

Doctoral Thesis

Manufacturing of Blends based on Biopolyesters and Polylactides: Process-induced Structure and Properties

Zpracování PBAT/PLA směsí: vliv procesních podmínek na strukturní a užitné vlastnosti

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RESUME

Process-induced changