], [35]; 2D (0); 3D (0). The original Leonov model was used only in: 1.5D (1 work) [25].

### 3.2.10 Phan-Thien-Tanner (PPT) model

The model was derived by Phan-Thien and Tanner [139] and Phan-Thien [140] from the Lodge-Yamamoto network theory, in which junctions are allowed to form and decay due to the flow. The model is given by the following equation

$$\underline{\underline{\tau}} + \lambda(\underline{\underline{\tau}}) \overset{\diamond}{\underline{\underline{\tau}}} = 2\eta_0 \underline{\underline{D}} \quad (51)$$

This model uses the Gordon–Schowalter convected time derivative of the stress tensor,  $\overset{\diamond}{\underline{\underline{\tau}}}$ , defined as

$$\overset{\diamond}{\underline{\underline{\tau}}} = \frac{\partial \underline{\underline{\tau}}}{\partial t} + (\mathbf{v} \cdot \nabla) \underline{\underline{\tau}} - \underline{\underline{\ell}}^T \underline{\underline{\tau}} - \underline{\underline{\tau}} \underline{\underline{\ell}}, \quad \underline{\underline{\ell}} = \underline{\underline{L}}_v - \xi_p \underline{\underline{D}} \quad (52)$$

and the relaxation time,  $\lambda(\underline{\underline{\tau}})$ , which is allowed to vary with the trace of the stress linearly (Eq. 53) or exponentially (Eq. 54).

*Linear PTT model:*

$$\lambda(\underline{\underline{\tau}}) = 1 + \frac{\varepsilon_p}{G} \text{tr}(\underline{\underline{\tau}}) \quad (53)$$

*Exponential PTT model:*

$$\lambda(\underline{\underline{\tau}}) = \exp \left[ \frac{\varepsilon_p}{G} \text{tr}(\underline{\underline{\tau}}) \right] \quad (54)$$



The model utilizes two parameters,  $\xi_p$  and  $\varepsilon_p$  (together with  $\lambda$  and  $G$ , where  $\eta_0 = \lambda G$ ). The linear PTT predicts an unrealistically monotonically increasing extensional viscosity, while the exponential PTT has the ability to give a maximum in the extensional viscosity, when plotted as a function of the extensional strain rate. Thus, the exponential PTT model is used more in modeling of the film casting process than its linear version; it has a good ability to describe shear as well as extensional rheology of linear as well as branched polymers and predicts non-zero values for  $N_1$  and  $N_2$ . On the other hand, steady uniaxial and planar extensional viscosities are predicted to be practically identical at high extensional strain rates, which might not be realistic for some polymers, as indicated in [127] (see Figure 19). This model has been shown to provide good agreement with experimental data for LDPE over a wide range of the take-up velocity and the air-gap length [109], [54] (see Figure 20). In this case, the exponential (6-7) mode PTT model using  $\varepsilon_p$  and  $\xi_p$  model parameters varying with each relaxation mode was necessary to adequately represent the measured shear and extensional data LDPEs reasonably. The discrete relaxation spectrum was determined from frequency-dependent loss and storage moduli, while  $\xi_p$  and  $\varepsilon_p$  parameters were identified from steady state shear viscosity vs. shear rate and steady uniaxial extensional viscosity vs. extensional strain rate.

This constitutive equation was utilized in the following studies within this review: 1.D (1 work) [79]; 1.5D (2 works) [47], [44]; 2D (7 works) [46], [105], [36], [108], [109], [54], [40]; 3D (1 work) [116]. The linear variant of the PTT model was used only in: 2D (1 work) [26].

### 3.2.11 Larson model

This model represents a differential approximation of the Doi-Edwards (DE) integral tube model proposed by Larson [141, 142], which includes non-affine motion and is given by the following expression:

$$\underline{\underline{\tau}} + \lambda \overset{\nabla}{\underline{\underline{\tau}}} + \frac{2}{3} \frac{\zeta \lambda}{G} \underline{\underline{D}} : \underline{\underline{\tau}} (\underline{\underline{\tau}} + G \underline{\underline{\delta}}) = 2\eta_0 \underline{\underline{D}} \quad (55)$$

where is  $\zeta$  the only non-linear model parameter (note that the best approximation of the DE model is achieved, if  $\zeta = 3/5$  [125]). The behavior of the model is comparable to the exponential PTT model meaning that it can capture shear and extensional viscosities for both linear and branched polymers, and  $N_I$  is predicted to be a non-zero value. However, the model unrealistically predicts  $N_2 = 0$ , steady uniaxial and planar extensional viscosities becomes comparable at high extensional strain rates, and  $\zeta$  must be varied with the relaxation time to capture uniaxial extensional strain hardening for commercial types of highly branched polyethylenes over a wide range of extensional strain rates. This model was successfully used in the film casting modeling for 4 different types of LDPEs, but 13 pairs of  $\lambda$ ,  $G$ ,  $\zeta$  were needed to describe the experimental reality [39]. The model provides reasonably good predictions for the neck-in and the film thickness (see Figure 21). This model also predicts that the increase in the uniaxial extensional strain hardening firstly, decreases  $NI$  in agreement with experimental observations (see Figure 22), and secondly, the edge beading region becomes narrower.

This constitutive equation was utilized in the following studies: 1.D (0); 1.5D (0) ; 2D (2 works) [39], [105]; 3D (0).

### 3.2.12 Integral constitutive equation of the K-BKZ type with PSM strain-memory function

The multi-mode Kaye-Bernstein-Kearsley-Zapas (K-BKZ) type of the integral model proposed by Papanastasiou, Scriven, Macosko (PSM) [143] and then modified by Luo and Tanner [144] to account for  $N_2$  through the parameter  $\theta$  has the following expression:

$$\underline{\underline{\tau}} = \frac{1}{1-\theta} \int_{-\infty}^t \sum_{j=1}^N \frac{G_j}{\lambda_j} \exp\left(-\frac{t-t'}{\lambda_j}\right) \frac{\alpha_j}{\alpha_j - 3 + \beta_j I_{\underline{\underline{C}}^{-1}} + (1-\beta_j) II_{\underline{\underline{C}}^{-1}}} \left[ \underline{\underline{C}}^{-1}(t') + \theta \underline{\underline{C}}(t') \right] dt' \quad (56)$$

where  $t$  and  $t'$  are the times present and past, respectively,  $I_{\underline{\underline{C}}^{-1}}$  and  $II_{\underline{\underline{C}}^{-1}}$  are the first and second invariant of the Finger strain tensor  $\underline{\underline{C}}^{-1}$ ;  $\alpha_j$ ,  $\beta_j$  and  $\theta$  are the parameters of the model. The K-BKZ model reduces to the upper-convected Maxwell model when  $\theta = 0$  and  $\alpha \rightarrow +\infty$ . The model has a very good ability to describe the shear viscosity,  $N_1$ ,  $N_2$  and uniaxial extensional viscosities for linear and branched polymers. The main disadvantage of the model is the inability to predict extensional strain hardening in planar extensional viscosity for branched polymers as LDPE [145] (see Figure 23).

The model showed a good agreement between experimental data and previous film casting simulations based on the Newtonian and UCM results in terms of film thickness, width and temperature [89]. Note that in order to capture the shear and extensional rheology of LDPE melts, the parameter  $\beta$  in Eq. 56 needs to be varied with the relaxation time, otherwise the extensional strain hardening is overpredicted at high extensional strain rates (see Figure 23).

This constitutive equation was utilized in the following studies: 1.D (1 work) [45]; 1.5D (1 work) [89]; 2D (0); 3D (0).

### 3.2.13 Hooke's Law with creep

Smith and Stolle considered the viscoelastic polymer melt to be an elastic material that is creeping. For such a description, they have suggested the incremental form of Hooke's law of the following form [146]:

$$\underline{\underline{\Delta\sigma}} = \underline{\underline{E}}(\underline{\underline{\Delta\varepsilon}} - \underline{\underline{\Delta\varepsilon}}^c) \quad (57)$$

where  $\underline{\underline{\Delta\sigma}}$  is the stress increment,  $\underline{\underline{E}}$  is the elasticity matrix,  $\underline{\underline{\Delta\varepsilon}}$  is the total strain increment and  $\underline{\underline{\Delta\varepsilon}}^c$  is the modified Perzyna creep strain increment that depends on the time step, the effective creep strain rate and the second invariant of the deviatoric stress tensor. This model was used to theoretically investigate the role of elasticity (by changing the relaxation time) and extensional strain hardening. It was found that the predictions were consistent with predictions of the conventional viscoelastic model i.e. that  $NI$  decreases with increased extensional strain hardening and the relaxation time. Note that the model uses only one constant relaxation time and modulus, and its ability to describe rheological experimental data has not been provided by the authors.

This constitutive equation was utilized in the following studies: 1.D (2 works) [77], [78]; 1.5D (0) ; 2D (1 work) [103]; 3D (0).

### 3.3 Boundary conditions for viscoelastic constitutive equations

If viscoelastic constitutive equations are used, the additional boundary stress condition at the die exit must be specified. This boundary condition is given by both, the flow in the die (upstream) and the extensional flow in the drawing length (downstream). Accurate determination of this additional boundary stress value therefore requires an intensive numerical

computation [46]. The following paragraphs summarize the approaches used to determine these types of boundary conditions (see also Tables 2–5).

Anturkar and Co [83] in their study, using the generalized upper-convected Maxwell model, estimated axial component of the stress tensor,  $\tau_{xx}$ , as the mean stress value for a fully-developed slit flow in an infinitely wide die. Silagy et al. [27] and [28] based on Denn et al. [147] and Demay et al. [88, 148] assumed two different stress states at the end of the die. In the first case, the extra stress in the machine direction,  $\tau_{xx}$ , equals zero, so the extra stresses are completely released due to the die swell, or the second, which assumed the mean value of the extra stress after flow in an infinite die with a rectangular cross section while the transversal extra stress,  $\tau_{yy}$ , is set to the value obtained from the Newtonian solution. They found that the initial stress conditions at the die exit had little effect on the final shape of the film, but calculations were performed only for low Deborah numbers. Iyengar and Co [84] took a different approach, and instead of specifying the axial stress component, set the ratio  $\tau_{zz}/\tau_{xx}$  to the value between two extreme cases for planar extensional flow and a fully-developed slit flow in the die noting that the true stress ratio should have lied in their range. Iyengar [149] then reported that both extreme cases with corresponding stress ratios provide very similar velocity and stress profiles. Debbaut et al. [113] in their viscoelastic study assumed an initial stresses of zero. Same as in Smith's work [8].

For a multilayer film casting analysis (based on a single-mode modified Giesekus model) Pis-Lopez and Co [76, 150] showed that if the aspect ratio (defined here as the ratio of the total film thickness at the die exit to the drawing distance  $2e_0/X$ ) is smaller than 0.05, the velocity and stress profiles converge to the same values, regardless of whether the initial stress condition is based on the assumption of a fully-developed slit flow or a fully-developed planar extensional flow. In another study using a multi-mode model approach, Denn [151] left the longest relaxation mode unspecified at the die exit, and rest of the modes were set with respect to this

mode. In contrast, Christodoulou et al. [46] concluded that the shortest mode should be left unspecified with the reasoning that the longest mode  $\tau_{xx(N)}^p$  is mainly determined by the flow inside the die, while the shorter modes  $\tau_{xx(j)}^p$  are determined by the external flow in the air gap.

Beris and Liu [152, 153] in their fiber spinning study for the single mode UCM viscoelastic liquid specified the die exit stress state through the ratio of normal to axial stress,  $\tau_{yy}/\tau_{xx}$ , and not each component separately. This value was approximated as the value below the homogeneous steady extensional flow at the effective extensional strain rate. For the viscoelastic multimode model, Denn [151] specified also  $\tau_{xx(j)}^p/\tau_{xx(N)}^p$  for  $j < N$  as an extra condition to  $\tau_{yy(j)}^p/\tau_{xx(j)}^p$  for all relaxation modes.

Devereux and Denn [154] proposed the same distribution between partial stresses as in the case of a fully-developed capillary flow with neglected radial partial stresses. The remaining initial stresses were adjusted to satisfy the downstream boundary condition (see Eq. 58).

$$\frac{\tau_{xx(j)}^p}{\tau_{xx(N)}^p} = \frac{\lambda_j G_j}{\sum_{j=1}^N \lambda_j G_j} \quad (58)$$

Note that Gagon and Denn [155] simplified the above-mentioned relationship for the wedge spectrum into

$$\frac{\tau_{xx(j)}}{\tau_{xx(N)}} = \frac{\lambda_j}{\lambda_N} \quad (59)$$

Barborik and Zatloukal [37, 38] recently defined the state of the stress at the die exit by the ratio of the second to the first normal stress difference,  $-N_2/N_1$ , calculated from the upstream side by using the viscoelastic modified Leonov constitutive equation.

$$-\frac{N_2}{N_1} = -\frac{\bar{\tau}_{zz}(0) - \bar{\tau}_{yy}(0)}{\bar{\tau}_{xx}(0) - \bar{\tau}_{zz}(0)} \quad (60)$$

It has been found that if the Deborah number is less than 0.1, the choice of initial stress conditions at the die exit has little effect on the steady-state calculations. However, at higher Deborah numbers, the die exit stress state, which may be influenced, for example, by extrusion die design, starts to significantly affect the neck-in phenomenon.

### 3.4 Overview on steady-state film casting modeling

Initial efforts have been made for a fiber spinning process in which the flow kinematics are mathematically similar when considered as a one-dimensional flow, for the Newtonian and Maxwell fluids by Gelder [156] and Fisher [157, 158], respectively. These studies focused on the draw resonance phenomenon, that Christensen [159] and Miller [160] first encountered, and who postulated that the nature of this phenomenon is not of viscoelastic, as it can also be observed in Newtonian fluids. Extending the kinematics of the process to two or three dimensions, both processes become different and one can observe phenomena in the film casting, which have no counterpart in the fiber spinning, i.e. neck-in and edge-beading. The above preliminary studies provided the background for extended studies on the extrusion film casting (EFC). Initial attempts to simulate EFC operations were devoted to investigating the stability of the process and determining the onset of draw resonance, rather than quantifying the extent of the neck-in phenomenon. The very first study of EFC process modeling in this way was carried out by Yeow [80] using numerical modeling. For the steady state solution, he used a one-dimensional isothermal model for Newtonian fluid (planar extensional free surface flow) and investigated the effect of introduced small two-dimensional perturbances on flow stability (namely transverse perturbations). The edge-effects, surface tension, aerodynamic drag and fluid inertia and gravity were neglected. A small curvature of the film was assumed along with a uniform axial stress and axial velocity over the film thickness. Due to the assumed free surface flow kinematics in the drawing section, the model could not capture

the edge-bead defect and the shrinkage in the width of the film, which was considered to be infinitely wide. The film thickness could only change in the machine direction.

Aird and Yeow [81] continued this mathematical background for the 1-dimensional (1D) model and extended the analysis for power-law fluids. Consequently, Anturkar and Co [83] and Iyengar and Co [75, 84] used isothermal generalized Upper-Convected Maxwell and Giesekus constitutive equations for linear and non-linear analysis in viscoelastic fluid simulations. The first isothermal attempts to model the necking phenomena were performed by Sergent [161] in 1977 and then by Cotto, Duffo and Barq [18, 87, 88] for non-isothermal conditions.

Another milestone work was set by Dobroth and Erwin [55] in 1986, who pointed out that there is a planar and uniaxial extensional deformation at the center of the film and at the edge, respectively, and that the extent of the edge-beads and the interrelated neck-in phenomenon is determined by the interplay between these two regions through an edge stress effect. This idea was consequently confirmed experimentally by Ito et al. [52]. The assumption of a planar-uniaxial distribution in the Dobroth and Erwin model can therefore be considered reasonable. Just note that, in the case of fiber spinning, only a uniaxial extensional flow can be observed.

Some authors have attempted to relate and quantify the gauge of the observed necking in terms of rheological parameters, such as shear, uniaxial and planar extensional viscosity. Many authors have reported that the strain hardening in uniaxial extension may reduce the extent of the necking phenomena [39, 72, 74, 113]. This idea was continued by Ito [53], who related the neck-in intensity to rheological parameters, such as the ratio of planar viscosities in axial and transverse directions, and derived an analytical equation for the edge line of Newtonian and Maxwell fluids. In accordance with the article by Dobroth and Erwin [55], who, as the first recognized deformation type in the drawing area, Shiromoto [7, 8, 33] recently presented the idea that the extent of the necking should not have been described by uniaxial extensional



viscosity only in addition to take-up length but as the ratio of planar and uniaxial extensional viscosities reflecting the deformation type in the central and edge portion of the film in the drawing section. In addition to performing non-isothermal viscoelastic simulations, they also proposed a theoretical model based on the force balance and the deformation type of the film in order to predict necking behavior.

More recently, the 2-dimensional (2D) membrane model has been introduced by d'Halewyu [112] and Debbaut [113] for Newtonian and viscoelastic fluids, respectively. This frequently used model is able to predict the dog-bone defect, i.e. development of edge-beads under stationary conditions.

Silagy et al. [27] improved and enriched the membrane model with a supplementary kinematic hypothesis originally provided by Narayanaswamy [162] in his paper on float glass stretching, and performed an extended isothermal study on the effect of processing conditions on film geometry and EFC stability analysis for Newtonian and Upper convected Maxwell (UCM) fluids. Due to the assumptions used in the flow kinematics, this model was able to cover the reduction of the film width, and thus predict the neck-in phenomenon, but still could not predict the edge-beading. This limitation was removed in their subsequent work [28], where a 2D isothermal membrane model combined with the Phan–Thien and Tanner (PTT) constitutive equation was developed and the obtained steady and transient stability results were compared with its 1.5D predecessor. In the following years, the 1.5D version of Sylagy's membrane model was used in many studies, and much work was done on EFC under non-isothermal conditions including crystallization effects (Lamberti et al. [91–93, 163], Lamberti and Titomanlio [92, 94, 95, 164], and Lamberti [67]). A three dimensional model for EFC simulation was further developed by Sakaki et al. [114]. To solve the model equations, it was necessary to use the finite element method. The problem was considered isothermal, steady state and Newtonian. A process parameter space was chosen to reflect the industrial processing conditions. The model

captured the development of both the neck-in and the edge beading and the effect of  $DR$ ,  $X$  and die width were investigated. They found that the intensity of the neck-in and the edge beading was affected by  $DR$  and  $X$  but not by the width of the die. The extent of the neck-in increased with increasing  $DR$  and  $X$ . Lately, this approach has been extended by Zheng et al. [115] for a non-isothermal steady Newtonian fluid. Kometani et al. [111] performed experimental and theoretical investigations of the effects of rheological properties on the neck-in in the film casting. For the two tested PP and LDPE materials, without significant differences in viscoelastic properties, except for extensional ones (LDPE showed a remarkable increase in extensional viscosity at high strain rates), the neck-in extent for PP under higher draw ratios increased over LDPE where the neck-in was constant and independent of the draw ratio. Based on these experiments, the authors concluded that neck-in phenomenon in film casting depends on the extensional rheological properties. Furthermore, they utilized a simulation based on three different rheological models (the Newtonian, Bird-Carreau and Giesekus model) to assess its applicability for film casting modeling. The results obtained by simulation based on the Giesekus model were in quantitative agreement with the experimental observations for both polymers, but the other two models used did not provide predictions well describing the measured data due to their inaccurate representation of extensional viscosities.

The influence of macromolecular architecture on the extent of the necking phenomenon has been investigated by Ito et al. [52, 53] (effects of the draw ratio and the take-up length on the necking for LDPE, HDPE and mLLDPE) and Baird et al. [68, 69] (effects of long chain branching and molecular weight distribution on the necking for LDPE, mLLDPE and Ziegler-Natta catalyzed LLDPE). Research on a multi-layer film casting considering Giesekus fluid has been performed in studies of Pis-Lopez and Co for steady state [76] and stability analysis [150].

Recently, Pol et al. [41, 42] and Chikhalikar et al. [43] published a number of articles in which they performed experimental and theoretical investigations of the effects of long chain

branching and molecular weight distribution on the extent of the necking phenomenon. For this purpose, they have used the 1.5D membrane model, originally proposed by Silagy [27], the multi-mode eXtended Pom-Pom constitutive equation and the multi-mode Rolie-Poly stretch constitutive equation, respectively, for the long chain branched (LDPE, PP) and the linear (HDPE, PP) polymers. Fixing the  $DR$  and  $X$ , they found that the extent of the necking was smaller for HDPE with a broader molecular weight distribution than for LLDPE with a narrower molecular weight distribution, and further that long chain branched LDPE necks-in to lower extent than linear HDPE or LLDPE (i.e. that increase in long chain branching is more effective in neck-in suppression than just broadening  $MWD$ ). In a subsequent study, Pol and Thete [98] switched from the 1.5D model that was used in their predecessor works on this topic to the two-dimensional model originally proposed by Ito et al. [53] incorporating UCM constitutive equation. In addition, they derived an analytical solution for low and high Deborah numbers. They found that while the film width of the modelled LLDPE continuously decreased with increased draw ratio, the film width for LDPE decreased with increased draw ratio for long take-up lengths and remained constant for shorter ones. This means that there is a locus of points in the attainable region that divides  $DR-De$  plane into sections where the dependence of the neck-in on the draw ratio has opposite trends.

In their latest work [44], they focused on the effects of the individual viscoelastic relaxation modes of a polymer melt on its behavior in polymer melt extrusion film casting process using UCM and PTT constitutive equations and the 1.5D isothermal membrane model. They found that experimental data for long-chain branched LDPE was better described using the UCM model, while the PTT model provided better simulation results for the experimental data with linear LLDPE.

Although the actual EFC manufacturing process involves complex 3-dimensional (3D) kinematics whose numerical simulation can be very challenging, many authors have shown that

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the EFC 1.5D membrane model (originally proposed by Silagy) is able to provide results that are in good agreement with experiment data (if appropriate constitutive equations are used).

## 4 CONCLUSIONS

Modeling of the steady-state film casting process began in 1974 using a 1D kinematic approach. 1.5D and 2D kinematic models were developed at 1990 and are currently the most commonly used (in more than 73%). A full 3D kinematic approach was introduced in 1996, but due to its complexity, its use is very rare (less than 5%) and usually only applies to viscous fluids. The following most commonly used simplifying assumptions are used to overcome numerical difficulties in steady-state cast film process modeling: reduction in dimensionality (1D, 1.5D and 2D), isothermal conditions – constant temperature field, non-transient description, incompressible fluid, constant density, excluded inertia effects, excluded effects of gravitational forces, constant boundary conditions, unrealistic or simplified constitutive equations, neglected aerodynamic drag, neglected surface tension, neglected die swell, neglected edge-effects, excluded crystallization (temperature, flow-induced), neglected the sag of the film in non-vertical installations (film curvature), effects of other devices (air knife, vacuum box, electrostatic pinning).

The following constitutive equations have already been used in the steady-state modeling of the extrusion cast film process: *Viscous* – Newtonian model, Generalized Newtonian model (utilizing Cross, Carreau and Carreau-Yasuda viscosity functions); *Elastic* – incremental Hooks law with creep. *Viscoelastic* – upper convected Maxwell model, generalized upper convected Maxwell model with deformation rate dependent relaxation time and viscosity, linear and exponential PTT model, Larson model, K-BKZ model with Papanastasiou-Scriven-Macosko (PSM) damping function, Leonov model modified by specific type of recoverable strain dependent dissipation function, Giesekus model, modified Giesekus with finite chain stretch, eXtended Pom-Pom model (XPP) and RoliePoly-Stretch model (RP-S).

Intensive experimental research on extrusion film casting in relation to the neck-in began in 1986 and continues to this day. Most of the research reported in the open literature focuses on

polyolefins such as LDPE, LLDPE and PP, while experimental studies for other polymers (PET, HDPE or polyethersulfone - PES) are very rare.

Based on experimental and theoretical studies presented in the open literature, the following variables were found to be the key with respect to the neck-in phenomenon: draw ratio, Deborah number (i.e. melt relaxation time, melt velocity at the die exit, air-gap), film cooling, ratio of the second and the first normal stress difference at the die exit, uniaxial extensional strain hardening and planar-to-uniaxial extensional viscosity ratio. Uniaxial and planar extensional viscosities can be considered as key material parameters, because during the polymer melt stretching in the post die area, the middle of the film undergoes planar elongation, while the material at the edge undergoes uniaxial elongation. However, measuring uniaxial and planar extensional viscosities at very wide deformation rates is still a very difficult task because the generation and control of extensional flows is difficult. Likewise, the role of draw ratio, heat transfer coefficients, flow induced crystallization, flow history (generated inside the extrusion die as  $-N_2/N_1$  ratio), molecular architecture of polymer melts on the film formation and its stability is still not fully understood yet due to the absence of relevant experimental and theoretical studies.

There is a currently attempt to relate flow stability with the molecular characteristics of the polymers used through advanced molecular constitutive equations such as XPP and RP-S. However, their use is still limited because these models do not allow realistic rheological description of polymers or polymer mixtures with an internal structure, which does not correspond to the molecular assumptions under which these constitutional equations were derived. Another major problem in experimental and theoretical flow stability research is the neglect of the influence of memory and flow history of polymer melts in extrusion heads on the stability of film formation in the post die area, as evidenced by recently published work.

Further research in the following areas can be expected to help significantly understand the formation stability of polymeric flat films:

- development and use of suitable rheological techniques for the determination of uniaxial and especially planar extensional viscosities at high deformation rates,
- development/modification and use of advanced constitutive equations with ability to handle measured extensional rheology data for polymers, polymer blends and filled polymers,
- elucidation of the role of polymer molecular architecture, flow history, heat transfer coefficients and flow induced crystallization,
- understanding the role of the draw ratio in film formation for branched polymers.

#### *Data availability statement*

Data sharing is not applicable to this review article as no new experimental data were generated.

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## 6 TABLES

**Table 1.** Research work devoted to the experimental investigation of the neck-in phenomenon.

YEAR	REFERENCE	MATERIAL	NOTE
1974	Kase [165]	PP	Experimental investigation of EFC aimed on process stability.
1986	Dobroth and Erwin [55]	LDPE	Enlightens the physical background of Edge-beads formation.
1989	Cotto et al. [18]	PP	Experimental and theoretical investigation of crystalline phase development during EFC and modelling.
1990	Barq et al. [22]	PET	The work is aimed on transient phenomena of draw resonance.
1990	Duffo et al. [100]	PP	Effect of roll temperature on crystallization in EFC process.
1991	Duffo et al. [87]	PP	Extended experimental and theoretical investigation of crystalline phase development during EFC and modelling.
1992	Barq et al. [88]	PET	Experimental and model results are compared, and influence of temperature is discussed.
1999	Acierno et al. [69]	PET; PP	Purely experimental study aimed on film temperature profiles and viscosity.
2000	Acierno et al. [90]	PET	Role of temperature profile on <i>NI</i> , minor importance of temperature if $X < 1/10L_0$ .
2000	Canning and Co [48]	LDPE; LLDPE	Purely experimental work deals with effect of rheology, <i>DR</i> and <i>MFR</i> on <i>NI</i> and <i>EB</i> .
2001	Canning et al. [70]	LDPE	Experimental work capturing velocity, width and thickness profile during EFC.
2001	Lamberti et al. [91]	iPP	Effect of processing cond. on film development including crystallization.
2001	Satoh et al. [39]	LDPE	Investigation of viscoelastic effects on <i>NI</i> and edge-beading, relates <i>NI</i> with <i>SH</i> in uni/pla extensional rate.
2002	Lamberti and Titomanlio [71]	iPP	Experimental investigation of EFC including width, velocity, temperature and crystallinity profiles.
2002	Lamberti et al. [92]	iPP	Experimental measuring and modelling of Hermans orientation factor and crystallinity. Evaluation of Ziabicki crystallization kinetics from measured film velocity, width and temperature profiles.
2002	Lamberti et al. [93]	iPP	Experimental measuring of film temperature profiles.
2002	Toft and Rigdahl [72]	LDPE; LLDPE; mLLDPE	Experimentally investigates the relationship between polymer elasticity and <i>NI</i> .
2003	Ito et al. [52]	LLDPE	Experimental oriented work using particle tracking. Confirmation of planar-uniaxial flows.
2003	Ito et al. [53]	LDPE; HDPE; LLDPE	Relates the <i>NI</i> extent to ratio of planar viscosities; axial to transverse.

2004	Uvieghara [68]	LLDPE	The experiment-oriented work, in which the effects of the Deborah number and the aspect ratio on the EFC were investigated.
2005 <sup>a</sup>	Agassant et al. [166]	-	Film casting review.
2005 <sup>a</sup>	Co [167]	-	Film casting review aimed on the draw resonance.
2005	Lamberti and Titomanlio [94]	iPP	Experimental part is accompanied by new cooling model with radiant heating.
2006	Aniunoh and Harrison [73]	PP	Effects of <i>DR</i> and die temperature on temperature, velocity and width profiles.
2006	Bourrigaud et al. [29]	LDPE	Effect of processing cond. on film development in coating process: divides <i>De-Dr</i> plane to attainable and unattainable regions.
2006	Lamberti and Titomanlio [95]	iPP	Effect of processing cond. on film development; film solidification within air gap.
2007	Aniunoh [107]	PP	Experimentally aimed study on how material properties and process conditions affect EFC.
2007	Kometani et al. [111]	PP; LDPE	Study aims on <i>NI</i> and <i>EB</i> investigation, utilized Giesekus equation was found to be the most suitable model describing the experimental data.
2007	Shin et al. [36]	LDPE; HDPE	Effects of temperature and extensional-thinning and -thickening.
2008	Kouda [74]	LDPE	Extrusion coating; linking neck-in degree with draw-down force.
2009	Seay and Baird [30]	LDPE; LLDPE; mLLDPE	Investigation of effects of <i>LCB</i> and <i>MWD</i> on <i>NI</i> via pom-pom model.
2009	McGrady et al. [65]	HDPE; LDPE	Effects of <i>LCB</i> and <i>MWD</i> on <i>NI</i> .
2010	Aniunoh and Harrison [66]	PP	Effects of <i>Mw</i> , <i>DR</i> and temperature on film formation
2010	Shiromoto et al. [109]	LDPE	Deals with extrusion lamination process. Relates the <i>NI</i> gauge to $\eta_{E,P}/\eta_{E,U}$ .
2010	Shiromoto et al. [54]	LDPE	Relates the <i>NI</i> gauge to $\eta_{E,P}/\eta_{E,U}$ .
2011	Lamberti [67]	iPP	Experimental Study designed to check capabilities of proposed <i>FIC</i> model.
2013	Pol et al. [41]	LDPE; HDPE; LLDPE	Effects of <i>LCB</i> and <i>MWD</i> on <i>NI</i> .
2014 <sup>a</sup>	Demay and Agassant [23]	-	Review targeting mainly transient instabilities during EFC.
2014	Pol et al. [42]	LDPE; HDPE; LLDPE	Effects of <i>LCB</i> on <i>NI</i> .
2014	Shiromoto [40]	LDPE	Effect of viscoelasticity on <i>NI</i> , relates the <i>NI</i> gauge to $\eta_{E,P}/\eta_{E,U}$ .
2015	Chikhalikar et al. [43]	PP	Effects of <i>LCB</i> on <i>NI</i> .
2015	Zhou et al. [96]	iPP	Effect of <i>DR</i> on crystallization and development of crystal morphology.
2016	Pol and Thete [98]	LDPE; LLDPE	Investigation of <i>NI</i> dependence on <i>De</i> and <i>DR</i> .
2020	Mu et al. [110]	PP	Influence of processing conditions on film geometry.

a) The state of art in EFC reviewing article

**Table 2.** Articles devoted to modelling the extrusion film casting process: 1D kinematic models. Unless otherwise stated, studies neglect the effects of flow induced crystallization, inertia, gravity, surface tension, aerodynamic drag, die swell, and film sag.

YEAR	REFERENCE	MATERIAL	EXPERIMENT	CONSTITUTIVE EQUATION	TRANSIENT SIMULATION	STRESS BOUNDARY CONDITION	NUMERICAL METHOD	TEMPERATURE	NOTE
1974	Yeow [80]	×	×	Newtonian	✓	×	Direct numerical scheme - 4 <sup>th</sup> order Runge-Kutta	×	First efforts of EFC modelling from its stability viewpoint.
1983	Aird and Yeow [81]	×	×	Generalized Newtonian (Power-law)	✓	×	?	×	Aims on investigation of stability EFC process.
1983	Minoshima and White [82]	×	×	Newtonian	✓	×	?	✓	Investigation of stability of EFC process by model incorporating thermal effects.
1988	Anturkar and Co [83]	×	×	Generalized UCM with deformation rate dependent $\eta$ and $\lambda$	✓	3	?	×	Stemming from (Yeow 1974) work, investigation of effects of viscoelasticity on the draw resonance phenomenon.
1990	Barq et al. [22]	PET	✓	Newtonian	✓	×	FDM	×	The work is aimed on transient phenomena of draw resonance.
1991 <sup>a</sup>	Alaie and Papanastasiou [45]	compared to PP	×	K-BKZ model with PSM damping function	×	0	FEM Galerkin	✓	Effects of die design and melt rheology, including temperature on film thickness in EFC process.
1993	Iyengar and Co [84]	×	×	mod. Giesekus with finite chain stretch	×	4	4 <sup>th</sup> order Runge-Kutta + adaptive step size control	×	Follows (Anturkar and Co 1988) work. Theoretical investigation of extensional viscosity curve and its effects on film velocity and stress profiles.
1994	Barq et al. [85]	compared to PET	×	UCM	✓	3	FDM	×	Theoretical investigation of limits of EFC process regarding to $DR_c$ .
1996	Iyengar and Co [75]	×	×	mod. Giesekus with finite chain stretch	✓	4	4 <sup>th</sup> order Runge-Kutta + adaptive step size control	×	Utilization of linear stability analysis to determination of $DR_c$ , examination of effects extensional thickening and thinning.

1996	Pis-Lopez and Co [76]	x	x	mod. Giesekus with finite chain stretch	x	4	4 <sup>th</sup> order Runge-Kutta + adaptive step size control	x	Steady-state analysis, multilayer film casting, study on the effect rheological properties and processing conditions on film velocity and stress profiles.
2000	S. Smith and Stolle [77]	x	x	Hooke's Law with creep	✓	x	Lagrangian FEM	x	Except few steady-state results, the work is focused on <i>DR</i> -determination viewed as a response problem.
2002	S. Smith and Stolle [78]	x	x	Hooke's Law with creep	✓	0	FEM + Newton Raphson	x	Comparison of Eulerian and Updated Lagrangian FE algorithms for film casting simulation.
2015	Polychronopoulos and Papathanasiou [26]	x	x	Newtonian; Linear PTT	x	0	OpenFOAM - FVM	x	Effect of draw ratio on die swell in film casting.
2020	Bechert [79]	x	x	Giesekus; exponential PTT; Generalized Newtonian (Carreau-Yasuda);	✓	4	AUTO-07P, MATLAB - CHEBFUN	x	Investigation of influence of viscoelastic, and in particular non-Newtonian, effects on the draw resonance instability in EFC.

a) Discrete relaxation spectra were used in this study.

*Legend to the stress boundary condition column on the die:*

**0** - all stress components are set to zero considering entire stress relaxation due to the die swell phenomenon.

**3** - the die exit stress state is given by the axial upstream extra stress component.

**4** - the thickness to axial extra stress component ratio for the upstream/down-stream side.

**Table 3.** Articles devoted to modelling the extrusion film casting process: 1.5D kinematic models. Unless otherwise stated, studies neglect the effects of flow induced crystallization, inertia, gravity, surface tension, aerodynamic drag, die swell, and film sag.

YEAR	REFERENCE	MATERIAL	EXPERIMENT	CONSTITUTIVE EQATION	MULTI-MODE APPROACH	STRESS BOUNDARY CONDITION	NUMERICAL METHOD	TEMPERATURE	CRYSTALLIZATION	NOTE
1989 <sup>a</sup>	Cotto et al. [18]	PP	✓	Newtonian	✗	✗	FDM	✓	✓	Experimental and theoretical investigation of crystalline phase development during EFC and modelling.
1990 <sup>a</sup>	Duffo et al. [100]	PP	✓	Newtonian	✗	✗	FDM	✓	✓	Effect of roll temperature on crystallization in EFC process.
1991 <sup>a</sup>	Agassant et al. [86]	✗	✗	Newtonian	✗	✗	FDM	✗	✗	Study deals with derivation of film width-variation model.
1991 <sup>a</sup>	Duffo et al. [87]	PP	✓	Newtonian	✗	✗	FDM	✓	✓	Extended experimental and theoretical investigation of crystalline phase development during EFC and modelling.
1992 <sup>ac</sup>	Barq et al. [88]	PET	✓	Newtonian	✗	✗	Runge-Kutta and Adams-Bashforth's method	✓	✗	Experimental and model results are compared, and influence of temperature is discussed.
1996 <sup>b</sup>	Silagy et al. [27]	✗	✗	Newtonian; UCM	✗	1	4 <sup>nd</sup> order Runge-Kutta; linear stability analysis	✗	✗	Brings afterward widely utilized 1.5D model based on (Narayanaswamy 1977) kinematics. Study of unattainable zone.
1999	Beaulne and Mitsoulis [89]	compared to PP; PET; LDPE;	✗	Newtonian; UCM; K-BKZ (PSM) model	✓	1	4 <sup>nd</sup> order Runge-Kutta; FEM;	✓	✗	Simulation result comparison with many related studies.
1999	Kwon [25]	Typical values for PE	✗	Leonov	✓	5	?	✗	✗	Qualitative simulation of anisotropic die swelling behavior of the extruded film
2000 <sup>a</sup>	Acierno et al. [90]	PET	✓	Newtonian	✗	✗	Ordinary single step Eulerian algorithm	✓	✗	Role of temperature profile on $Nl$ ; minor importance of temperature if $X < 1/10L_0$ .
2001	Lamberti et al. [91]	iPP	✓	Generalized Newtonian (Cross)	✗	✗	Euler's method	✓	✓	Effect of processing cond. on film development including crystallization.

2002	Lamberti et al. [93]	iPP	✓	Generalized Newtonian (Cross)	✗	✗	FDM	✓	✓	Experimental measuring of film temperature profiles.
2003	Ito et al. [53]	LDPE; HDPE; LLDPE	✓	Newtonian; UCM	✗	✗	Shooting method	✗	✗	Relates the <i>NI</i> extent to ratio of planar viscosities; axial to transverse.
2005	Lamberti and Titomanlio [94]	iPP	✓	UCM	✗	✗	✗	✓	✗	Experimental part is accompanied by new cooling model with radiant heating.
2005	Barot and Rao [99]	compared to iPP	✗	UCM	✗	✗	Variable order method	✓	✓	Modelling of crystallization during EFC process.
2006	Bourrigaud et al. [29]	LDPE	✓	UCM	✗	0	Num. integration+ Newton method	✗	✗	Effect of processing cond. on film development in coating process: divides <i>De-DR</i> plane to attainable and unattainable regions.
2013	Pol et al. [41]	LDPE; HDPE; LLDPE	✓	XPP; RP-S	✓	2	MATLAB - ode15s	✗	✗	Effects of <i>LCB</i> and <i>MWD</i> on <i>NI</i> .
2014	Pol et al. [42]	LDPE; HDPE; LLDPE	✓	XPP; RP-S	✓	2	MATLAB - ode15s	✓	✗	Effects of <i>LCB</i> on <i>NI</i> .
2015	Chikhalikar et al. [43]	PP	✓	XPP; RP-S	✓	1	MATLAB - ode15s	✓	✗	Effects of <i>LCB</i> on <i>NI</i> .
2015	Zhou et al. [96]	iPP	✓	Generalized Newtonian (Cross)	✗	✗	FDM/Monte Carlo	✓	✓	Effect of <i>DR</i> on crystallization and development of crystal morphology.
2016 <sup>bc</sup>	Bechert et al. [97]	✗	✗	Newtonian	✗	✗	AUTO-07P	✗	✗	Study reveals a stabilizing effect of <i>NI</i> , gravity and inertia on EFC process from draw resonance viewpoint.
2016	Pol and Thete [98]	LDPE; LLDPE	✓	UCM	✗	0	Finite difference scheme; time marching scheme	✗	✗	Investigation of <i>NI</i> dependence on <i>De</i> and <i>DR</i> .
2017	Barborik et al. [37]	compared to LDPE	✗	mod. Leonov	✗	5	4 <sup>th</sup> order Runge-Kutta	✗	✗	Investigation of the role of planar to uniaxial extensional viscosity ratio, extensional strain hardening and Deborah number.
2017 <sup>b</sup>	Dhadwal et al. [47]	✗	✗	exponential PTT	✓	1	MATLAB - ode15s; linear stability analysis	✗	✗	Effects of melt relaxation modes on process stability.
2017	Thete et al. [44]	LDPE; LLDPE	✓	UCM; exponential PTT	✓	2	MATLAB - ode15s	✗	✗	Effects of melt relaxation modes on <i>NI</i> .

2018	Barborik and Zatloukal [38]	compared to LDPE	✗	mod. Leonov	✗	5	4 <sup>th</sup> order Runge-Kutta	✗	✗	Study aimed on the effect of the second to first normal stress difference at the die exit and Deborah number.
2019	Barborik and Zatloukal [35]	compared to iPP	✗	mod. Leonov	✗	5	4 <sup>th</sup> order Runge-Kutta	✓	✓	Theoretical investigation of crystalline phase development during EFC.

- a) The model prediction capabilities are limited only to the final value of the film width  $L(X)$  to be in the range of  $L_0 > L(X) > L_0 - \sqrt{2}X$ .
- b) The study also includes a transient simulation.
- c) The effects of inertia and gravity are included in the model.

*Legend to the stress boundary condition column on the die:*

**0** - all stress components are set to zero considering entire stress relaxation due to the die swell phenomenon.

**1** - at least one stress component is given by the Newtonian solution for the downstream side, independent of the type of constitutive equation used.

**2** - two extra stress components are set manually without further justification.

**5** - ratio of the second to the first normal stress difference,  $-N_2/N_1$ , calculated from the upstream side by using a viscoelastic constitutive equation.

**Table 4.** Articles devoted to modelling the extrusion film casting process: 2D kinematic models. Unless otherwise stated, studies neglect the effects of surface tension, aerodynamic drag, die swell, and film sag.

YEAR	REFERENCE	MATERIAL	EXPERIMENT	MODEL DIMENSIONALITY	CONSTITUTIVE EQUATION	MULTI-MODE APPROACH	TRANSIENT SIMULATION	STRESS BOUNDARY CONDITION	NUMERICAL METHOD	TEMPERATURE	CRYSTALLIZATION	FLOW-INDUCED CRYSTALLIZATION	INERTIA	GRAVITY	NOTE
1990	D'Halewyu et al. [112]	x	x	2D	Newtonian	x	x	x	FEM; FVM	x	x	x	x	x	First 2D numerical model of EFC.
1995	Debbaut et al. [113]	x	x	2D	Newtonian; Generalized Newtonian (Power-law); UCM; Giesekus	x	x	0	FEM	x	x	x	x	x	Examining edge-beading in the light of viscoelastic flow behavior.
1997	W. S. Smith [12]	compared to PP	x	1D & 2D	Newtonian	x	x	x	Eulerian FEM	✓	x	x	✓	✓	Aims on effects of inertia, gravity and temperature on film formation; model comparison.
1998	Silagy et al. [28]	compared to LLDPE; LDPE	x	1.5D & 2D	Newtonian; UCM	x	✓	0, 1	2 <sup>nd</sup> order Runge-Kutta; FEM; linear stability analysis for 2D	x	x	x	x	x	Effect of <i>DR</i> , <i>A</i> and <i>De</i> on <i>NI</i> , comparison of models results; investigation of unattainable region.
1999	Silagy et al. [101]	compared to PES	x	1D & 2D	Newtonian	x	✓	x	FEM – Galerkin method; linear stability analysis for 1D and 2D	x	x	x	x	x	Effect of <i>DR</i> and <i>A</i> on <i>NI</i> , and onset of draw resonance.
2000	Christodoulou et al. [46]	PET	x	2D	exponential PTT	✓	✓	multi	FEM EVSS	x	x	x	x	x	Effect of stress boundary conditions on computational stability; determination of critical <i>DR</i> .



2000	S. Smith and Stolle [102]	Typical values for PET; LDPE	x	2D	Newtonian	x	x	x	FEM	✓	x	x	✓	✓	Investigation of factors that contribute <i>NI</i> reduction (self-weight, <i>EB</i> , cooling).
2001	Sato et al. [39]	LDPE	✓	2D	Newtonian; Larson	✓	x	1	Galerkin FEM	✓	x	x	x	x	Investigation of viscoelastic effects on <i>NI</i> and edge-beading, relates <i>NI</i> with <i>SH</i> in uni/pla extensional rate.
2001	W. S. Smith [8]	Typical values for PP	x	1.D; 1.5D & 2D	Generalized Newtonian (Power-Law); UCM	x	✓	0	UL FEM	✓	x	x	x	x	Model comparison – Eulerian vs Lagrangian approach.
2003	S. Smith and Stolle [103]	Typical values for PP	x	2D	Hooke's Law with creep	x	✓	0	UL FEM	✓	x	x	x	x	Based on Lagrangian description of motion, <i>DR<sub>c</sub></i> determined from response problem.
2003	Sollogoub et al. [104]	compared to PET	x	2D	Newtonian	x	x	x	Quadrangle continuous Galerkin FEM	✓	x	x	x	✓	Investigation of effects of <i>HTC</i> on film development.
2006	Kajiwara et al. [105]	x	x	2D	Newtonian; Generalized Newtonian (Cross); Larson; exponential PTT	x	x	1	Galerkin FEM	x	x	x	x	✓	Relates <i>NI</i> to ratio of uniaxial to planar extensional viscosity.
2006	Sollogoub et al. [106]	x	x	2D	UCM	x	x	0	Quadrangle continuous Galerkin FEM	✓	x	x	x	x	Effect of <i>HTC</i> and VE on film formation.
2007	K. Aniunoh [107]	PP	✓	2D	Giesekus	x	x	x	Matlab	✓	✓	✓	x	x	Experimentally aimed study on how material properties and process

															conditions affect EFC.
2007	Kometani et al. [111]	PP; LDPE	✓	2D	Newtonian; Generalized Newtonian (Bird Carreau); Giesekus	✗	✗	✗	Polyflow FEM	✗	✗	✗	✗	✗	Study aims on <i>NI</i> and <i>EB</i> investigation, utilized Giesekus equation was found to be the most suitable model describing the experimental data.
2007	Shin et al. [36]	LDPE; HDPE	✓	2D	exponential PTT	✗	✓	1	FEM ALE	✓	✗	✗	✗	✗	Effects of temperature and extensional-thinning and -thickening.
2010	Lee and Kim [108]	✗	✗	2D	exponential PTT	✗	✓	✗	Galerkin FEM	✗	✗	✗	✗	✗	Investigation of high aspect ratios causing the highly oriented molecular structures.
2010	Shiromoto et al. [109]	LDPE	✓	2D	exponential PTT	✗	✗	✗	ANSYS, Polyflow FEM	✓	✗	✗	✗	✗	Deals with extrusion lamination process. Relates the <i>NI</i> gauge to $\eta_{E,P}/\eta_{E,U}$ .
2010	Shiromoto et al. [54]	LDPE	✓	2D	exponential PTT	✗	✗	✗	ANSYS, Polyflow FEM	✓	✗	✗	✗	✗	Relates the <i>NI</i> gauge to $\eta_{E,P}/\eta_{E,U}$ .
2014	Shiromoto [40]	LDPE	✓	2D	Generalized Newtonian (Carreau-Yasuda); exponential PTT	✓	✗	✗	ANSYS, Polyflow FEM	✓	✗	✗	✗	✗	Effect of viscoelasticity on <i>NI</i> , relates the <i>NI</i> gauge to $\eta_{E,P}/\eta_{E,U}$ .
2020	Mu et al. [110]	PP	✓	2D	Generalized Newtonian (Carreau)	✗	✗	✗	FEM – with standard Galerkin formulation	✓	✗	✗	✗	✗	Influence of processing conditions on film geometry.

*Legend to the stress boundary condition column on the die:*

**0** - all stress components are set to zero considering entire stress relaxation due to the die swell phenomenon.

**1** - at least one stress component is given by the Newtonian solution for the downstream side, independent of the type of constitutive equation used.

**Table 5.** Articles devoted to modelling the extrusion film casting process: 3D kinematic models. Unless otherwise stated, studies neglect the effects of crystallization, flow induced crystallization, surface tension, aerodynamic drag, die swell, and film sag.

YEAR	REFERENCE	MATERIAL	EXPERIMENT	CONSTITUTIVE EQATION	STRESS BOUNDARY CONDITION	NUMERICAL METHOD	TEMPERATURE	INERTIA	GRAVITY	NOTE
1996	Sakaki et al. [114]	×	×	Newtonian	×	Streamline FEM	×	×	×	Effects of $DR$ , $X$ , $L_0$ on $NI$ and $EB$ .
2006	Zheng et al. [115]	compared to PET	×	Generalized Newtonian (Carreau)	×	FEM	✓	×	✓	Includes Shear-thinning, self-weight, viscous dissipation.
2007	Zheng et al. [116]	compared to PET	×	exponential PTT	1	Polyflow FEM	×	✓	✓	Effects of strain-hardening and elasticity on the final film shape.

*Legend to the stress boundary condition column on the die:*

**1** - at least one stress component is given by the Newtonian solution for the downstream side, independent of the type of constitutive equation used.

**Table 6.** Influence of material and extrusion film casting process variables on the extent of neck-in phenomenon (opposite trends are highlighted in bold).

YEAR	REFERENCE	MATERIAL	DEPENDENT VARIABLE	INDEPENDENT VARIABLE								
			Neck-in	Strain hardening in uniaxial extension	Mw, MWD ( $\lambda$ )	Draw ratio (adjusted via chill roll speed, $v_x(X)$ )	Draw ratio (adjusted via melt velocity at the die exit, $v_x(0)$ )	$v_x(X)$ and $v_x(0)$ (draw ratio is kept constant)	Take-up length	Temperature	$-N_2/N_1$ (if $De$ is high)	$\eta_{E,P}/\eta_{E,U}$
1986	Dobroth and Erwin [55]	LDPE	↑	—	—	—	↑	—	↑	—	—	—
1999	Acierno et al. [69]	PP, PET	↑	—	—	↑	—	—	↑	—	—	—
2000	Acierno et al. [90]	PET	↑	—	—	—	↑	—	↑	—	—	—
2000	Canning and Co [48]	LDPE, LLDPE	↑	↓	—	↓ <sup>c</sup> (LDPE) ↑ (LLDPE)	—	↓	—	—	—	—
2001	Canning et al. [70]	LDPE	↑	—	—	—/↑	—	—	—	—	—	—
2001	Satoh et al. [39]	LDPE	↑	↓	—	↑/∩ <sup>c</sup>	—	↓	—	—	—	—
2001	Lamberti et al. [91]	iPP	↑	—	—	↑	↓ <sup>a</sup>	—	—	—	—	—
2002	Lamberti et al. [93]	iPP	↑	—	—	—	↓ <sup>a</sup>	—	—	—	—	—
2002	Lamberti and Titomanlio [71]	iPP	↑	—	—	—	↓ <sup>a</sup>	—	—	—	—	—
2002	Lamberti et al. [92]	iPP	↑	—	—	—	↓ <sup>a</sup>	—	—	—	—	—
2002	Toft and Rigdahl [72]	LDPE, LLDPE, mLLDPE	↑	↓	↓	↓ <sup>b</sup>	—	—	↑	↑	—	—
2003	Ito et al. [53]	LDPE, LLDPE, HDPE	↑	—	—	↑ (LLDPE, HDPE) ↓ <sup>c</sup> (LDPE)	—	—	↑	—	—	—
2003	Ito et al. [52]	mLLDPE	↑	—	—	↑	—	—	—	—	—	—
2005	Seyfzadeh et al. [168]	PET	↑	—	—	↑	↑	—	—	—	—	—
2006	Aniunoh and Harrison [73]	PP	↑	—	—	↑	—	—	↑	↑	—	—
2006	Lamberti and Titomanlio [95]	iPP	↑	—	—	—	—	—	—	↑	—	—
2007	Aniunoh and Harrison [169]	PP	↑	—	↓	—	—	—	—	—	—	—
2007	Kometani et al. [111]	PP, LDPE	↑	—	—	↑ (PP) —/↑ <sup>c</sup> (LDPE)	—	—	—	—	—	—
2008	Kouda [74]	LDPE	↑	↓	—	—	—	—	—	—	—	—
2009	McGrady et al. [65]	HDPE	↑	↓	—	—	—	—	—	—	—	—
2009	Seay and Baird [30]	LDPE, LLDPE, mLLDPE	↑	↓	↓	↑	—	—	—	—	—	—
2010	Aniunoh and Harrison [66]	PP	↑	—	↓	↑	—	—	↑	↑	—	—

2010	Shiromoto et al. [109]	LDPE	↑	—	—	—	—	↓	↑	—	—	↑
2010	Shiromoto et al. [54]	LDPE	↑	↓	—	—	—	↓	↑	—	—	—
2013	Pol et al. [41]	LDPE, LLDPE, mLLDPE, HDPE	↑	↓	↓	↑	—	—	↑	—	—	—
2014	Pol et al. [42]	LDPE, LLDPE	↑	↓	↓	↑	—	—	—	—	—	—
2015	Chikhalikar et al. [43]	PP	↑	↓	↓	↑	—	—	—	—	—	—
2015	Zhou et al. [96]	iPP	↑	—	—	↑	—	—	—	—	—	—
2018	Barborik and Zatloukal [38]	LDPE	↑	—	—	↑	—	—	—	—	↑	↑
2020	Mu et al. [110]	PP	↑	—	—	↑	—	—	↑	—	—	—

- a) The observed opposite trend is due to the crystallization of the polymer in the drawing region.
- b) Used LLDPE and mLLDPE (octene/hexene comonomer types) have a high Trouton ratio ( $>20$ ), indicating the presence of *LCB*.
- c) The effect of the draw ratio on the neck-in is complex for LDPEs (most likely due to the non-monotonic dependence of the uniaxial and planar extensional viscosities on the extensional strain rate that is typical for branched polymers).

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## 7 LIST OF SYMBOLS

Latin Symbols	Meaning	Unit
A	Aspect ratio	1
a	Parameter in Cross and Carreau-Yasuda model	1
B	Bead ratio	1
b	Dissipation term in modified Leonov model	s <sup>-1</sup>
b <sub>c</sub>	Chain extensibility parameter in modified Giesekus model	1
$\underline{\underline{c}}$	Recoverable Finger tensor	1
$\underline{\underline{C}}^{-1}$	Finger strain tensor	1
$\overset{\circ}{\underline{\underline{c}}}$	Jaumann (corotational) time derivative of the recoverable Finger strain tensor in modified Leonov model	s <sup>-1</sup>
D	Deformation rate tensor	s <sup>-1</sup>
De	Deborah number	1
DR	Draw ratio	1
DR <sub>c</sub>	Critical draw ratio	1
$\underline{\underline{E}}$	Elasticity matrix in Hooke's Law	Pa
e	Half-thickness of the film at any $x$ location	m
e <sub>0</sub>	Die half-gap (half-thickness of the film at the die exit)	m

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$\underline{\underline{e}}_p$	Irreversible rate of strain tensor in modified Leonov model	$s^{-1}$
$f(x)$	Rate of deformation in transverse y-direction	$s^{-1}$
$G$	Linear Hookean elastic modulus	Pa
$G'$	Storage modulus	Pa
$G''$	Loss modulus	Pa
$g(x)$	Rate of deformation in thickness z-direction	$s^{-1}$
HTC	Heat transfer coefficient	$J \cdot s^{-1} \cdot K^{-1} \cdot m^{-2}$
$h_f^{edge}$	Edge final film thickness	mm
$h_f^{center}$	Center final film thickness	mm
$\underline{\underline{\Pi}}_D$	Second invariant of deformation rate tensor	$s^{-1}$
$\underline{\underline{I}}_c$	First invariant of recoverable Finger tensor	1
$\underline{\underline{\Pi}}_c$	Second invariant of recoverable Finger tensor	1
$\underline{\underline{I}}_{C^{-1}}$	First invariant of the Finger strain tensor	1
$\underline{\underline{\Pi}}_{C^{-1}}$	Second invariant of the Finger strain tensor	1
$i$	Index $i$ , noting the spatial direction	1
$J_E^0$	Linear steady-state elastic compliance	$Pa^{-1}$
$j$	Relaxation mode identification number	1
$L$	Half-width of the film at any $x$ location	m
$\underline{\underline{L}}_v$	Velocity gradient tensor	$s^{-1}$

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$L_0$	Half-width of the die (half-width of the film at the die exit)	m
$m$	Flow consistency index in Power-law model	$\text{Pa}\cdot\text{s}^n$
MFR , $\dot{m}$	Mass flow rate	$\text{kg}\cdot\text{h}^{-1}$
$M_n$	Number average molar mass	$\text{g}\cdot\text{mol}^{-1}$
$M_w$	Mass average molar mass	$\text{g}\cdot\text{mol}^{-1}$
$N$	Presents the highest value available	–
$n$	Flow behavior index in GNM	1
$n_L$	Non-linear Leonov model parameter	1
$n'$	Adjustable parameter in relaxation time function in Generalized UCM model	1
$NI$	Maximum attainable neck-in	m
$N_1$	First normal stress difference	Pa
$N_2$	Second normal stress difference	Pa
$-N_2/N_1$	Stress state at the die exit	1
$q$	Number of arms in the XPP model	1
$T$	Melt temperature	$^{\circ}\text{C}$
$t$	Present time in K-BKZ model	s
$t'$	Past time in K-BKZ model	s
$v_x$	Axial velocity component of the film at any $x$ location	$\text{m}\cdot\text{s}^{-1}$



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$v_x(0)$	Axial velocity component at the die exit (velocity in the machine direction)	$\text{m}\cdot\text{s}^{-1}$
$v_x(X)$	Chill roll speed (Take-up rate)	$\text{m}\cdot\text{s}^{-1}$
$v_y$	Velocity component of the film in transverse y-direction at any $x$ location	$\text{m}\cdot\text{s}^{-1}$
$v_z$	Velocity component of the film in thickness z-direction at any $x$ location	$\text{m}\cdot\text{s}^{-1}$
$W$	Elastic potential in modified Leonov model	Pa
$X$	Take-up length (drawing distance, air gap)	m
$x, y, z$	Spatial coordinates in axial, transverse and thickness direction, respectively	m
$x$	Position in axial x-direction	m
$Z$	Function in modified Giesekus model	1
<b>Greek Symbols Meaning</b>		<b>Unit</b>
$\alpha$	Anisotropy parameter in Giesekus and XPP model	1
$\alpha_i$	Parameter in K-BKZ model	1
$\beta$	Non-linear Leonov model parameter	1
$\beta_c$	Convective constraint release coefficient in RP-S model	1
$\beta_i$	Parameter in K-BKZ model	1

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$\dot{\gamma}$	Shear strain rate	$s^{-1}$
$\Delta \underline{\underline{\varepsilon}}$	Total strain increment in Hooke's Law	1
$\Delta \underline{\underline{\varepsilon}}^c$	Modified Perzyna creep strain increment in Hooke's Law	1
$\Delta \underline{\underline{\sigma}}$	Stress increment in Hooke's Law	Pa
$\underline{\underline{\delta}}$	The Kronecker delta, unit tensor	1
$\delta_0$	Fitting scalar parameter in RP-S model	1
$\varepsilon_p$	Parameter in PTT model	1
$\dot{\varepsilon}_p$	Extensional strain rate	$s^{-1}$
$\eta, \eta_s$	Steady shear viscosity	Pa·s
$\eta_0$	Newtonian viscosity, zero-shear viscosity	Pa·s
$\eta_b$	Steady biaxial extensional viscosity	Pa·s
$\eta_p$	Polymer viscosity in Giesekus model	Pa·s
$\eta_\infty$	Infinite-shear-rate viscosity, solvent viscosity in Giesekus model	Pa·s
$\eta_{E,P}, \eta_P$	Steady planar extensional viscosity	Pa·s
$\eta_{E,U}, \eta_U$	Steady uniaxial extensional viscosity	Pa·s
$\eta_{E,U,max}$	Maximal steady uniaxial extensional viscosity	Pa·s
$\frac{\eta_{E,U,max}}{3\eta_0}$	Uniaxial extensional strain hardening	1

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$\theta$	Parameter in K-BKZ model	1
$\Lambda$	Backbone tube stretch in XPP model	1
$\lambda, \lambda_1$	Melt relaxation time	s
$\bar{\lambda}$	Average relaxation time	s
$\lambda_0$	Adjustable parameter in relaxation time function in Generalized UCM model	s
$\lambda_{0b}$	Orientation relaxation time in XPP model	s
$\lambda_{0s}$	Stretch relaxation time in XPP model	s
$\lambda_d$	Reptation relaxation time in RP-S model	s
$\lambda_r$	Rouse relaxation time in RP-S model	s
$\lambda_t$	Adjustable parameter in relaxation time function in Generalized UCM model	s
$\underline{\underline{\lambda(\tau)^{-1}}}$	relaxation time tensor in XPP model	s
$\nu$	Non-linear Leonov model parameter	1
$\xi$	Non-linear Leonov model parameter	1
$\xi_p$	Parameter in PTT model	1
$\zeta$	Non-linear model parameter in Larson model	1
$\underline{\underline{\tau}}$	Extra stress tensor	Pa
$\underline{\underline{\diamond \tau}}$	Gordon–Schowalter convected time derivative of the stress tensor in PTT model	Pa·s <sup>-1</sup>

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$\nabla$	Upper-convected time derivative of the stress tensor	$\text{Pa}\cdot\text{s}^{-1}$
$\tau_{=p}$	Polymer contribution to the stress tensor in Giesekus model	Pa
$\tau_{=s}$	Solvent contribution to the stress tensor in Giesekus model	Pa
$\tau_{xx}$	Normal stress in axial x-direction	Pa
$\bar{\tau}_{xx}$	Dimensionless normal stress in axial x-direction	1
$\tau_{yy}$	Normal stress in transverse y-direction	Pa
$\bar{\tau}_{yy}$	Dimensionless normal stress in transverse y-direction	1
$\tau_{zz}$	Normal stress in thickness z-direction	Pa
$\bar{\tau}_{zz}$	Dimensionless normal stress in thickness z-direction	1
$\nabla$	Gradient operator	1

#### Latin Abbreviations and Acronyms

Unit

1D, 1.5D, 2D and 3D	Model dimensionality, e.g. 1D – One-dimensional model	–
CHEBFUN	Framework within the MATLAB	–
DE	Doi-Edwards integral tube model	–
EB	Edge-beading phenomenon	–

EFC	Extrusion film casting	–
FDM	Finite difference method	–
FE	Finite element	–
FEM	Finite element method	–
FEM ALE	FEM the arbitrary Lagrangian Eulerian method	–
FEM EVSS	FEM the elastic viscous stress splitting	–
FIC	Flow-induced crystallization	–
FVM	Finite volume method	–
GNM	Generalized Newtonian model	–
HDPE	Material: high-density polyethylene	–
iPP, PP	Material: isotactic polypropylene	–
K – BKZ	Kaye-Bernstein-Kearsley-Zapas constitutive model	–
LCB	Long chain branching	–
LDPE	Material: low-density polyethylene	–
LLDPE	Material: linear low-density polyethylene	–
MWD	Molecular weight distribution	–
mLLDPE	Material: linear metallocene-catalyzed low-density polyethylene	–
ode15s	Differential equation solver within MATLAB software	–
PBAT	Material: poly(3-hydroxybutyrate-co-3-hydroxyvalerate)	–

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PBS	Material: polybutylene succinate	—
PES	Material: polyethersulfone	—
PET	Material: polyethylene terephthalate	—
PLA	Material: polylactide	—
PP	Material: polypropylene	—
PS	Material: polystyrene	—
PSM	Papanastasiou-Scriven-Macosko damping function for K-BKZ constitutive model	—
PTT	Phan–Thien and Tanner constitutive model	—
RP–S	Rolie-Poly Strech constitutive model	—
SH	Strain hardening	—
$\text{tr}(\ )$	Denotes the trace of a matrix	—
UCM	Upper convected Maxwell constitutive model	—
UL FEM	Updated Lagrangian FEM	—
VE	Viscoelasticity	—
XPP	eXtended Pom–Pom constitutive model	—

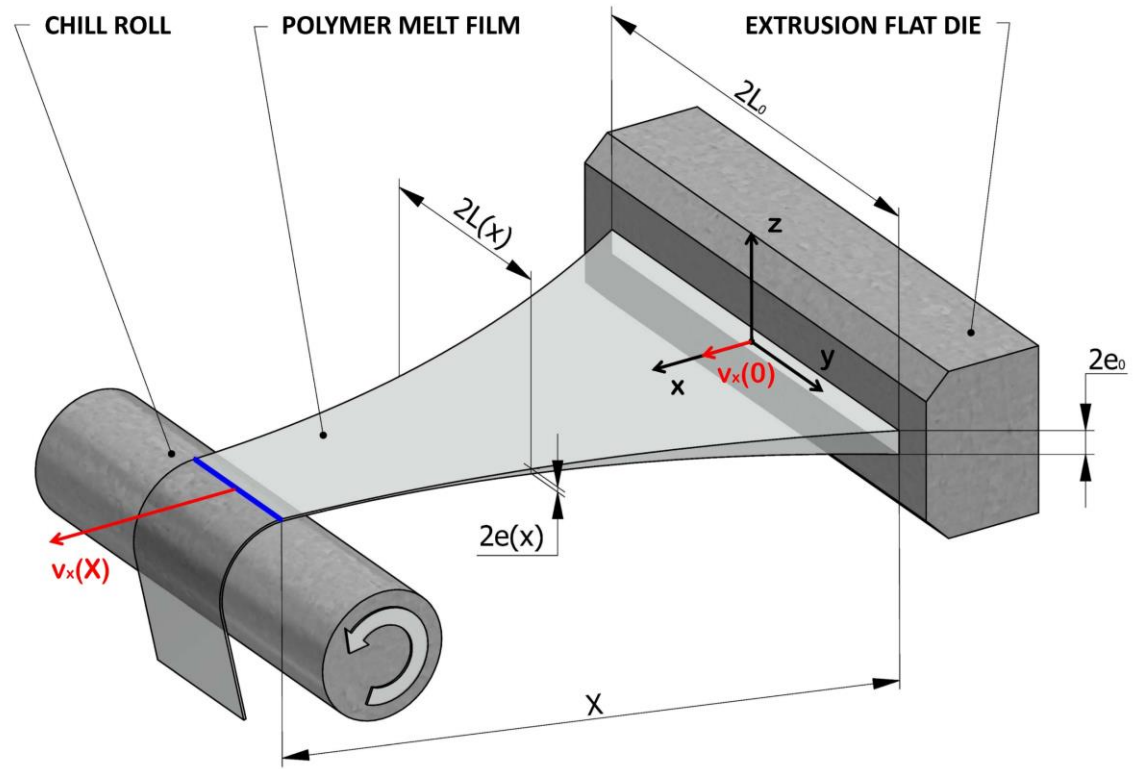


Fig. 1: Schematics of the extrusion film casting kinematics. Reproduced from [T. Barborik, and M. Zatloukal, in *AIP Conference Proceedings* 1843, 030010 (American Institute of Physics Inc., Zlin, Czech Republic July 26-27, 2017., 2017)], [170], with the permission of AIP Publishing.

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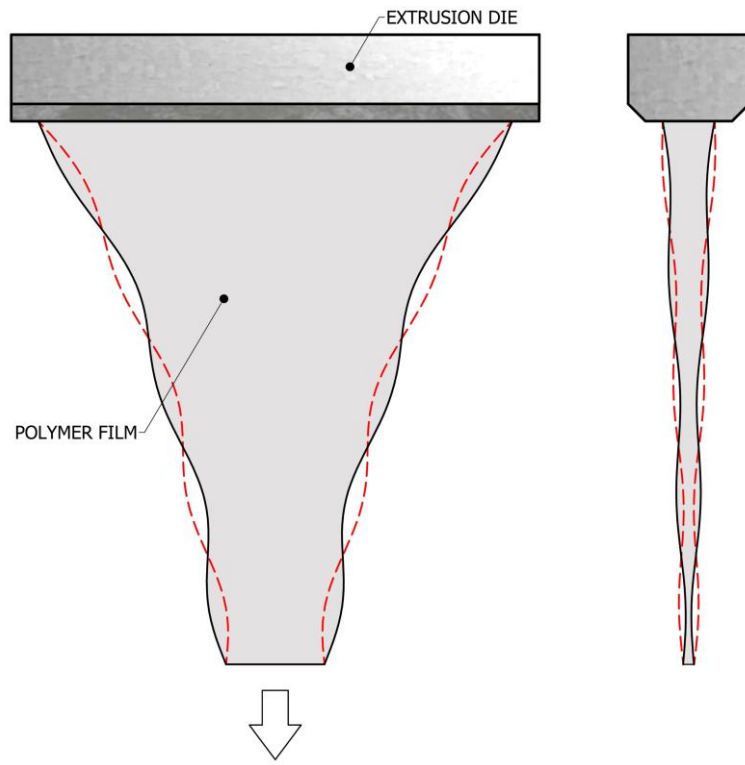


Fig. 2: Visualization of the effect of draw resonance on film width and thickness.



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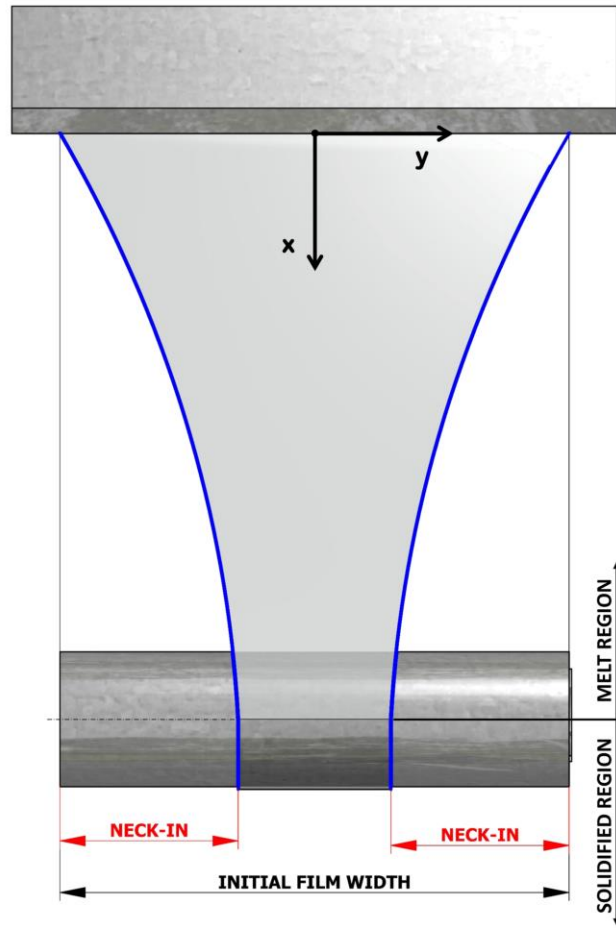


Fig. 3: Visualization of the neck-in phenomenon during extrusion film casting.  
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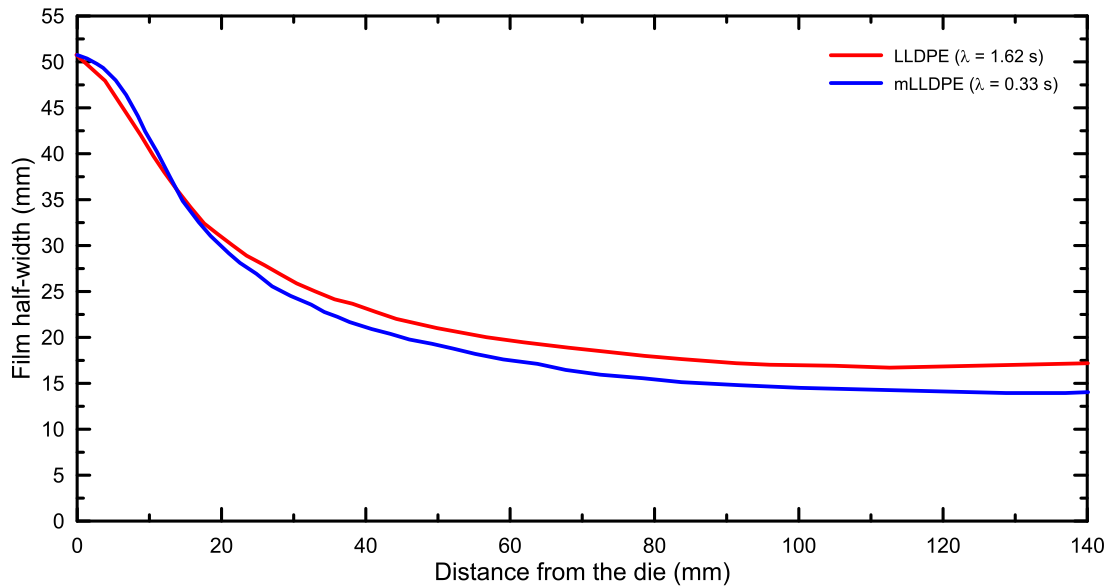


Fig. 4: Effect of Carreau relaxation time due to increased MWD on the neck-in for linear metallocene-catalyzed low-density polyethylene (mLLDPE,  $\lambda = 0.33$  s at  $150$  °C,  $M_w = 110\,000$  g/mol,  $M_w/M_n = 2.04$ ,  $LCB/10\,000\,C=0$ ) and Ziegler-Natta-catalyzed linear low-density polyethylene LLDPE ( $\lambda = 1.62$  s,  $M_w = 122\,700$  g/mol,  $M_w/M_n = 3.44$ ,  $LCB/10\,000\,C=0$ ). The process conditions for both samples were following: die width was  $101.6$  mm and its gap size was  $0.57$  mm, take-up length equal to  $141.4$  mm, a temperature of  $150$  °C, an extrusion shear rate of  $8.62\text{ s}^{-1}$  and a drawdown ratio of 15. The flow direction is oriented left to right here. Selected and digitalized from [30].

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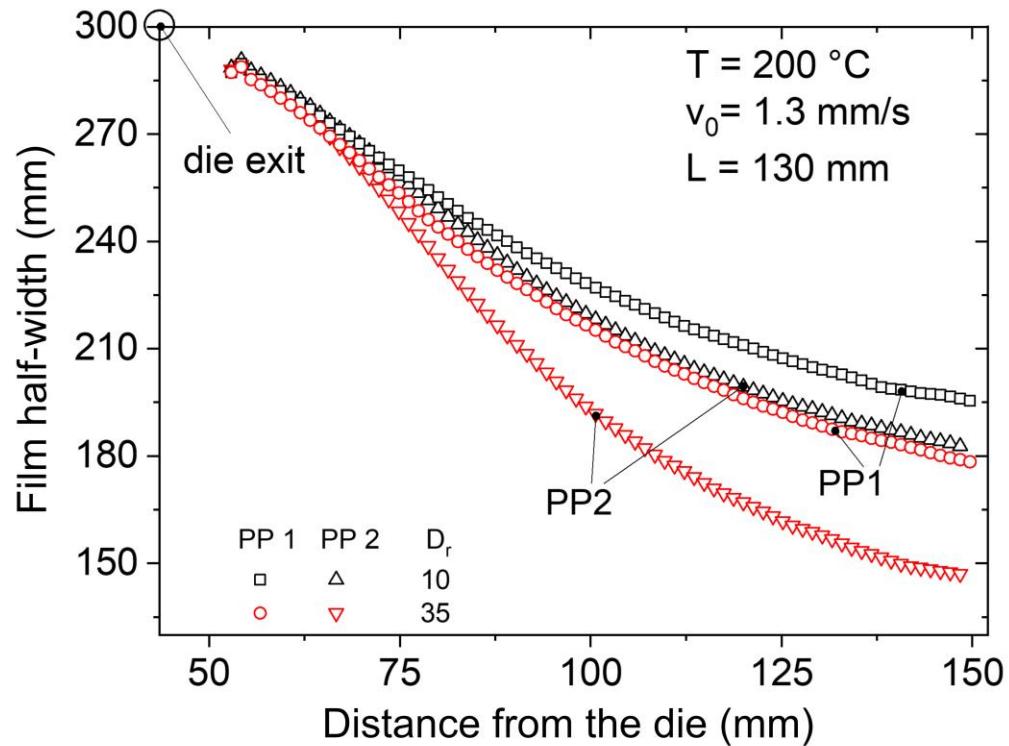


Fig. 5: Effect of the longest relaxation time (i.e.  $\lambda = \eta_0 J_E^0$ , where  $J_E^0$  is the linear steady-state elastic compliance) due to the increased molecular weight on the neck-in for linear polypropylenes PP1 ( $\lambda = 15.5$  s at 230 °C,  $M_w = 527\,000$  g/mol,  $M_w/M_n = 5.3$ ) and PP2 ( $\lambda = 4.2$  s at 230 °C,  $M_w = 359\,000$  g/mol,  $M_w/M_n = 6.9$ ) at two draw down ratios (10 and 35). The process conditions for both samples were as follows: the velocity at the die exit was 1.3 mm/s, the take-up length was 130 mm, the temperature was 200 °C [31–34]. Here, the flow direction is oriented from left to right. Reproduced from [H. Münstedt, in AIP Conference Proceedings (eds. Zatloukal, M. & Musil, J.) 2107, 030001 (American Institute of Physics Inc., Zlin, Czech Republic, 2019)], [34], with the permission of AIP Publishing.

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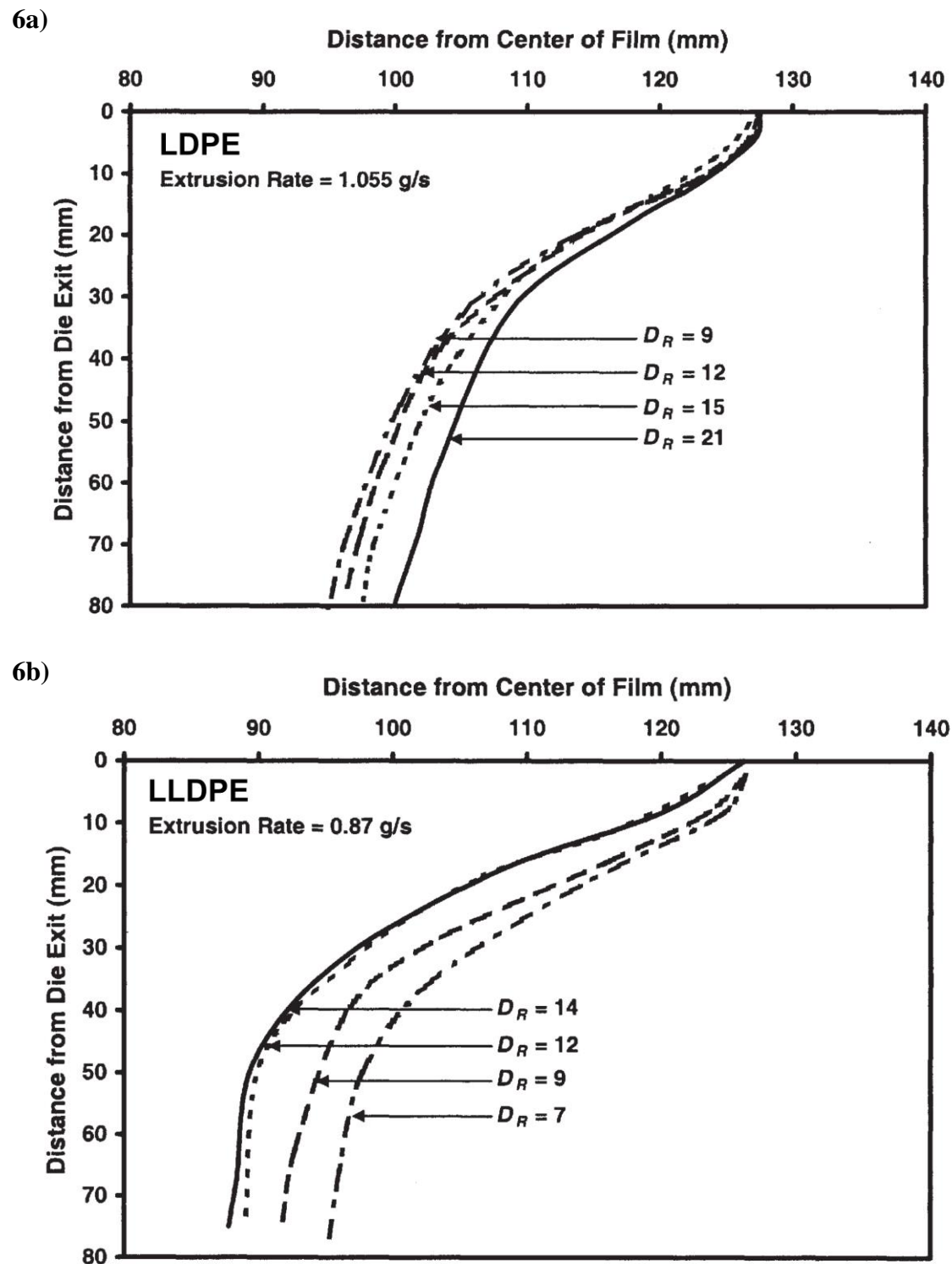
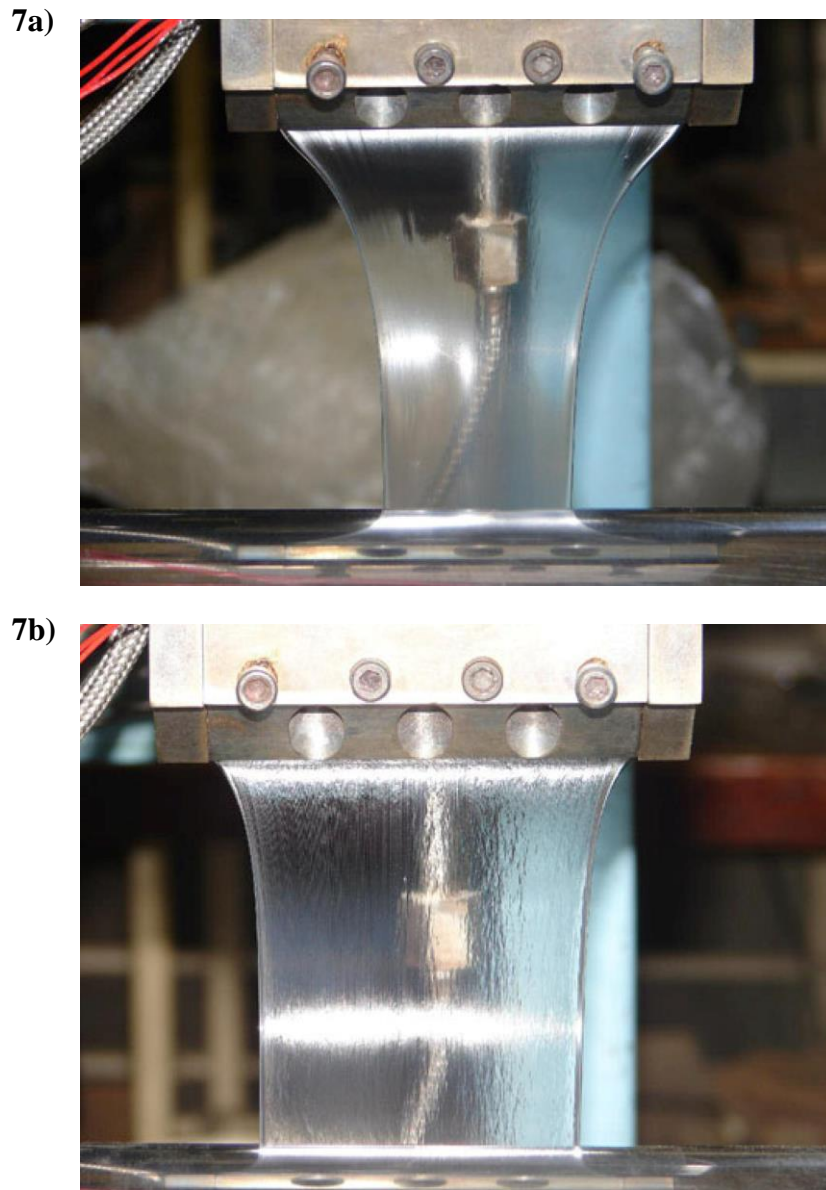


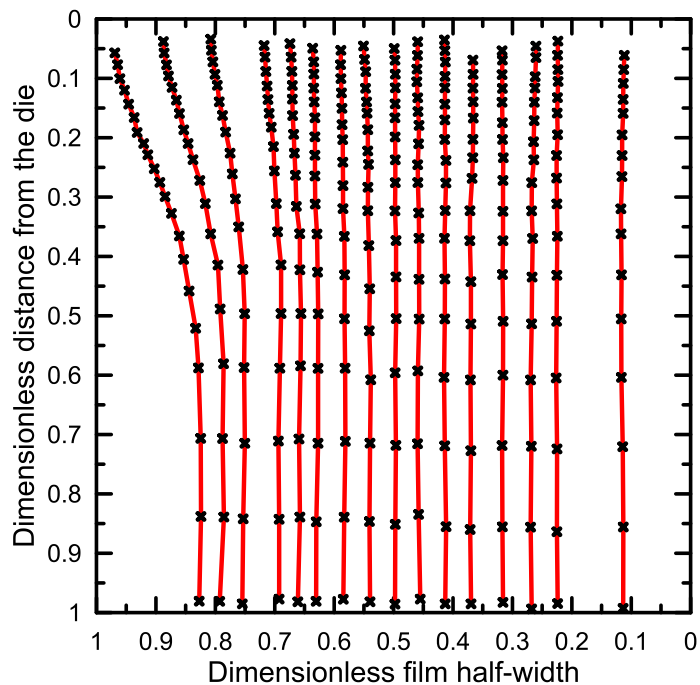
Fig. 6: Effect of long chain branching on the neck-in development at different draw ratios, experimental data: 6a) LDPE with LCB; 6b) LLDPE without LCB. The process conditions for both samples were following: the die width was 254 mm and the temperature 240 °C. Here the flow direction is oriented from top to bottom. Reproduced with permission from *J. Plast. Film Sheeting* 16 (3), 188 (2000) [48]. Copyright 2000, SAGE Publications.

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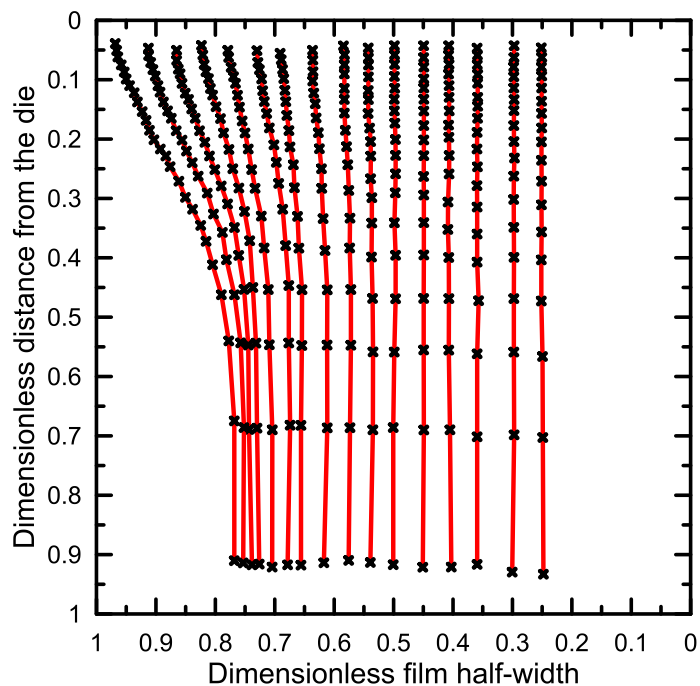


*Fig. 7: Effect of increased long chain branching on the neck-in: 7a) linear metallocene-catalyzed low-density polyethylene (mLLDPE); 7b) highly branched low-density polyethylene (LDPE). The process conditions for both samples were as follows: the die width was 101.6 mm and its gap size 0.57 mm, the take-up length 141.4 mm, the temperature 150 °C, the extrusion shear rate  $1.33 \text{ s}^{-1}$  and the drawdown ratio 10. Here, the flow direction is oriented from top to bottom. Courtesy of professor Donald G. Baird for his permission to reprint this figure from [171].*

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DR = 4.4



DR = 12.2

Fig. 8: Visualization of the polymer flow field in the air gap during film casting of linear metallocene-catalyzed low-density polyethylene (mLLDPE,  $M_w = 57\,200$  g/mol,  $M_w/M_n = 2.26$ ), in which the flow direction is from top to bottom. The process conditions were as follows: the velocity at the die exit was 10.7 mm/s, the take-up length equal to 150 mm, the temperature 190 °C. Selected and digitalized from [52].

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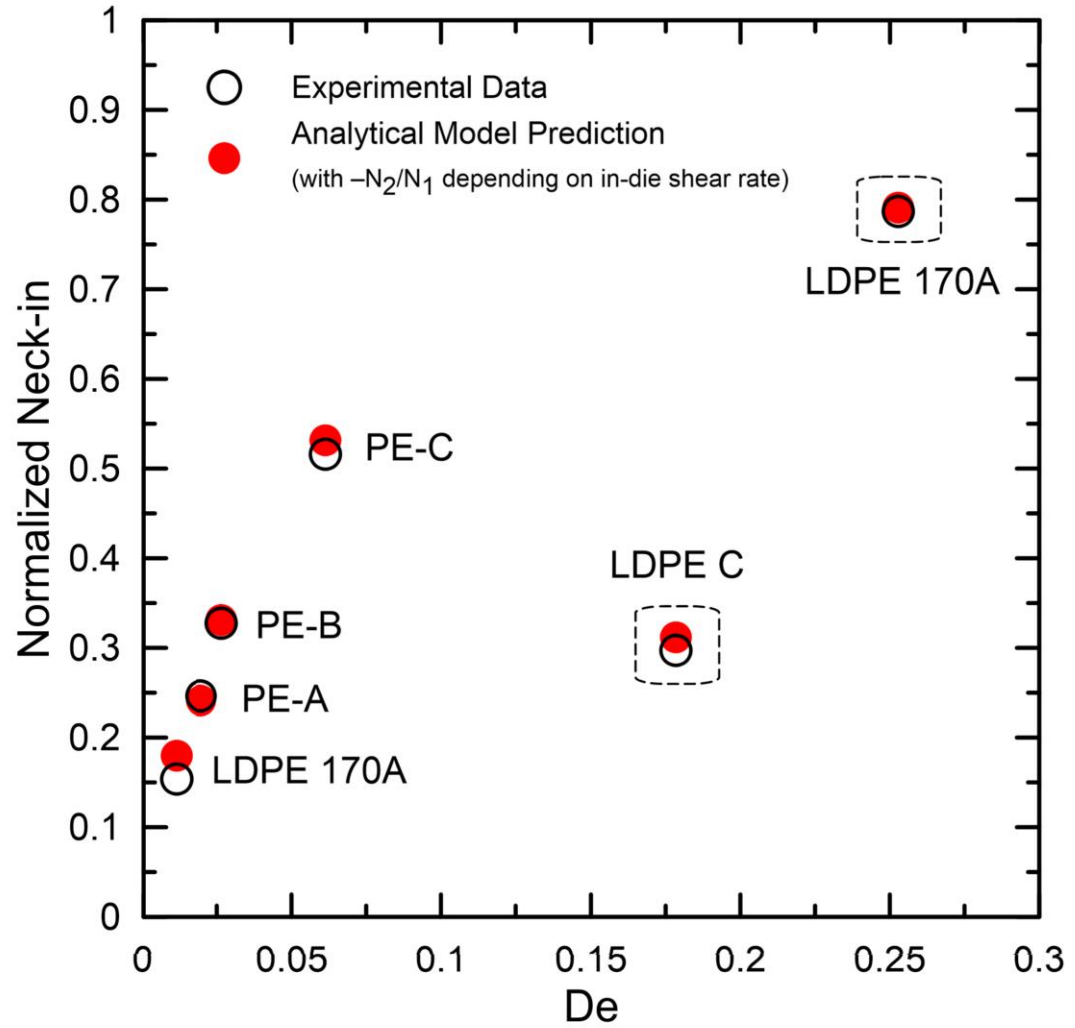
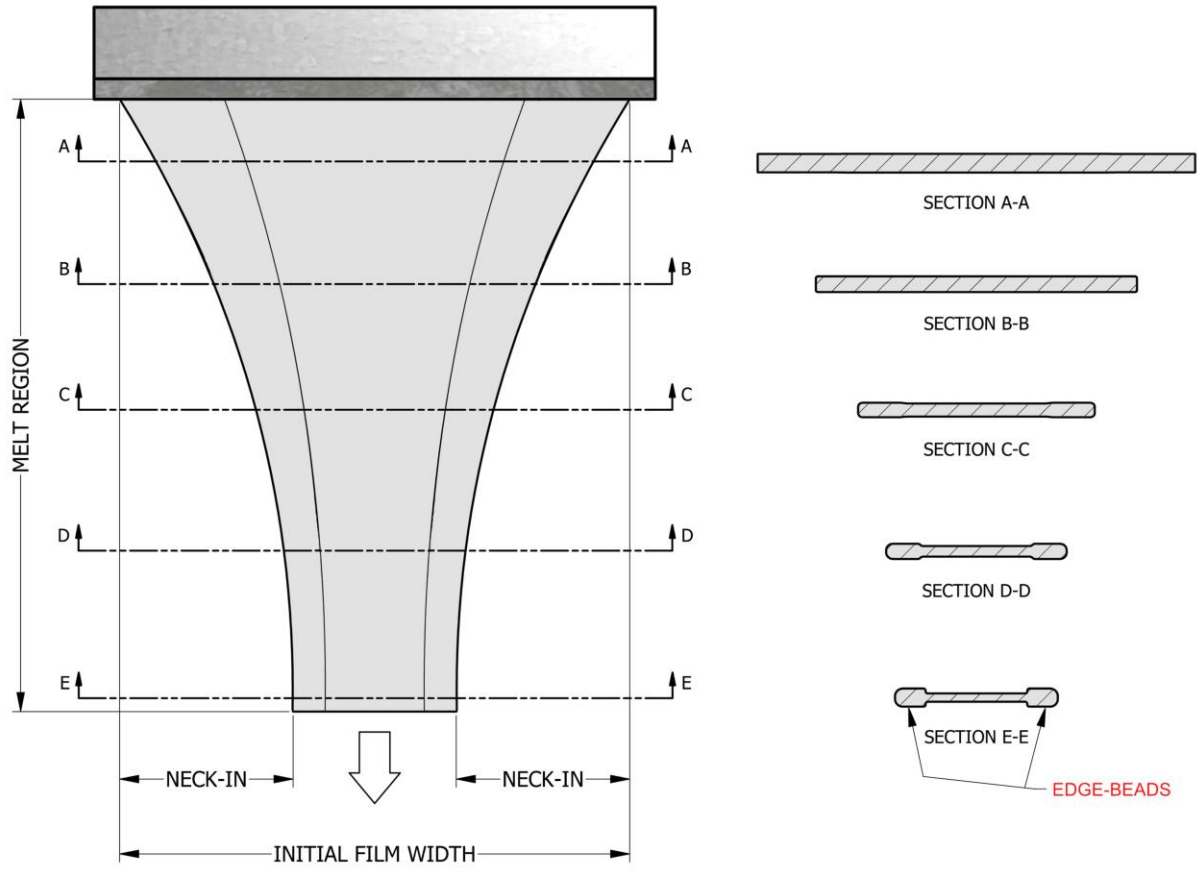


Fig. 9: Maximum attainable normalized neck-in (i.e.  $NI/X$ ) for different LDPEs. Experimental data and proposed analytical model predictions (Eq. 6) are presented here with open and filled symbols, respectively. Reproduced with permission from J. Nonnewton. *Fluid Mech.* 255, 39 (2018) [38]. Copyright 2018, Elsevier.



*Fig. 10: Schematic illustration of the extrusion film casting process with the indicated film cross-sectional development (formation of edge-beads) in the air-gap. The curves within the film represent the borders between the planar extensional flow (center area) and the uniaxial extensional flow (edge areas).*

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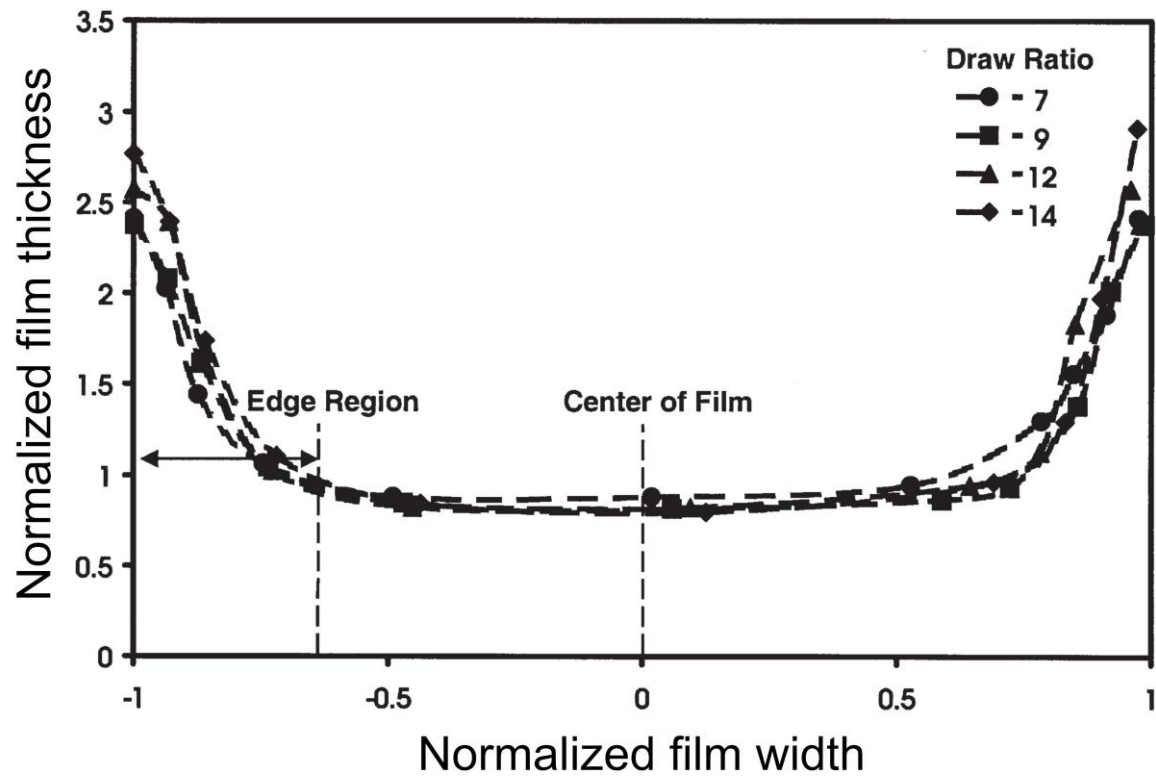


Fig. 11: Evolution of edge-beading at different draw ratios, experimental data for LLDPE without LCB (corresponding film width profiles are provided in Figure 6b). Process conditions were following: the die width was 254 mm and the temperature 240 °C. Reproduced with permission from *J. Plast. Film Sheeting* 16 (3), 188 (2000) [48]. Copyright 2000, SAGE Publications.

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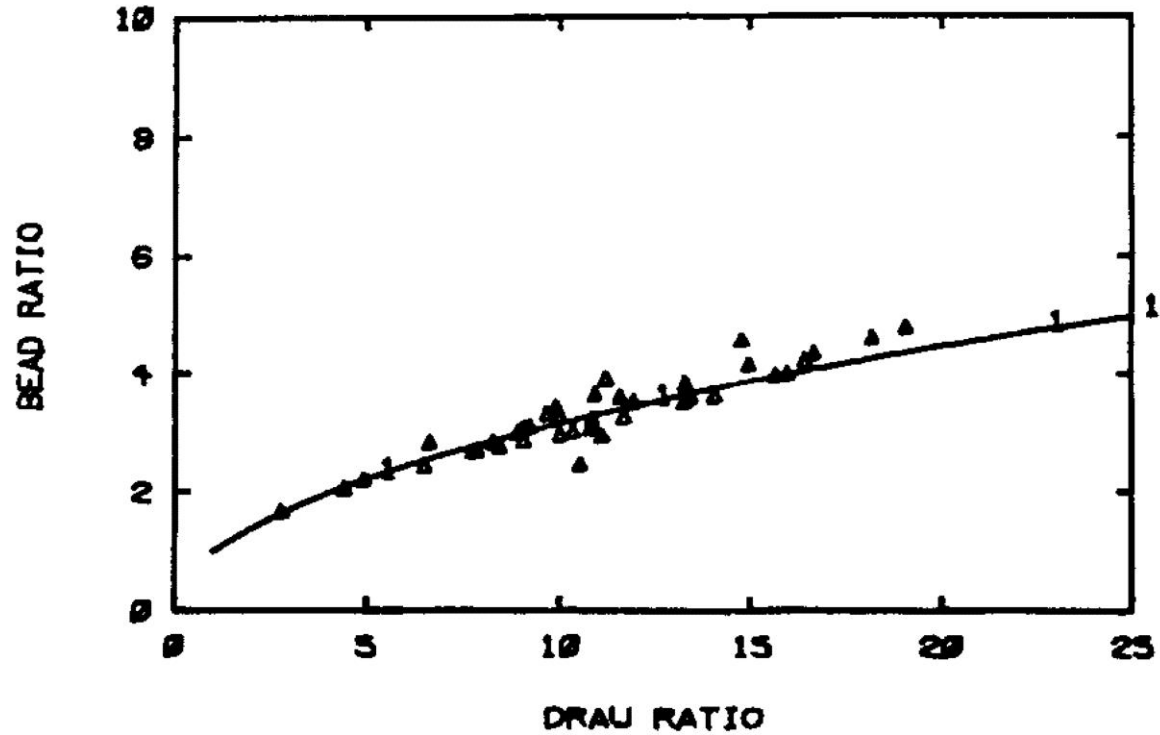


Fig. 12: Bead ratio as a function of DR for LDPE at 177 °C. The symbols represent experimental data, while the line represents the theoretical value predicted by the Eq. 7. Reproduced with permission from *Polym. Eng. Sci.* 26 (7), 462 (1986) [55]. Copyright 1986, John Wiley and Sons.

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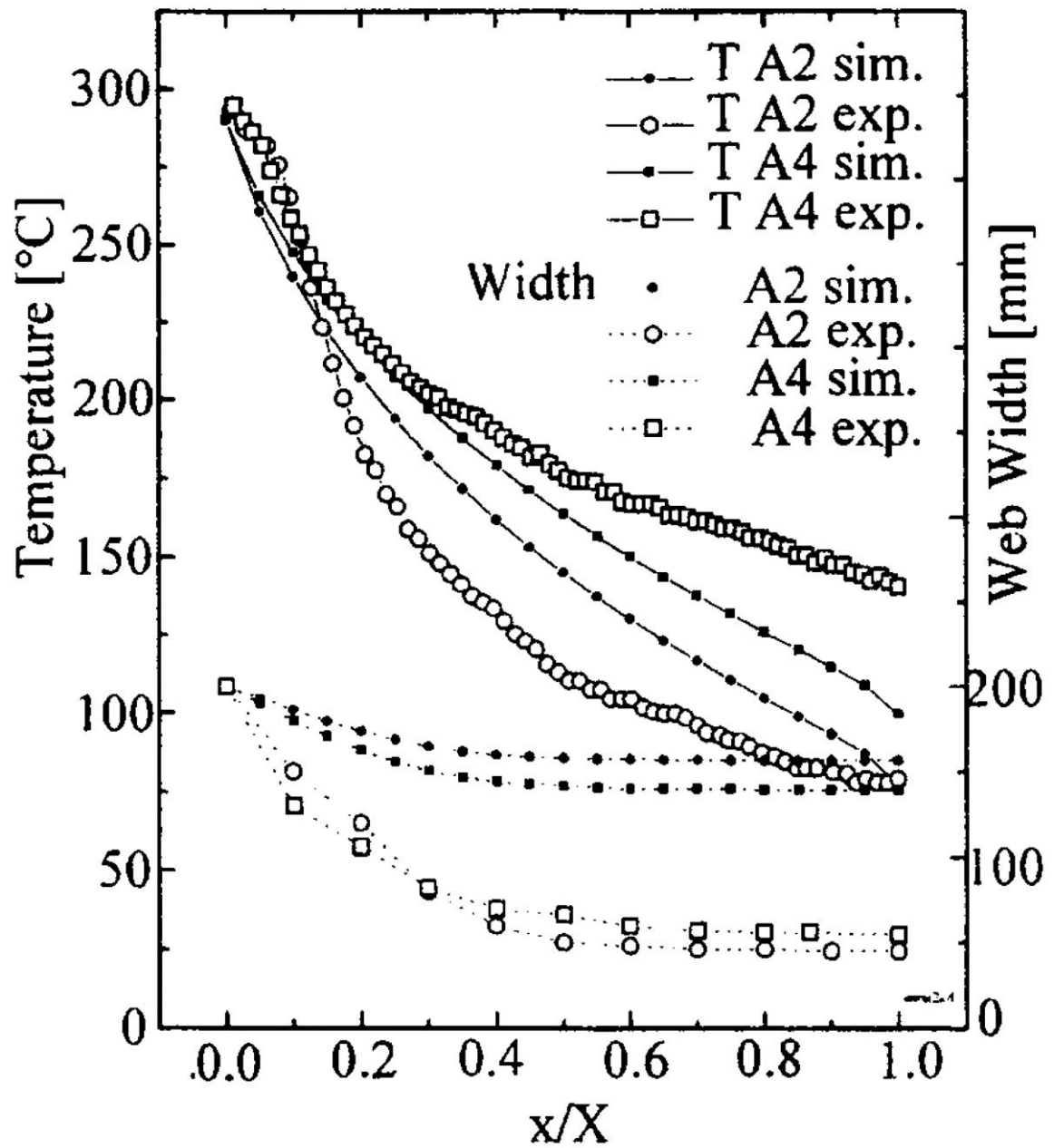


Fig. 13: Film temperature and width against dimensionless drawing distance for PET for DR = 10 (A2 experiment) and DR = 20 (A4 experiment), die width = 200 mm and  $X = 150$  mm. The simulations are based on the Newtonian constitutive equation and 1.5D kinematic model. Reproduced with permission from *Polym. Eng. Sci.* 40 (1), 108 (2000) [90]. Copyright 2000, John Wiley and Sons.

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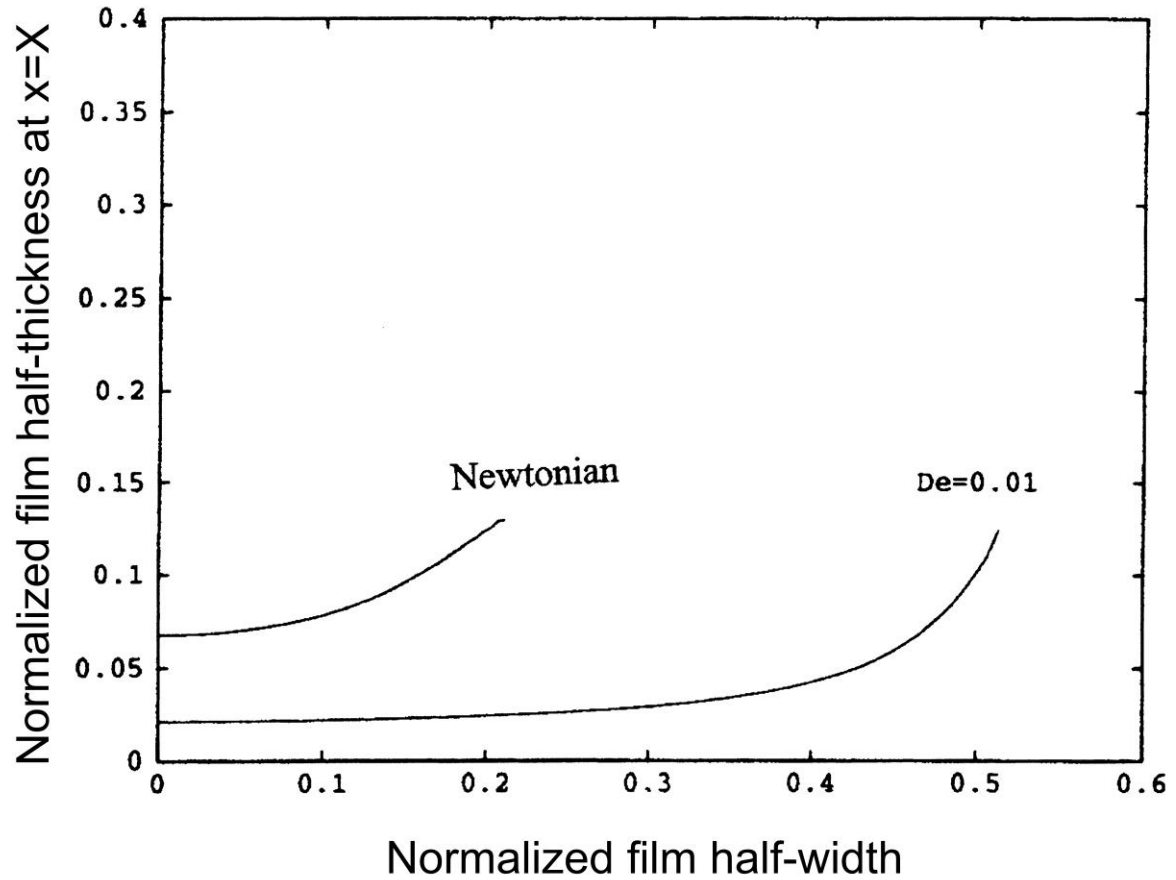


Fig. 14: Effect of elasticity on the edge-beading for  $DR = 50$  predicted by the 2D film casting model considering Newtonian and UCM constitutive equations. The vertical axis represents the dimensionless film thickness (actual thickness divided by the die gap) and the horizontal axis represents the dimensionless film width (actual distance from the film center divided by the die width). Reproduced with permission from J. Nonnewton. *Fluid Mech.* 79 (2–3), 563 (1998) [28]. Copyright 1998, Elsevier.

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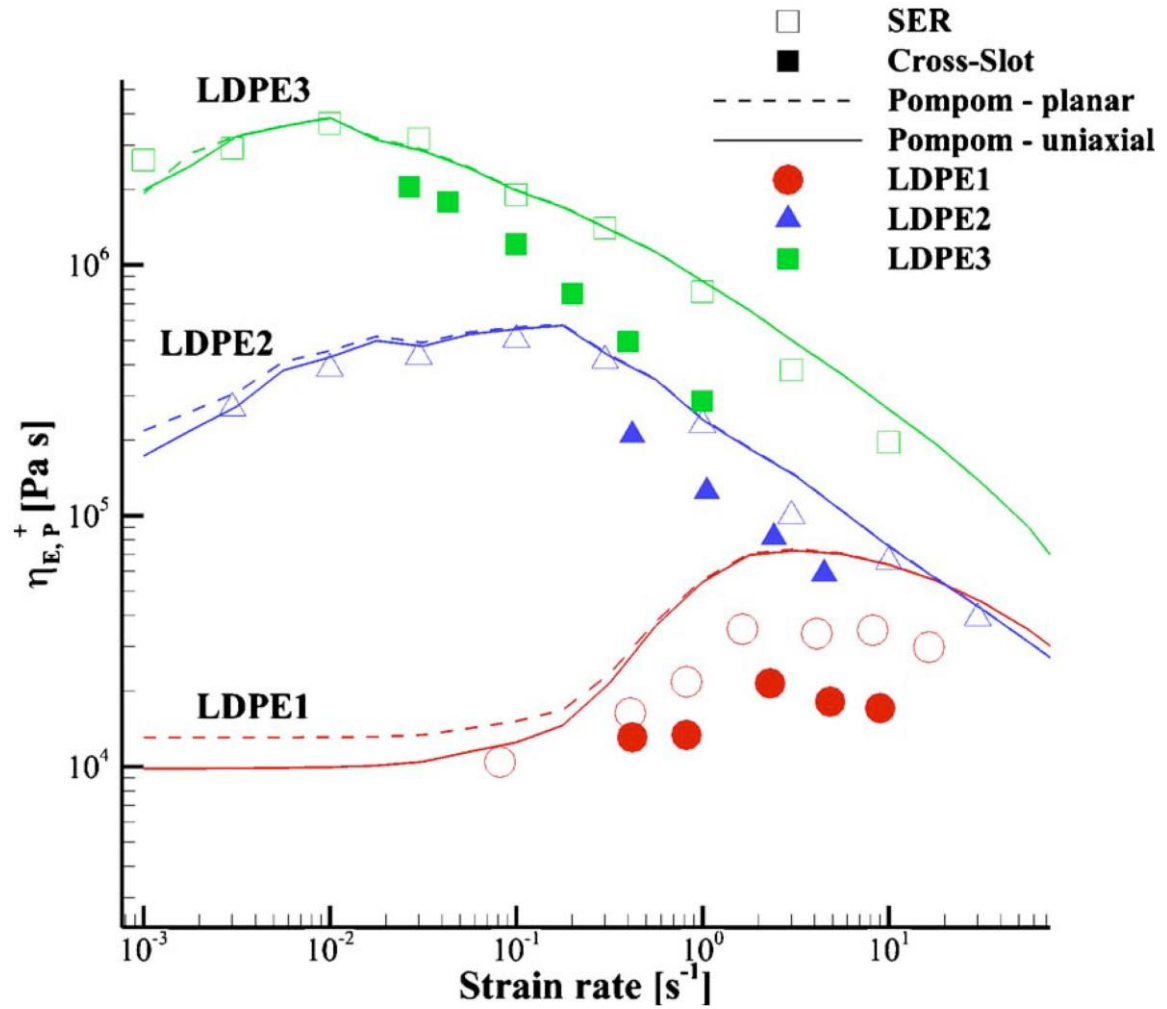


Fig. 15: Comparison between experimentally measured uniaxial (open symbols) and planar (closed symbols) extensional viscosities together with the multimode Pom-Pom model predictions for the uniaxial (solid lines) and planar (dashed lines) extensional viscosity for the LDPE series. Reproduced from [D. Auhl, D. M. Hoyle, D. Hassell, T. D. Lord, O. G. Harlen, M. R. Mackley, and T. C. B. McLeish, “Cross-slot extensional rheometry and the steady-state extensional response of long chain branched polymer melts,” *J. Rheol.* 55 (4), 875 (2011)], [127], with the permission of AIP Publishing.

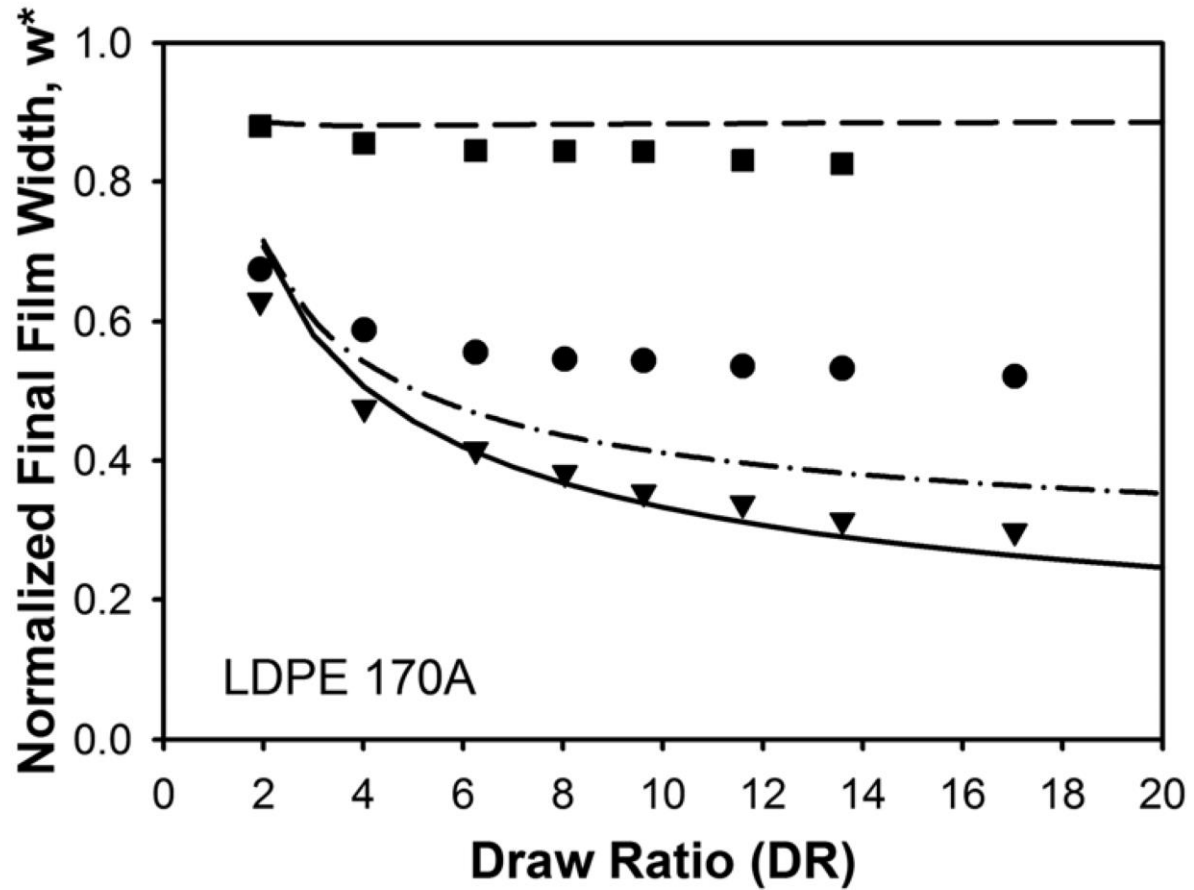


Fig. 16: Normalized film width vs. draw ratio for branched LDPE at three different air-gaps (squares:  $X = 10\text{mm}$ ; circles:  $X = 90\text{mm}$ ; inverted triangles:  $X = 228\text{mm}$ ; die width =  $100\text{ mm}$ ). The symbols represent experimental data and the lines are predictions based on the XPP constitutive equation and the 1.5D kinematic model. Reproduced from [H. V. Pol, S. S. Thete, P. Doshi, and A. K. Lele, "Necking in extrusion film casting: The role of macromolecular architecture," *J. Rheol.* 57 (2), 559 (2013)], [41], with the permission of AIP Publishing.

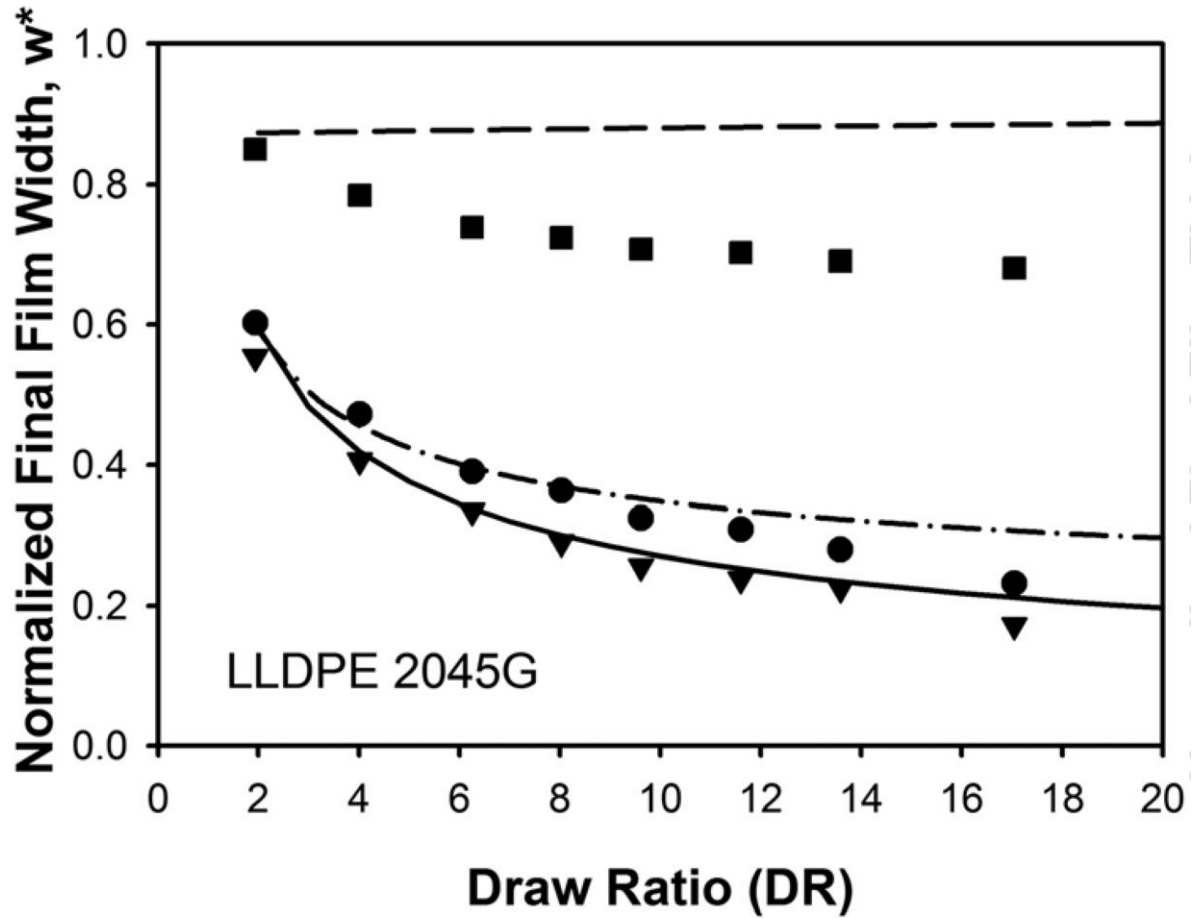
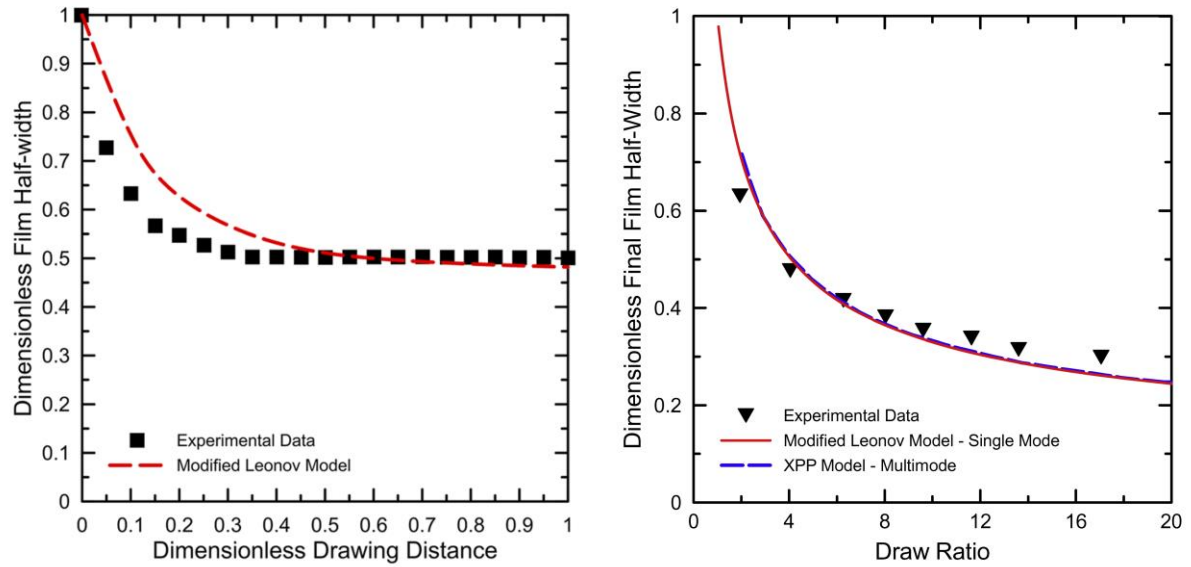


Fig. 17: Normalized film width vs. draw ratio for linear LLDPE at three different air-gaps (squares:  $X = 10$  mm; circles:  $X = 90$  mm; inverted triangles:  $X = 228$  mm; die width = 100 mm). The symbols represent experimental data and the lines are predictions based on the RP-S constitutive equation and the 1.5D kinematic model. Reproduced from [H. V. Pol, S. S. Thete, P. Doshi, and A. K. Lele, "Necking in extrusion film casting: The role of macromolecular architecture," *J. Rheol.* 57 (2), 559 (2013)], [41], with the permission of AIP Publishing.

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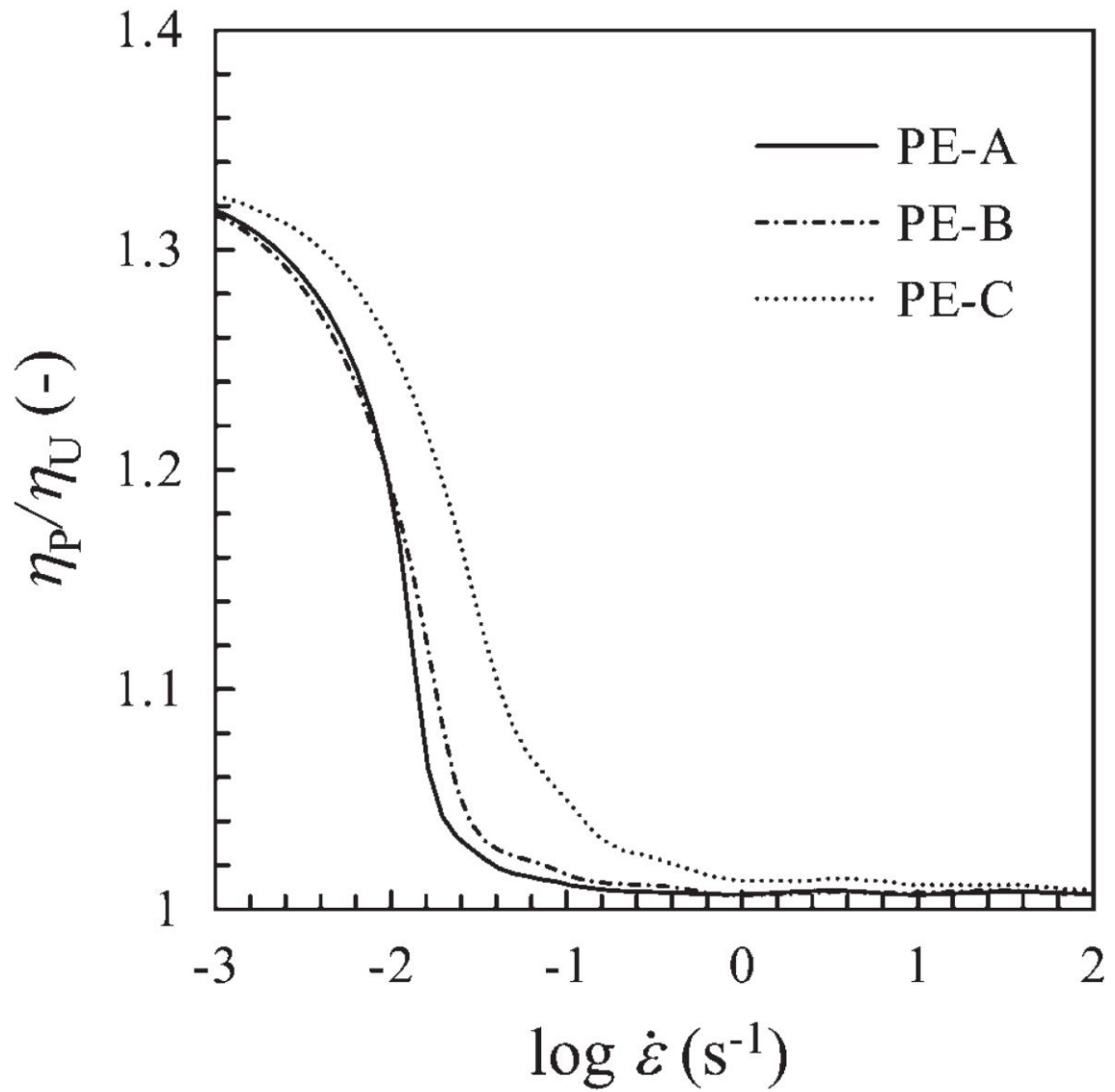
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*Fig. 18: Normalized film width vs. drawing distance at  $DR = 34.7$ , die width = 200 mm and  $X = 400$  mm for linear PP (left) and normalized film width vs. DR at die width = 100 mm at  $X = 230$  mm for branched LDPE (right). The symbols represent experimental data and the lines are predictions based on the modified Leonov constitutive equation and the 1.5D kinematic model. Left figure reproduced from [T. Barborik, and M. Zatloukal, “Effect of Heat Transfer Coefficient, Draw Ratio and Die Exit Temperature on the Production of Flat Polypropylene Membranes,” *Phys. Fluids* 31 (5), 053101 (2019)], [35], with the permission of AIP Publishing. Right figure reproduced with permission from *Int. J. Heat Mass Transf.* 111, 1296 (2017) [37]. Copyright 2017, Elsevier.*



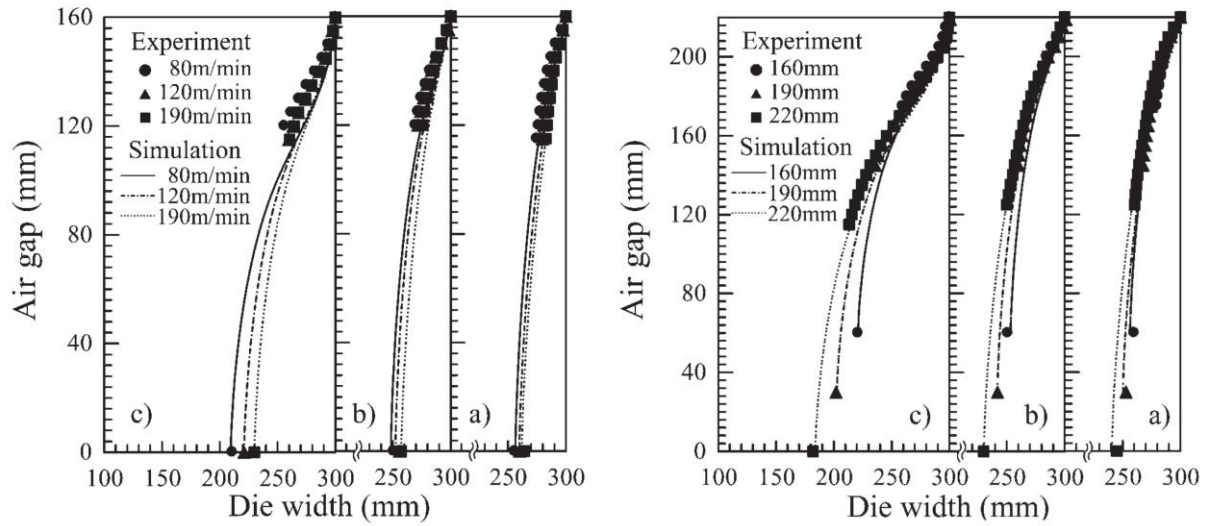
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*Fig. 19: The planar to uniaxial extensional viscosity ratio predicted by the PTT model for three different LDPEs plotted as a function of the extensional strain rate. Reproduced with permission from Polym. Eng. Sci. 50 (1), 22 (2010) [109]. Copyright 2010, John Wiley and Sons.*

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*Fig. 20: Comparison of the film edge shapes for three LDPEs with different uniaxial extensional strain hardening (a-high, b-middle, c-low uniaxial extensional strain hardening) where die width = 600 mm, and DR was kept the same, equal to 40 by proper adjusting of  $v_x(0)$  and  $v_x(X)$ . Left: the role of take-up velocities (80, 120 and 190 m/min),  $X = 160$  mm; Right: the role of air-gaps (160, 190 and 220 mm) at the fixed take-up velocity (120 m/min). The symbols represent experimental data and the lines are predictions based on the multimode PTT constitutive equation and the 2D kinematic model. Reproduced with permission from Polym. Eng. Sci. 50 (1), 22 (2010) [109]. Copyright 2010, John Wiley and Sons.*

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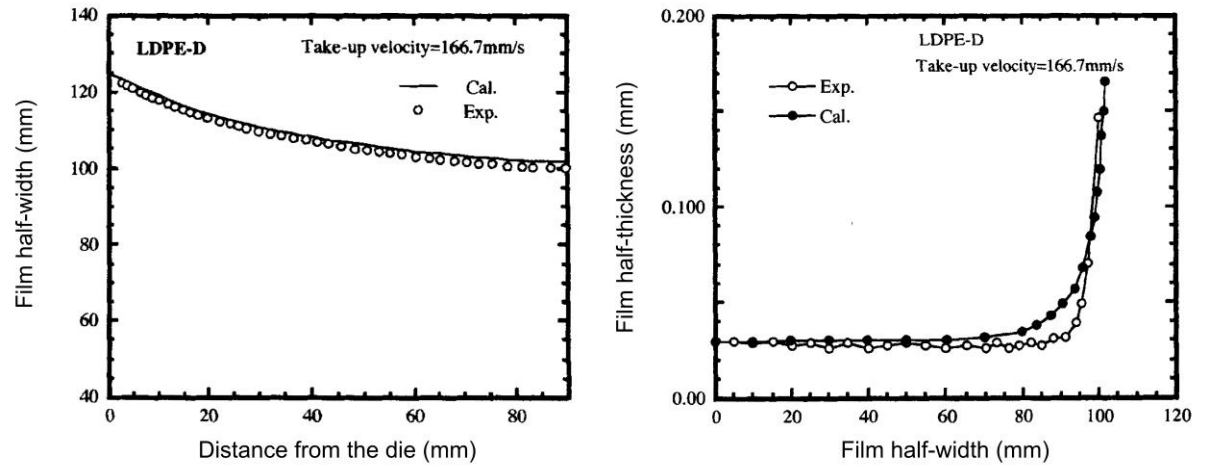


Fig. 21: Film width vs. drawing distance (left) and final film thickness across the film width (right) at  $DR = 32.7$ , die width = 0.25 m and  $X = 0.09$  m for branched LDPE (right). Open circles represent experimental data and lines (or lines with filled circles) are predictions based on the Larson constitutive equation and the 2D kinematic model. Reproduced with permission from *Polym. Eng. Sci.* 41 (9), 1564 (2001) [39]. Copyright 2001, John Wiley and Sons.

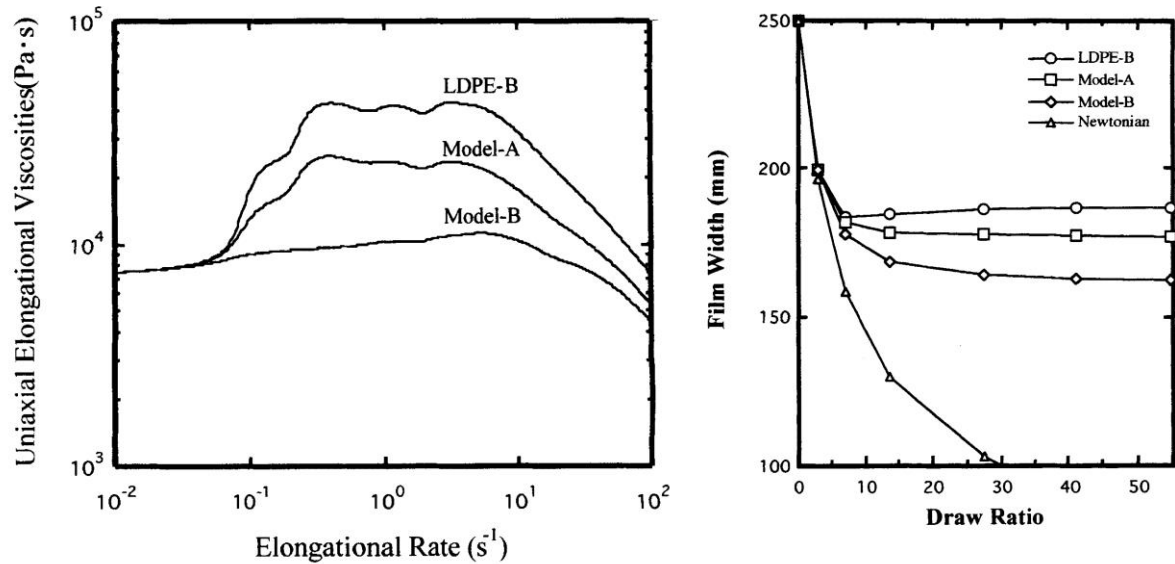


Fig. 22: Uniaxial extensional viscosity vs. extensional strain rate predicted by the multimode PTT constitutive equation for LDPE-B: high; Model-A: middle; Model-B: low uniaxial strain hardening (left) and related predictions for film width vs. DR including comparison with corresponding Newtonian predictions by using the 2D kinematic model (right), where  $DR = 32.7$ , die width = 250 mm and  $X = 90$  mm. Reproduced with permission from *Polym. Eng. Sci.* 41 (9), 1564 (2001) [39]. Copyright 2001, John Wiley and Sons.

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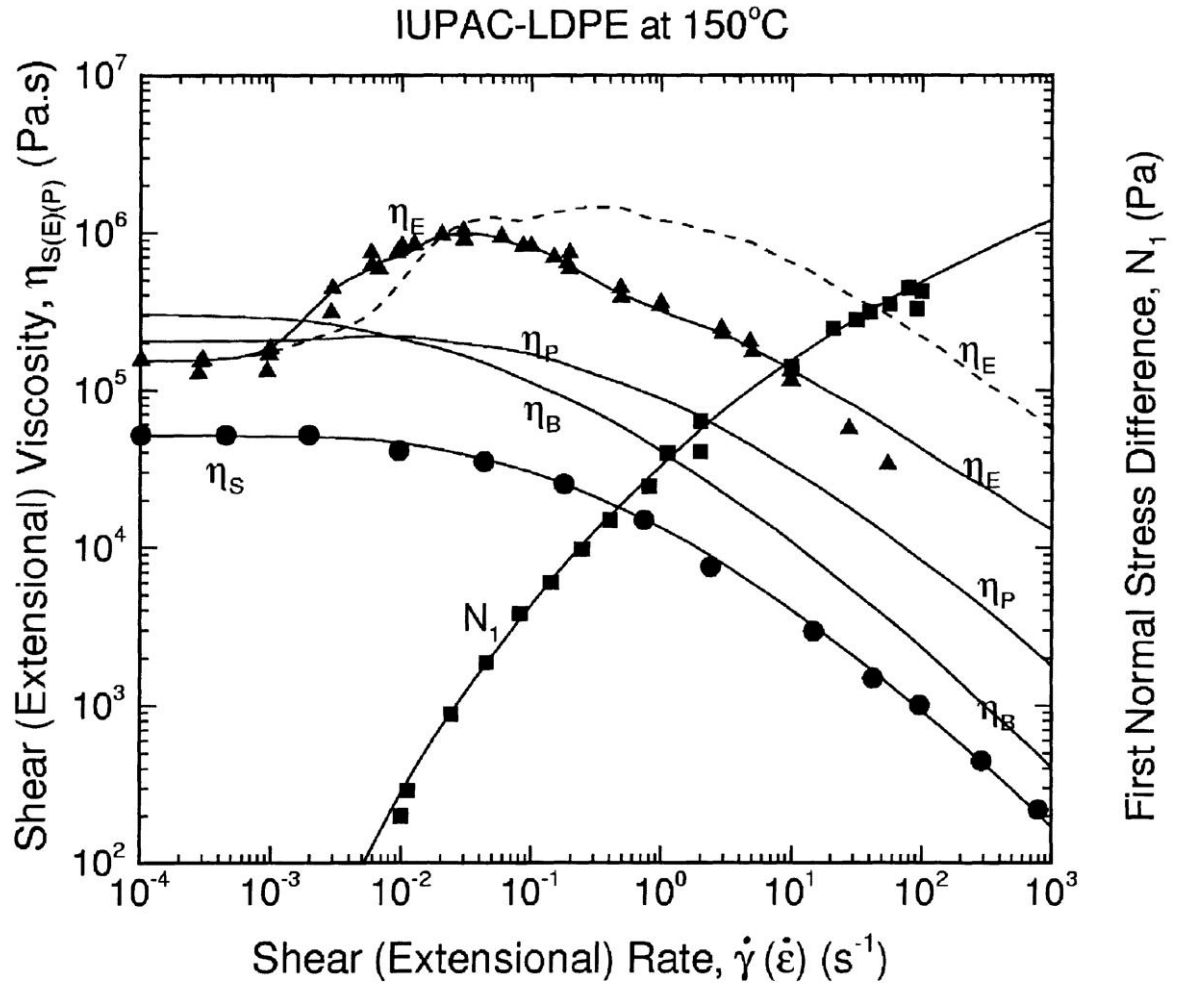
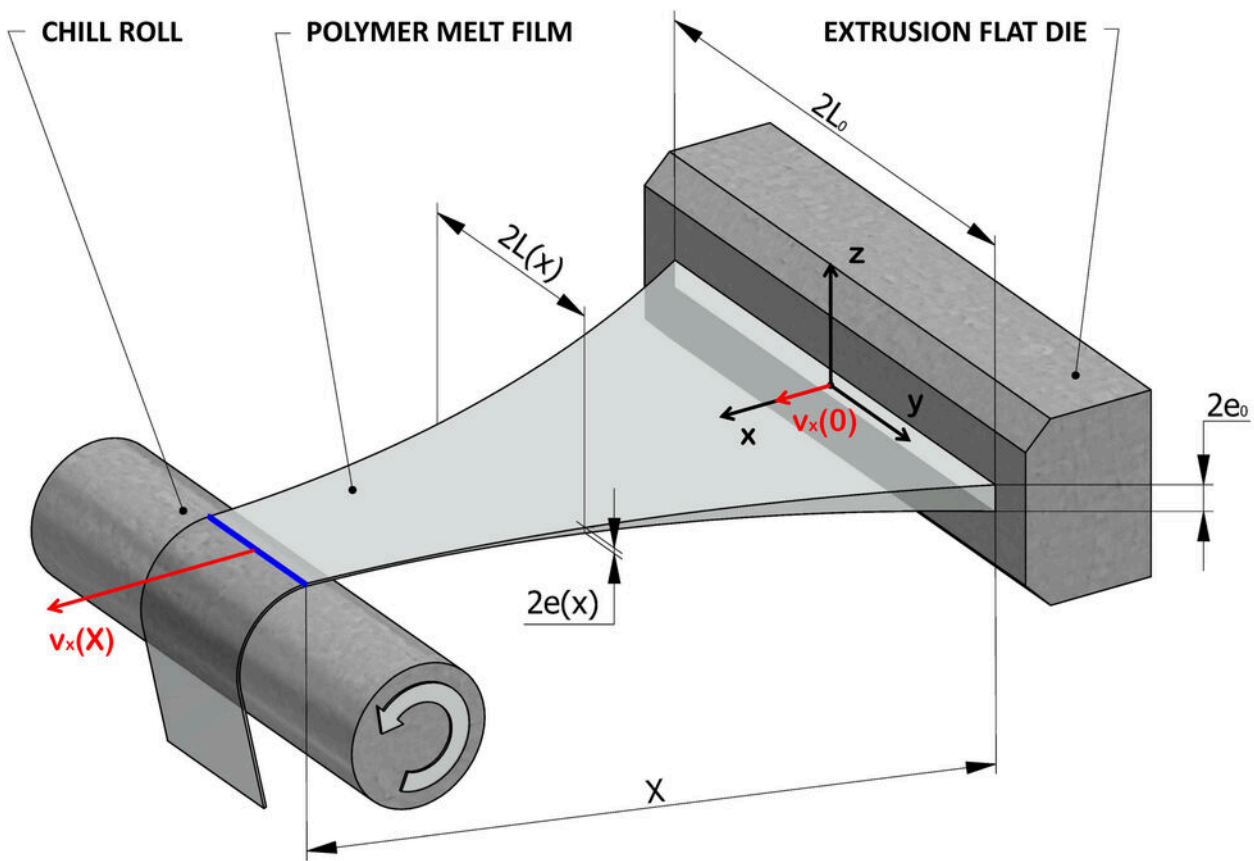
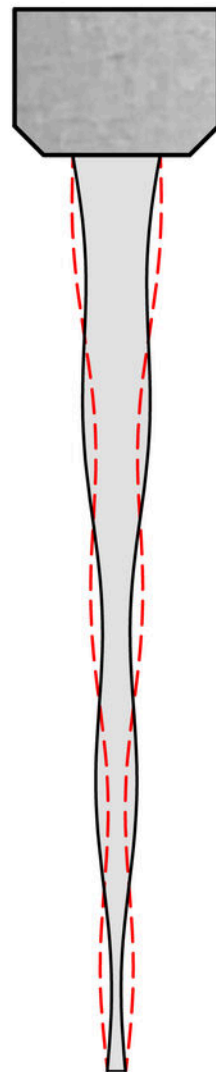
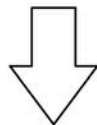
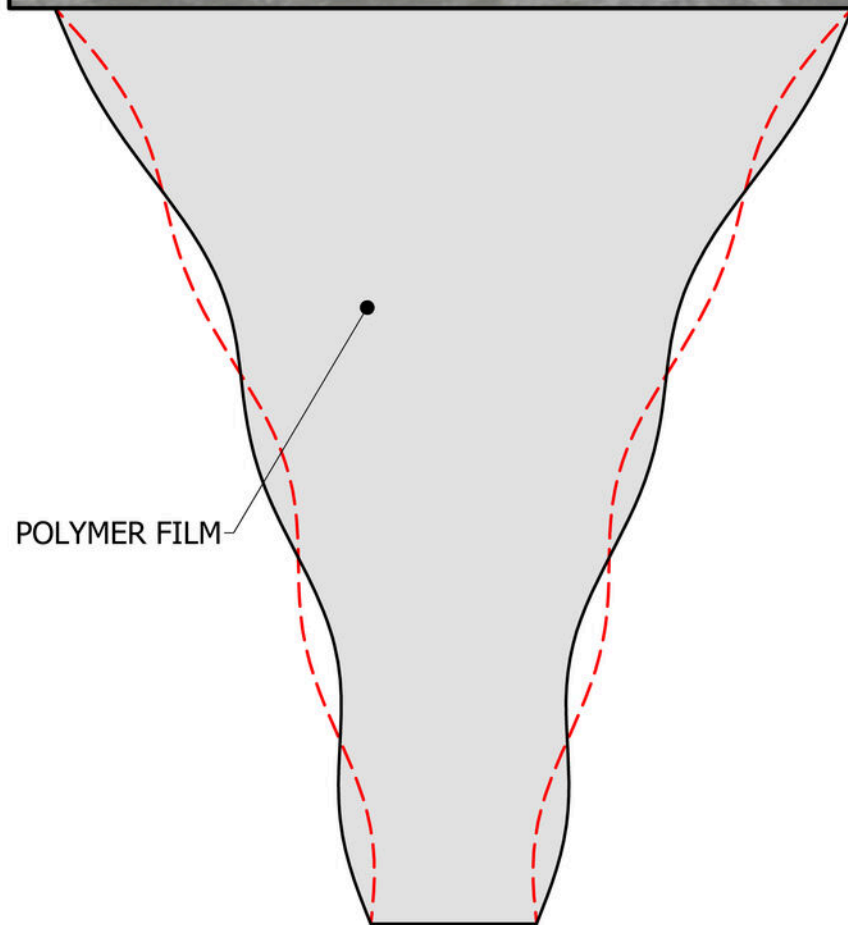
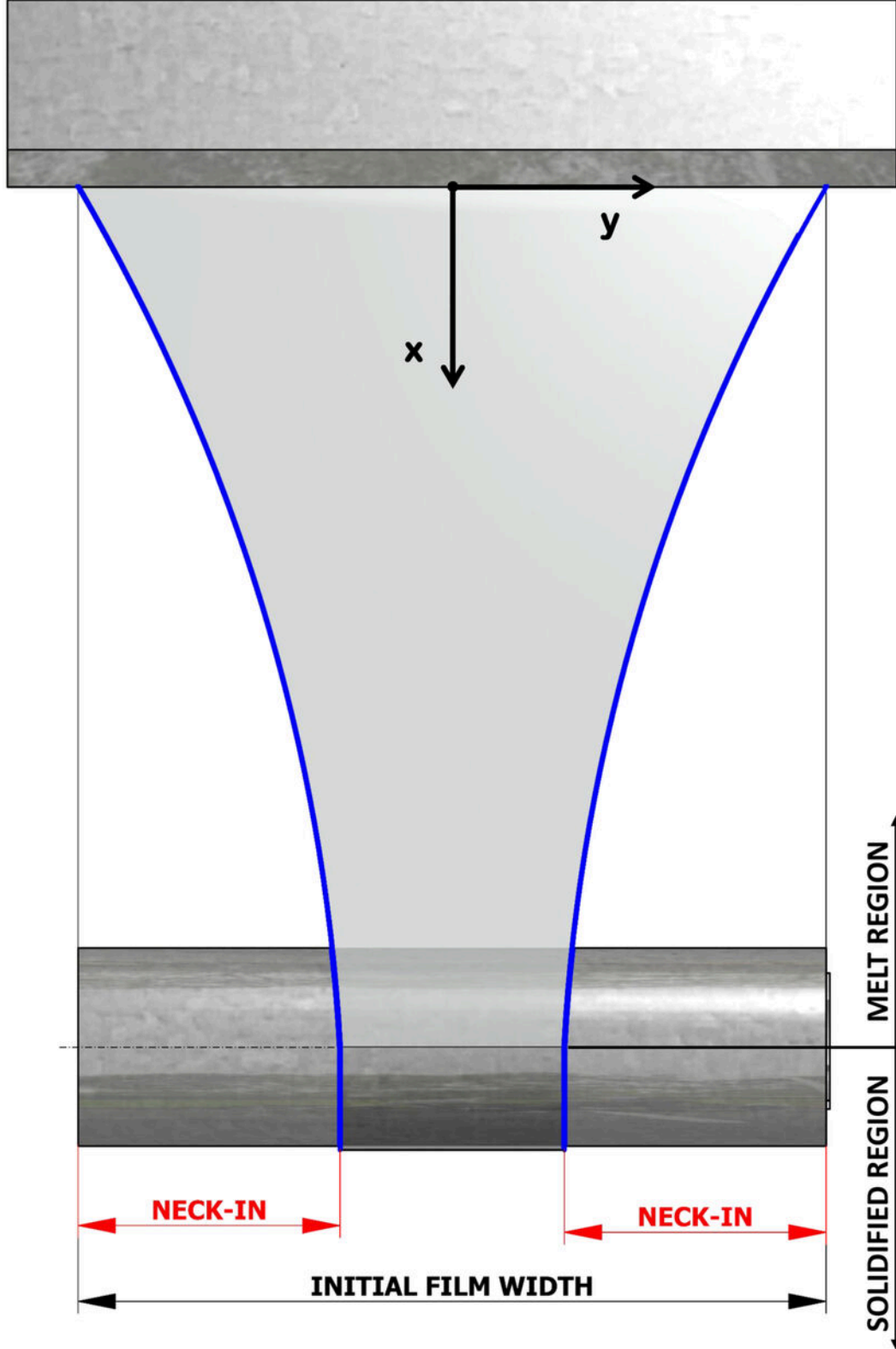


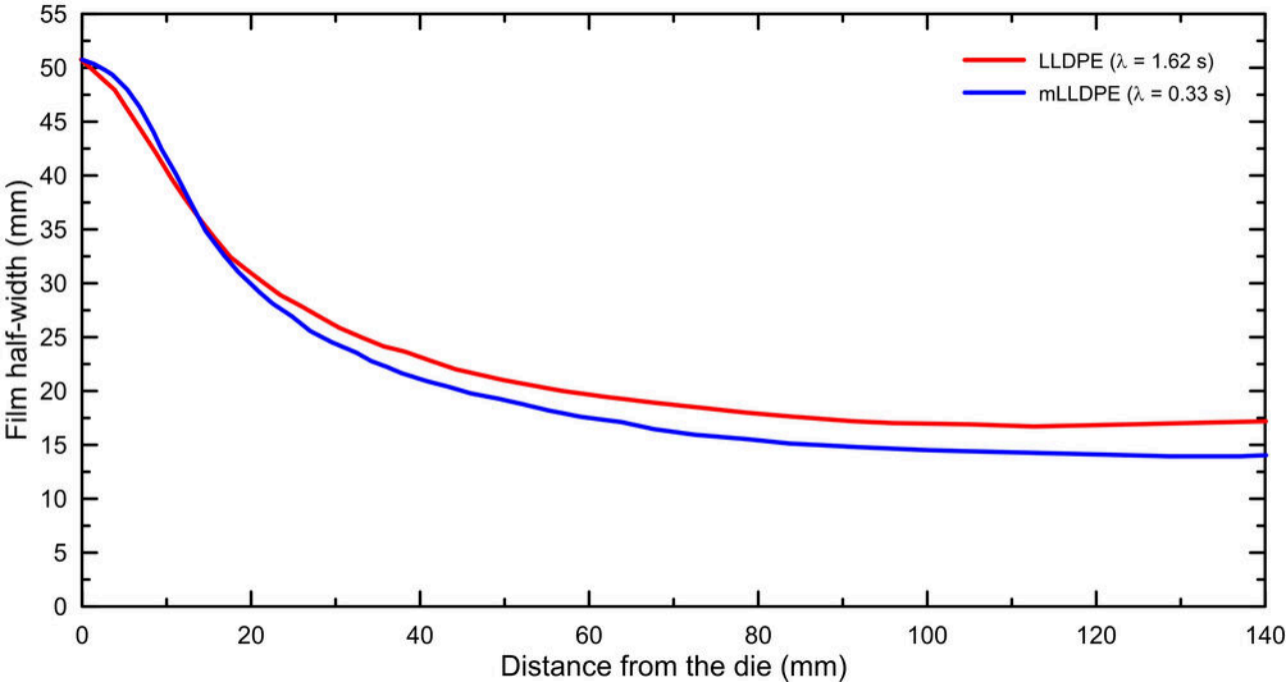
Fig. 23: Prediction of steady shear viscosity ( $\eta_s$ ), first normal stress difference ( $N_1$ ) uniaxial ( $\eta_E$ ), planar ( $\eta_P$ ) and biaxial ( $\eta_B$ ) extensional viscosities by using integral constitutive equation of the K-BKZ type with PSM strain-memory function (Eq. 56) for IUPAC-LDPE melt A at 150 °C. Solid symbols visualize experimental data. Solid lines represent model predictions when  $\beta$  parameter in Eq. 56 is allowed to vary with the relaxation time, the dashed line corresponds to the uniaxial extensional viscosity predicted by using a single value of  $\beta$ . Reproduced with permission from J. Nonnewton. *Fluid Mech.* 97 (1), 13 (2001) [145]. Copyright 2001, Elsevier.

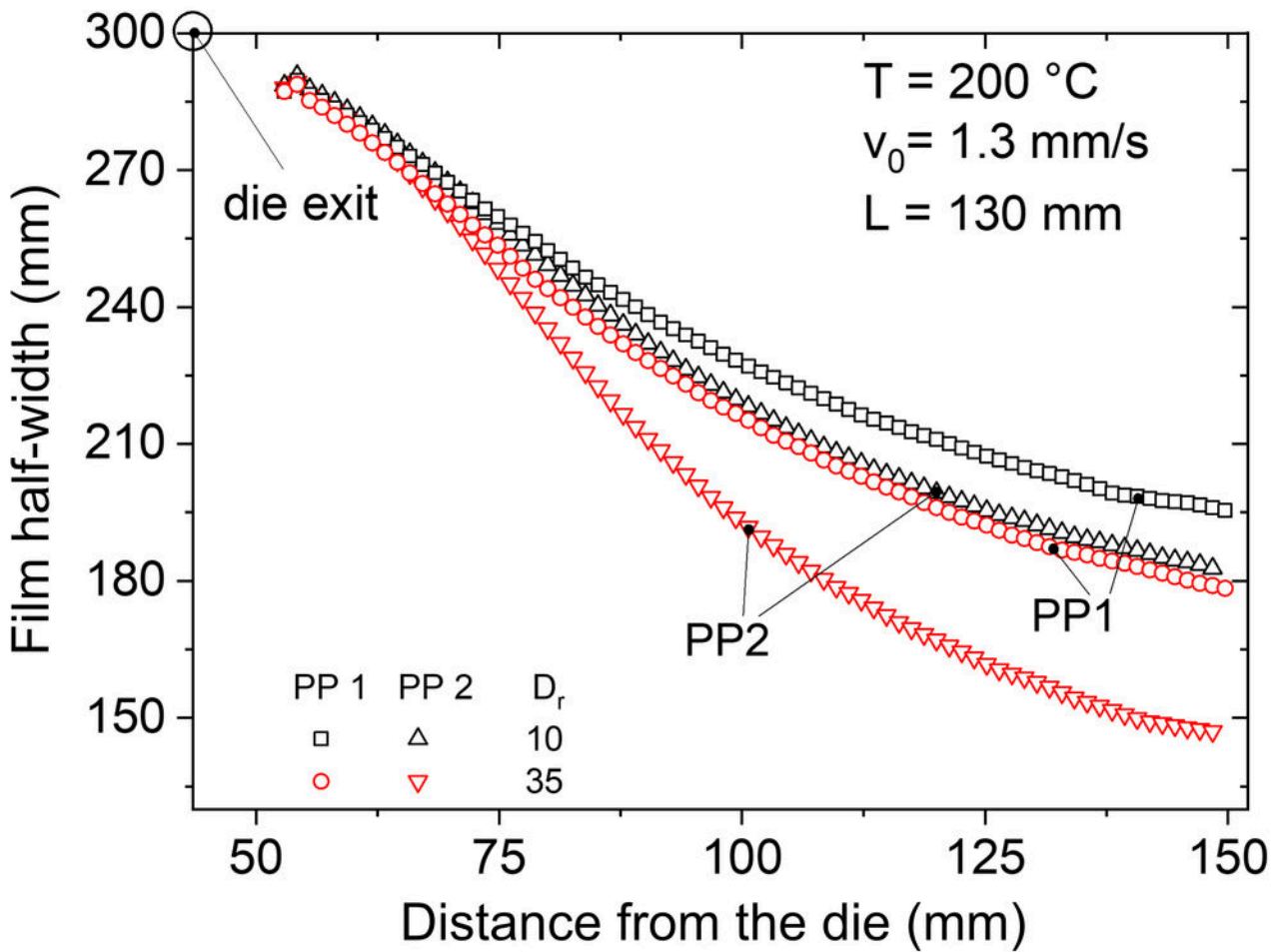


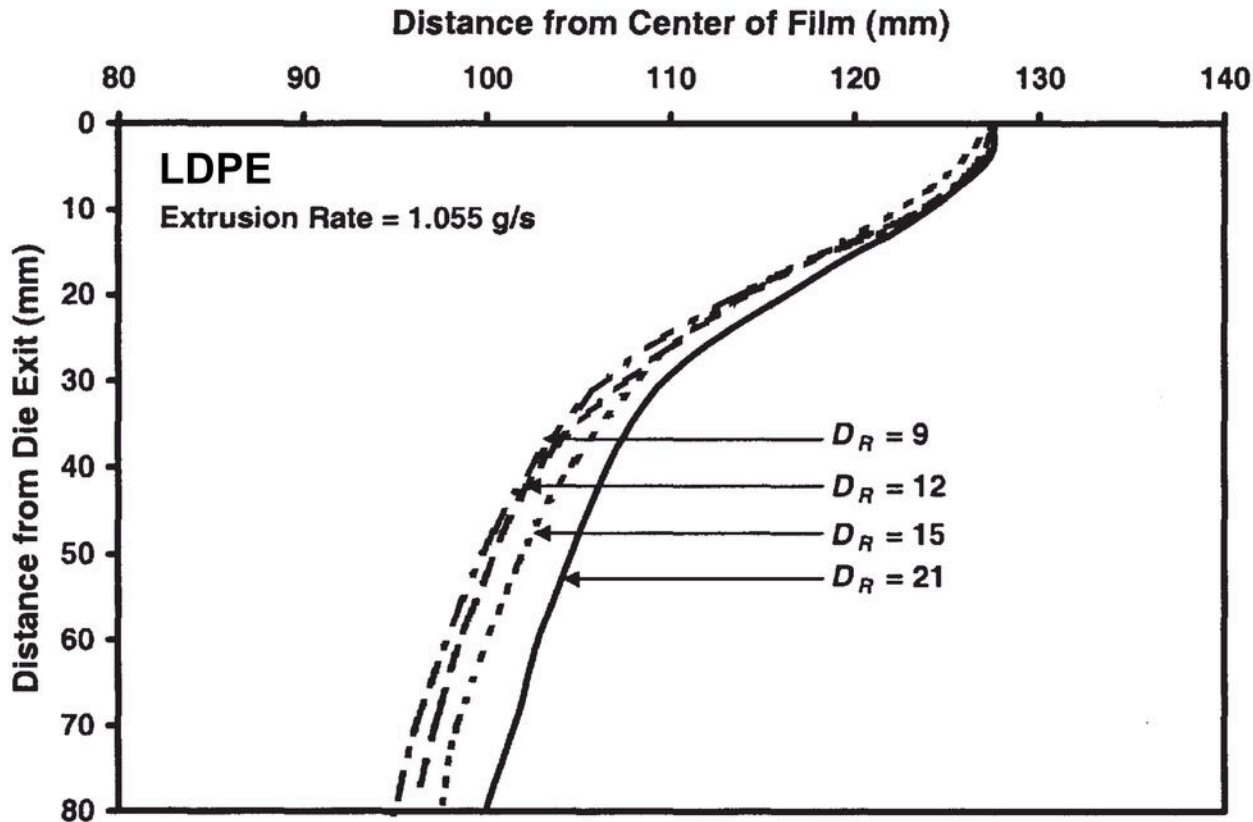












Distance from Center of Film (mm)

80

90

100

110

120

130

140

0

LLDPE

Extrusion Rate = 0.87 g/s

10

20

30

40

50

60

70

80

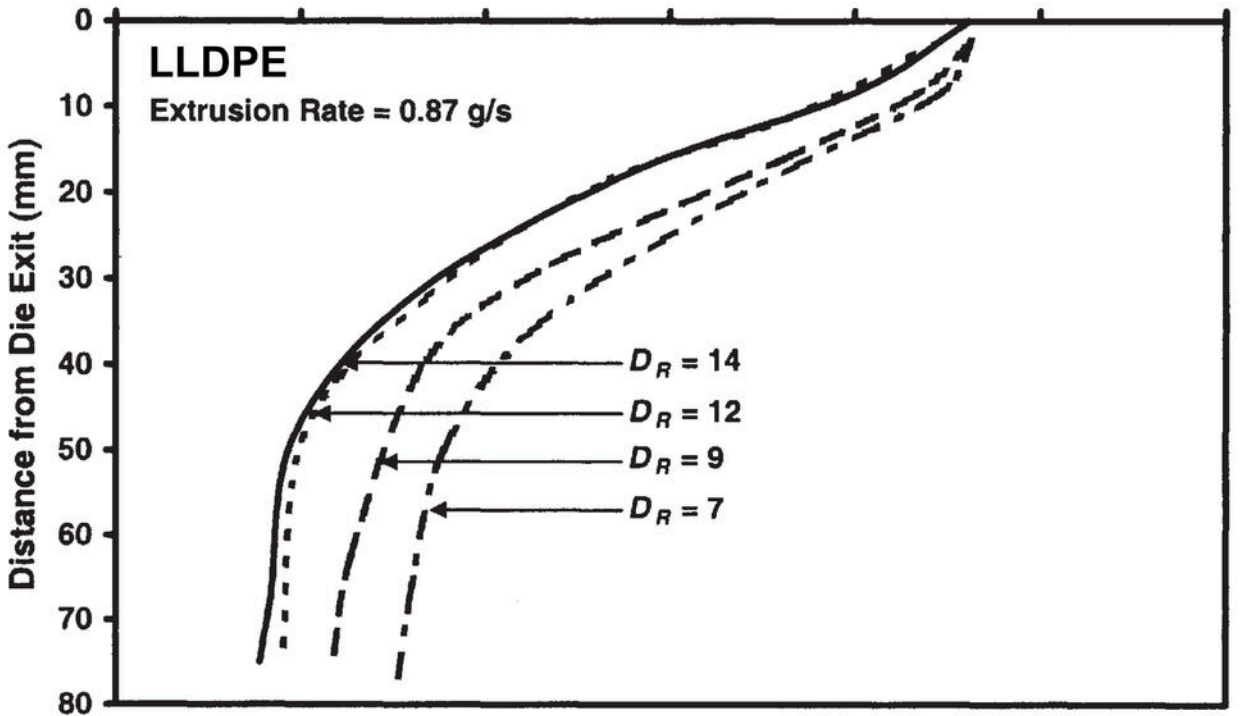
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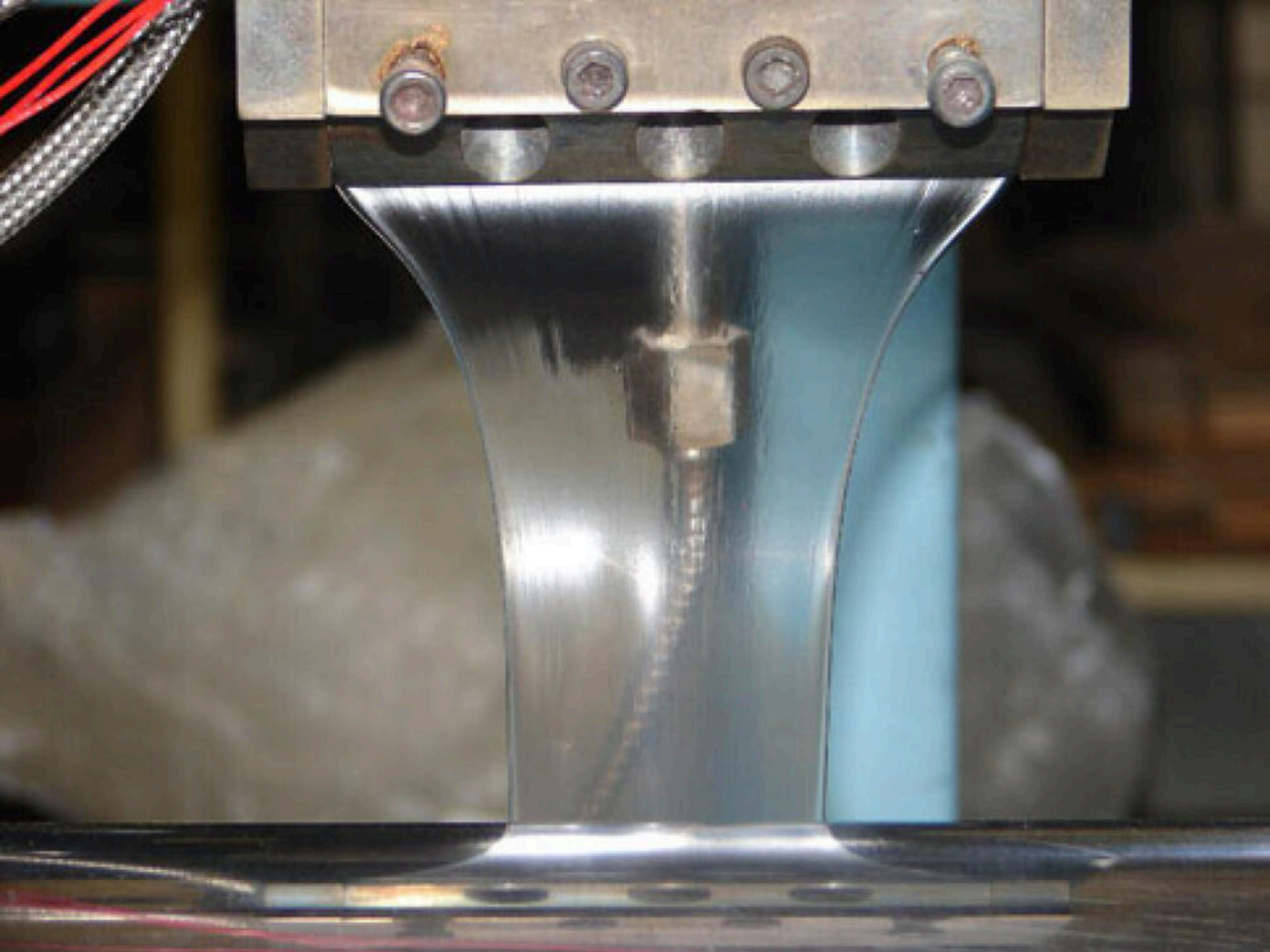
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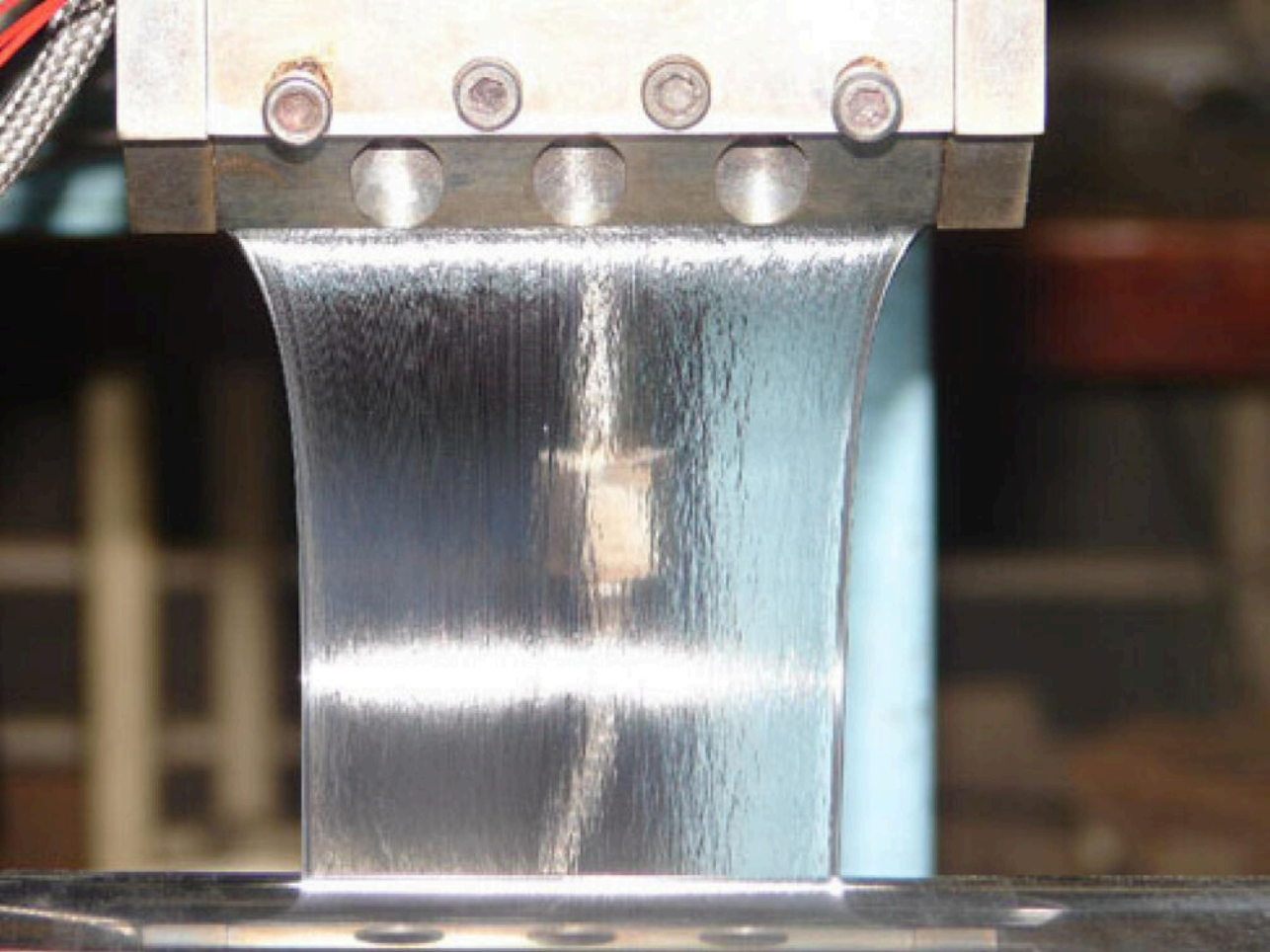
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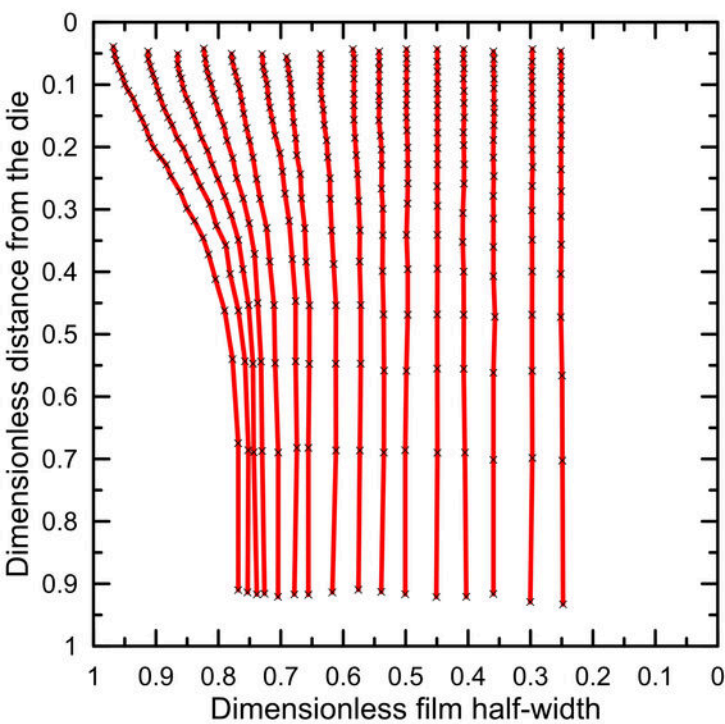
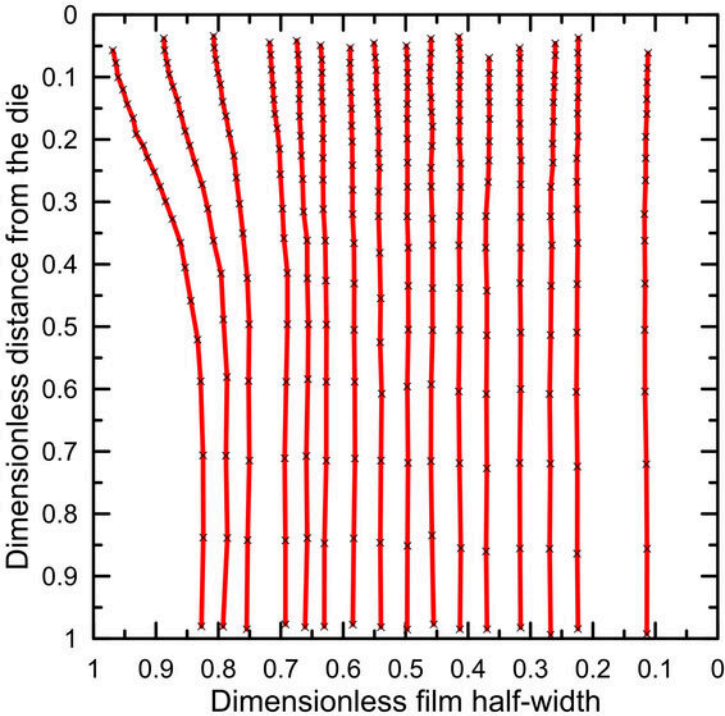
$D_R = 7$

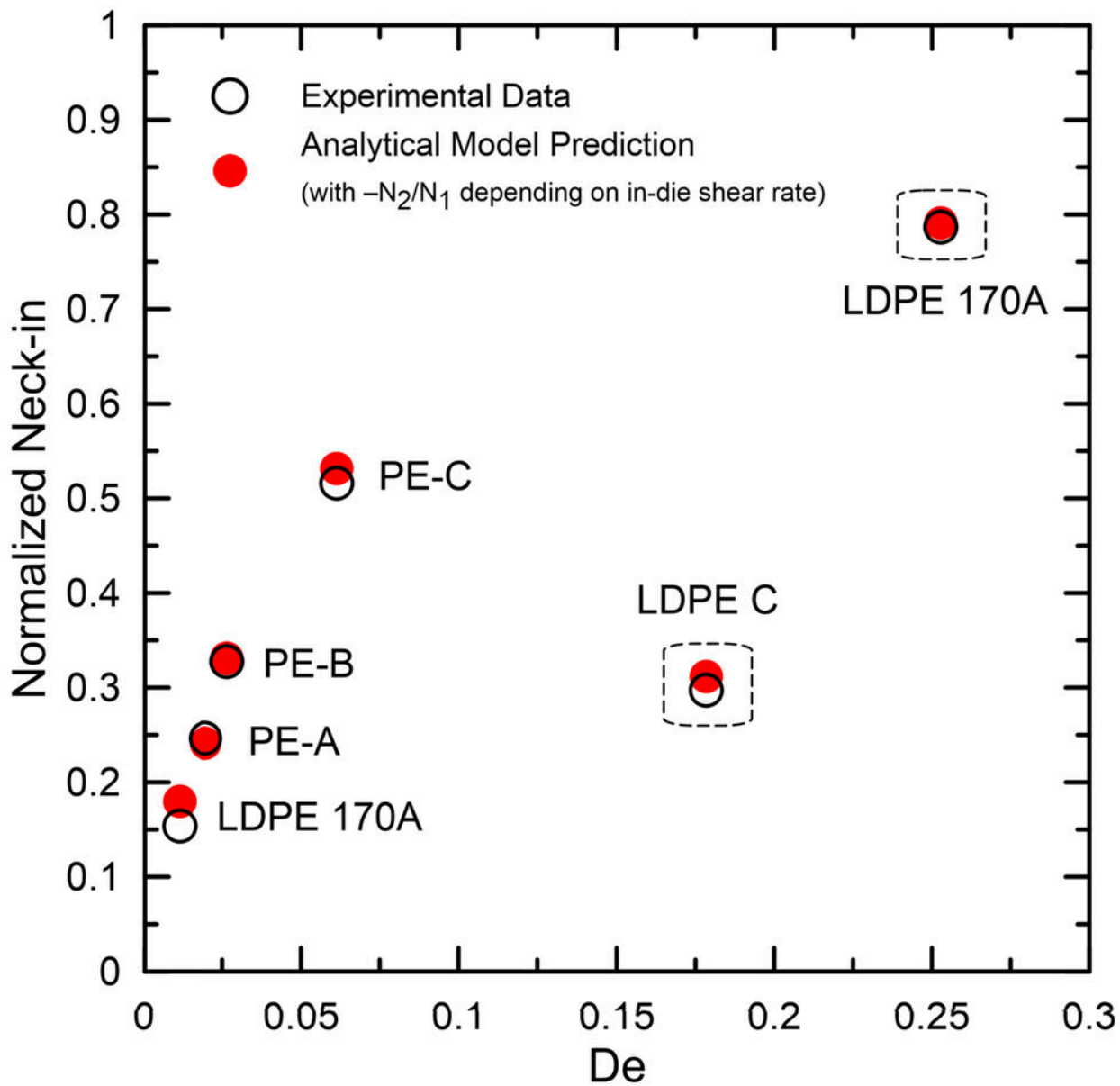
Distance from Die Exit (mm)



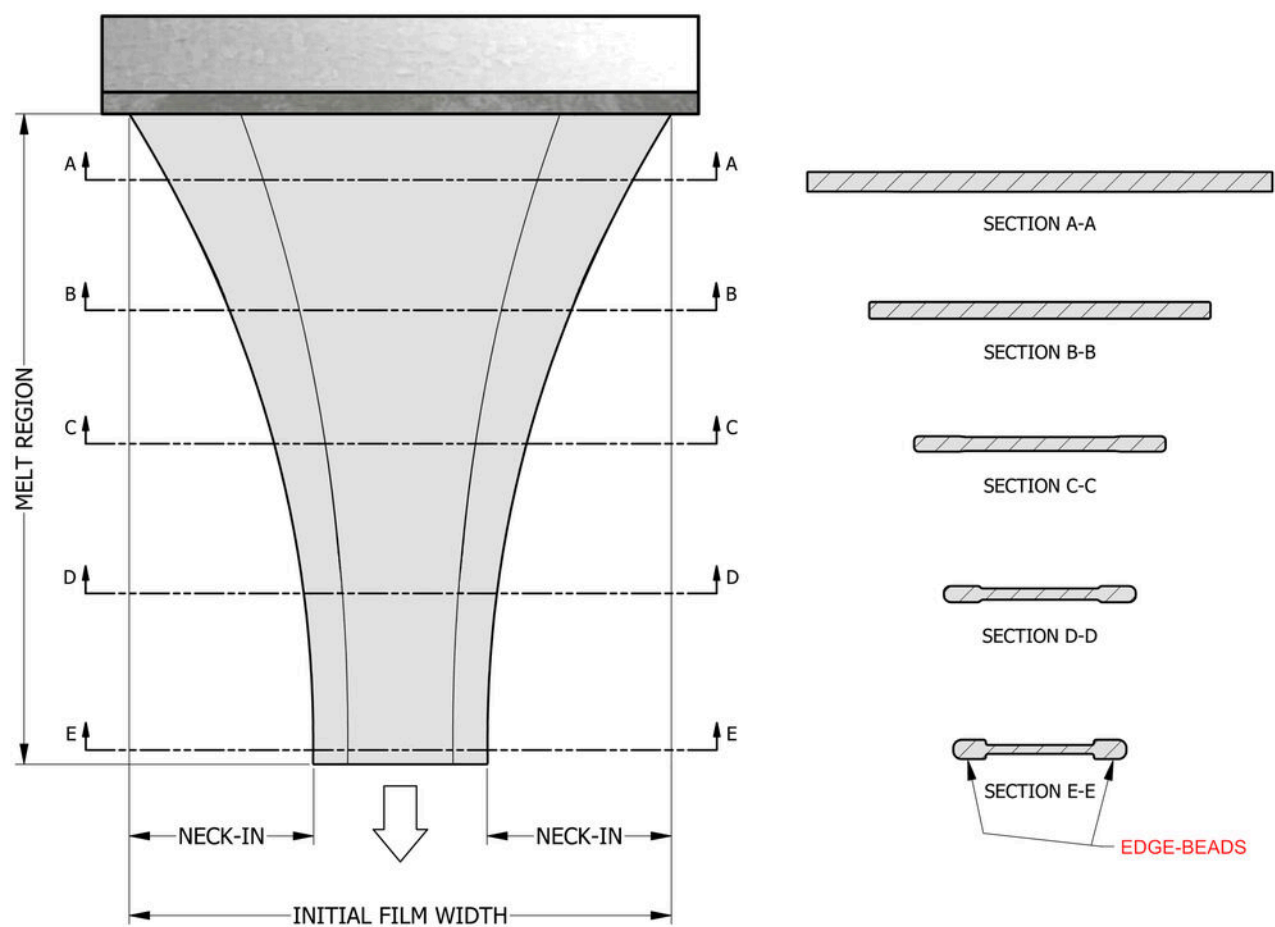




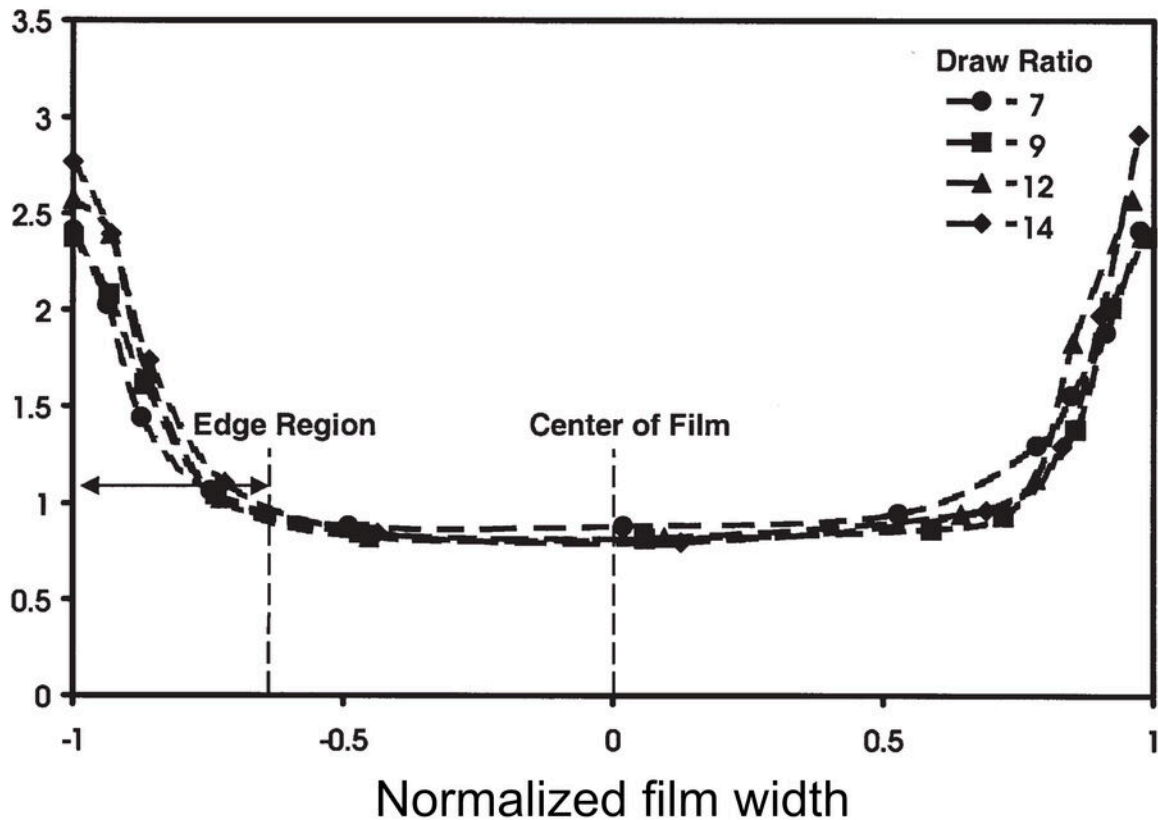




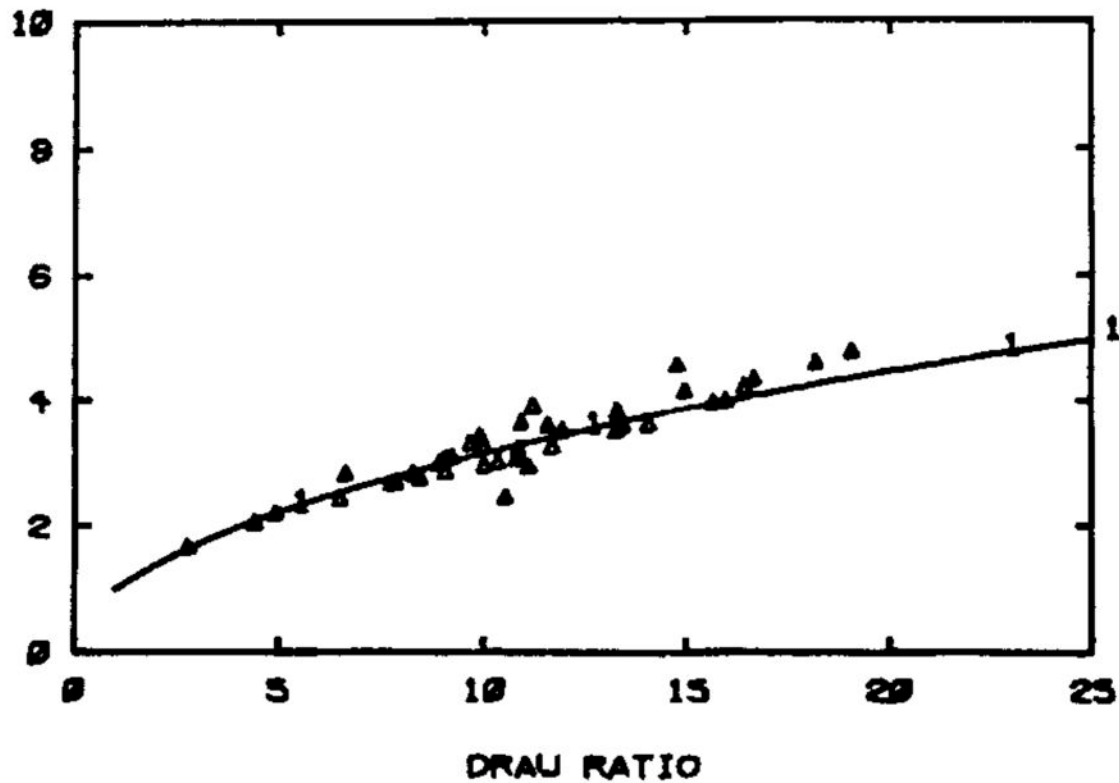


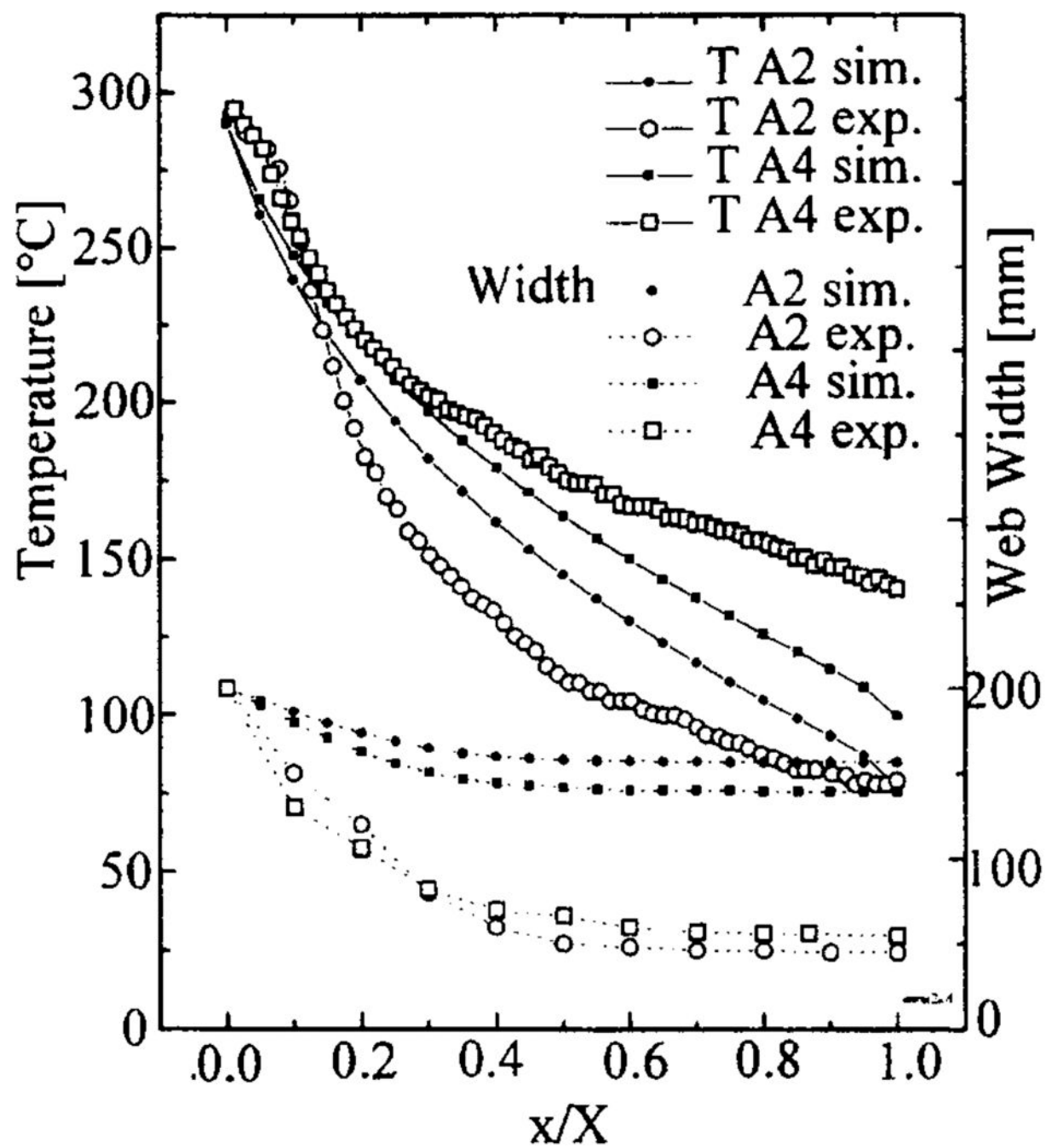


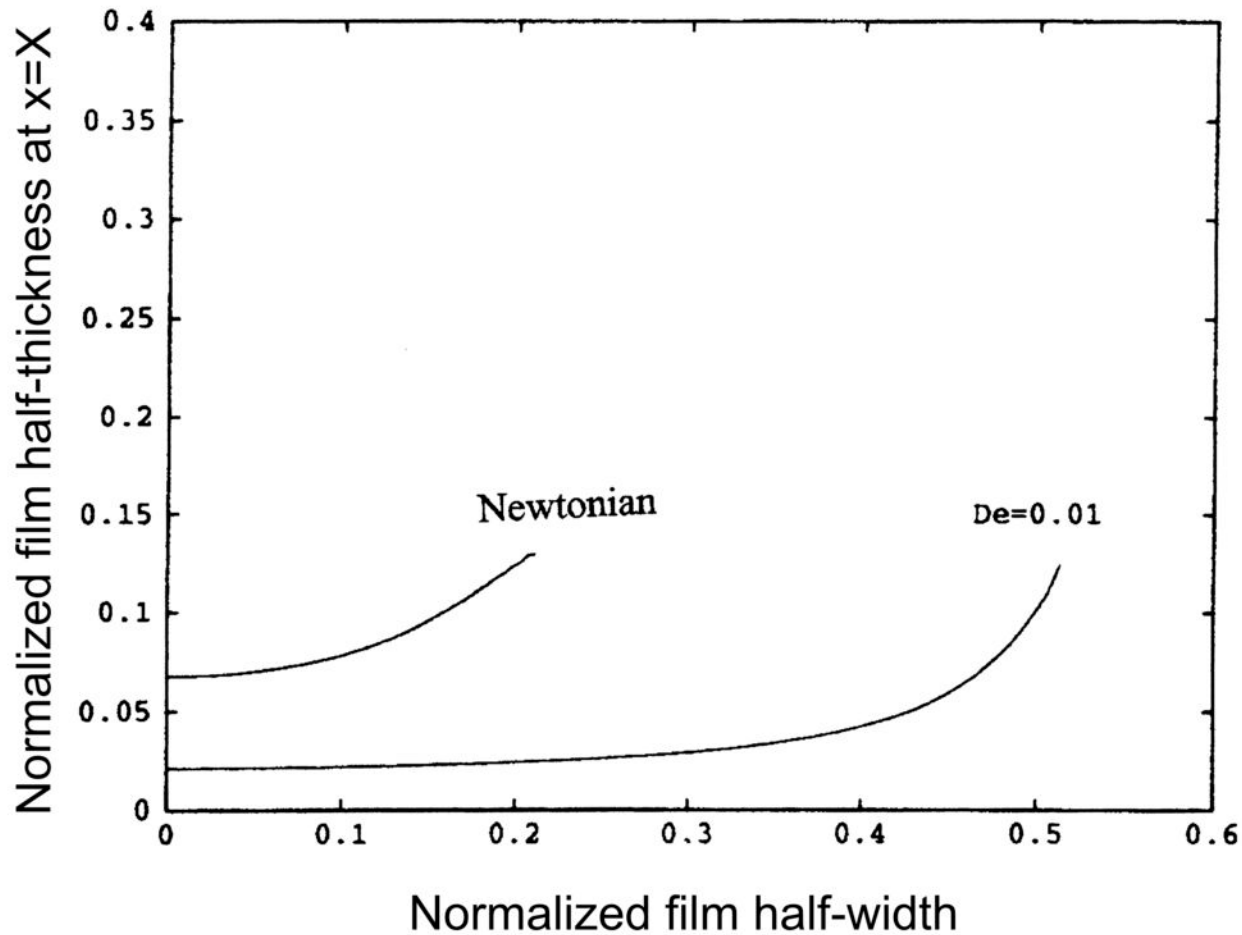
Normalized film thickness

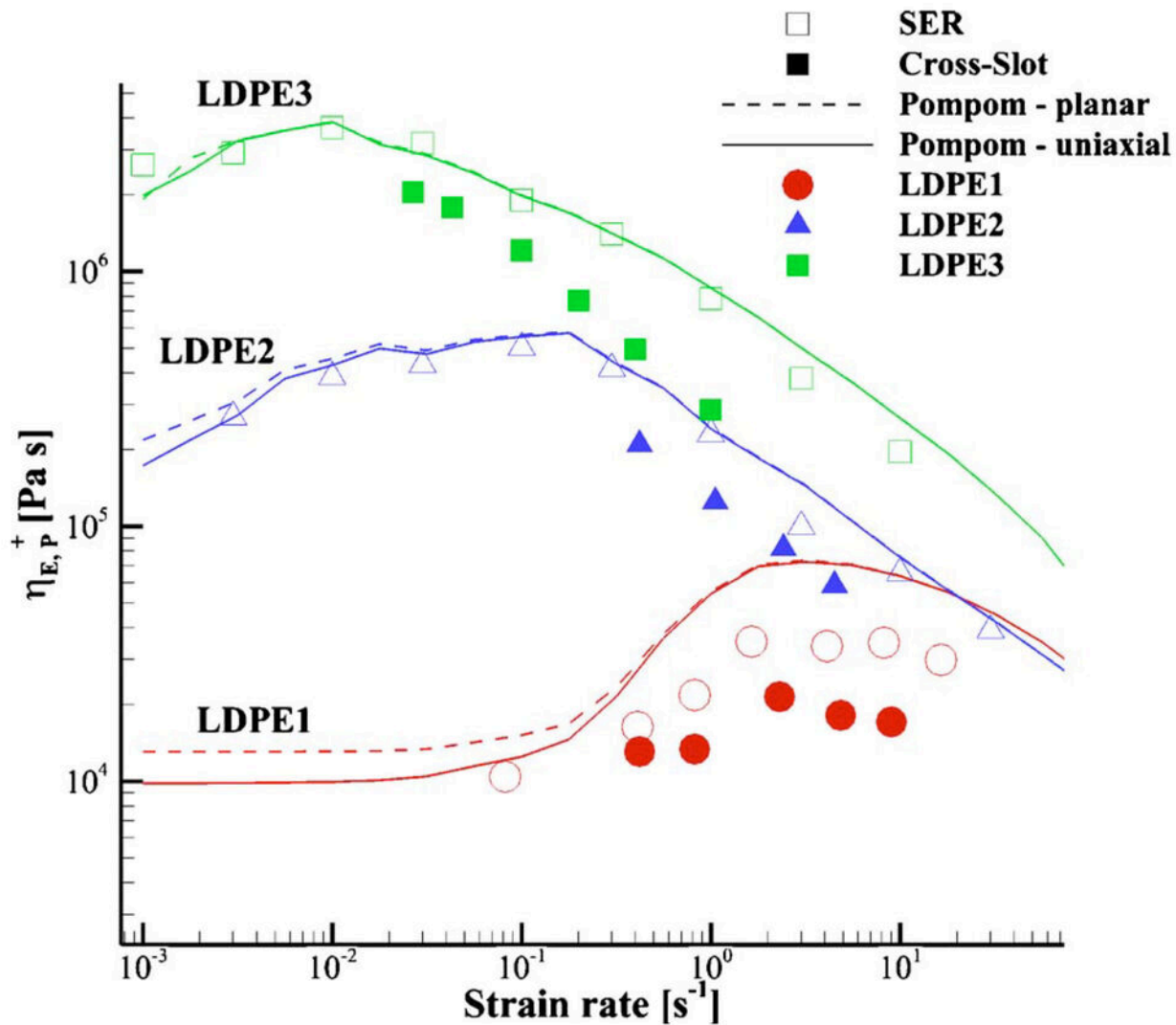


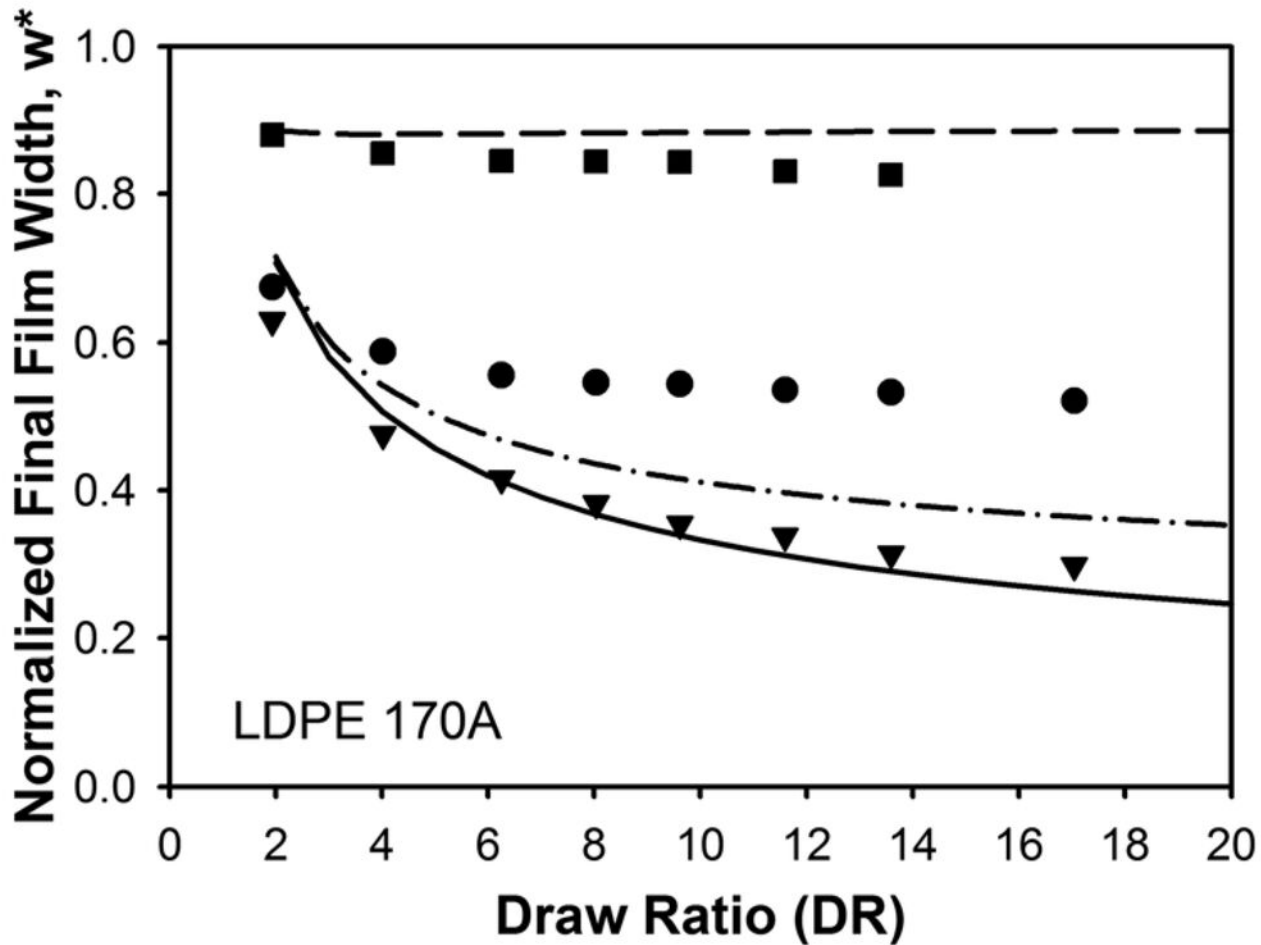
BEAD RATIO

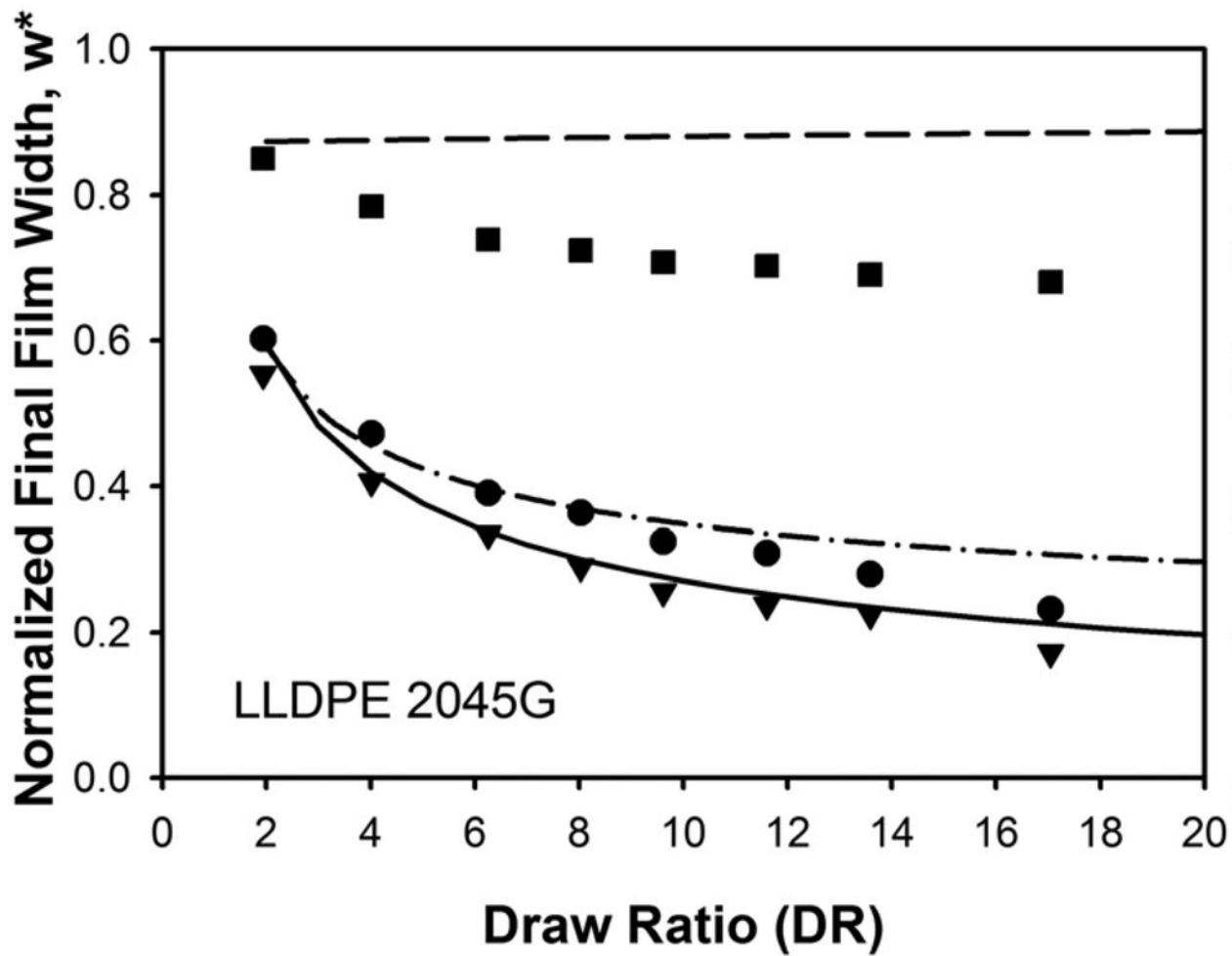




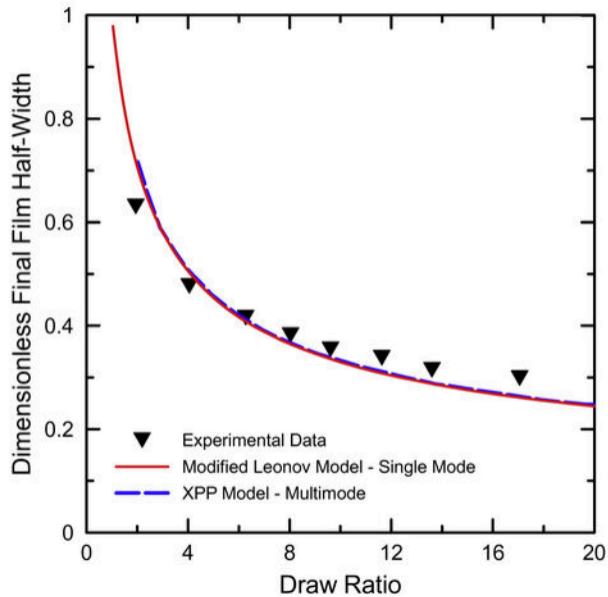
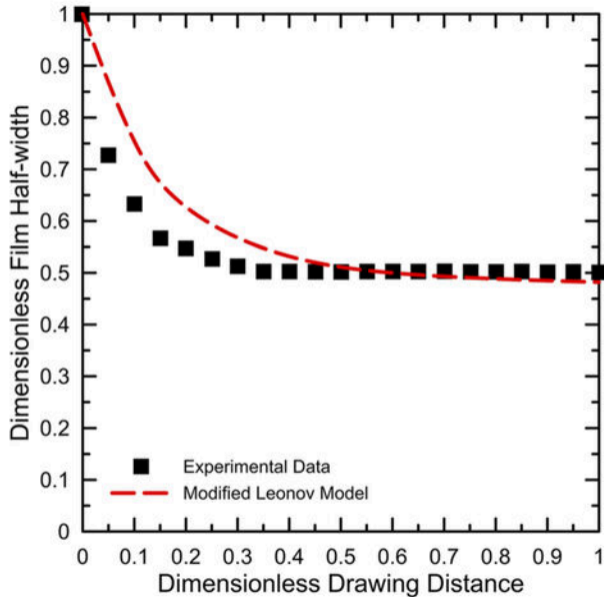


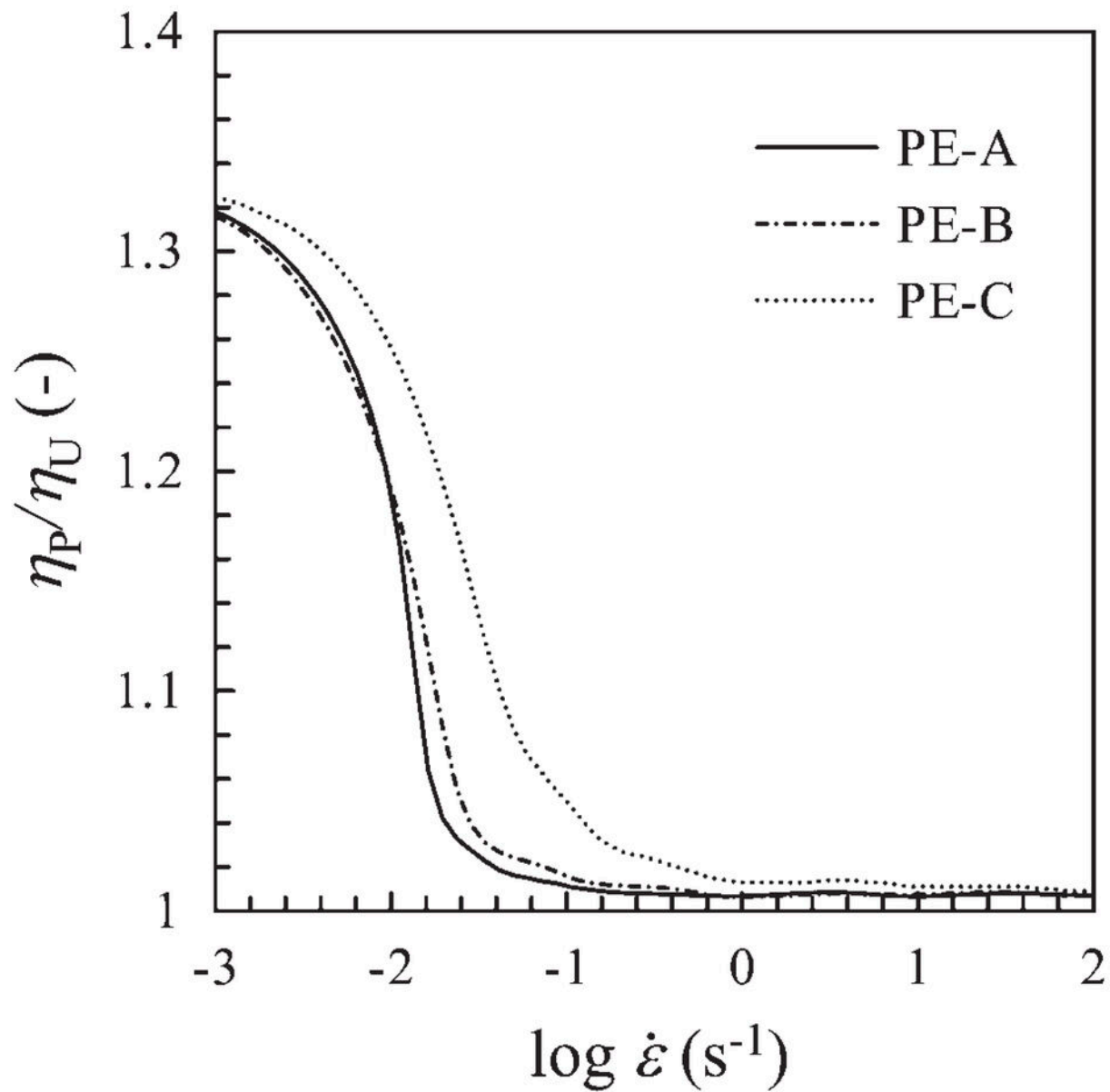


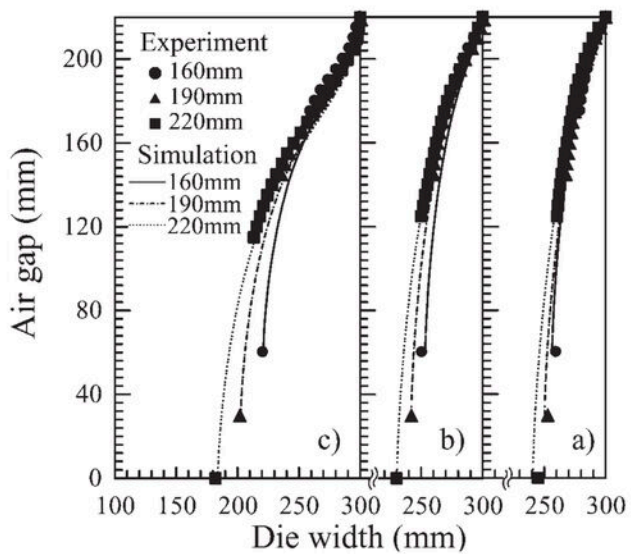
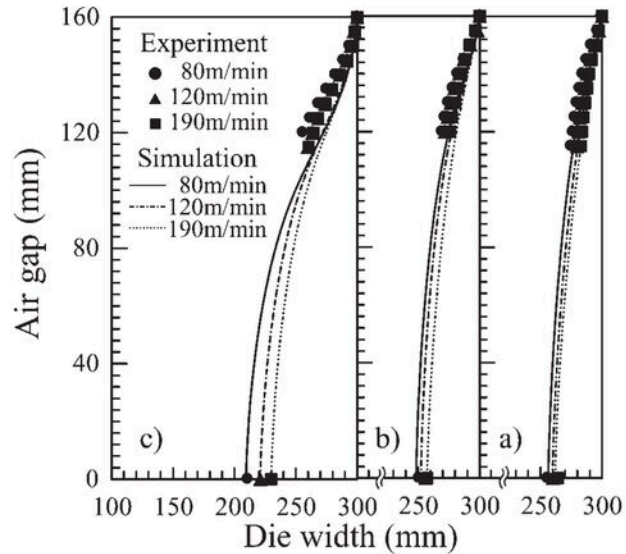


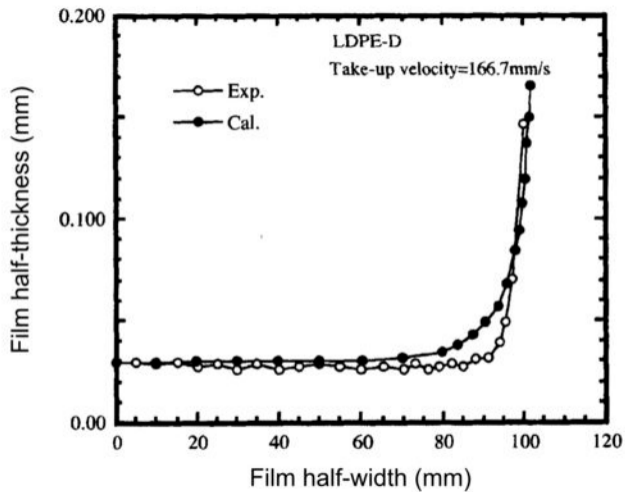
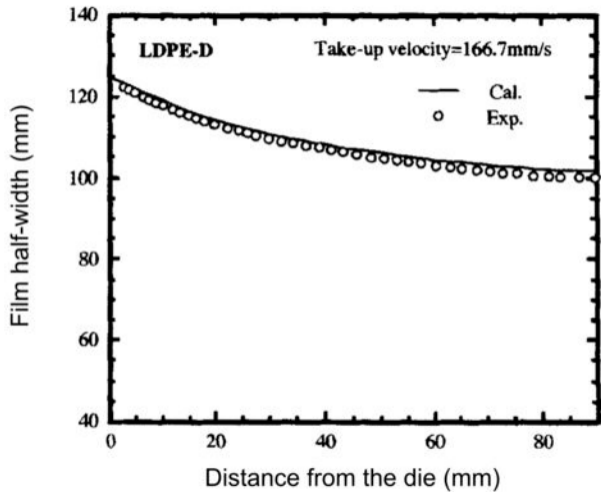


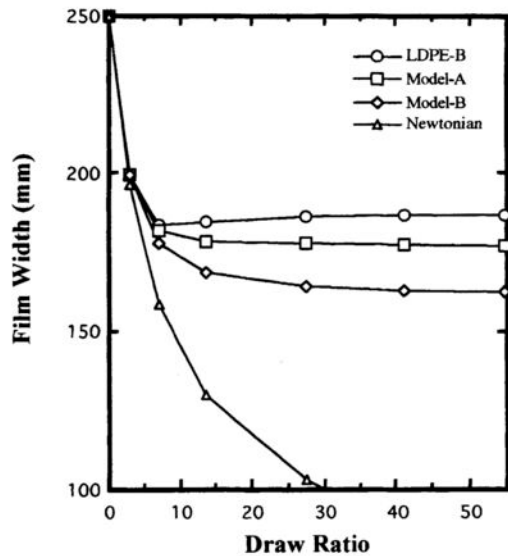
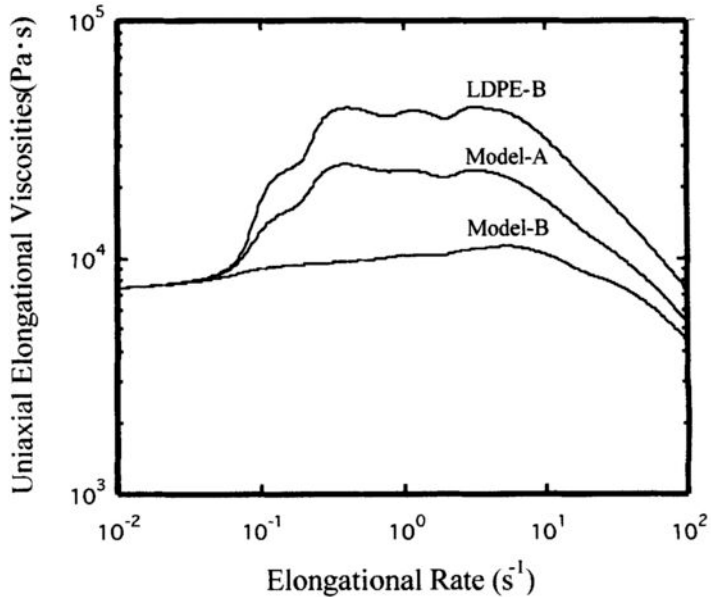












## IUPAC-LDPE at 150°C

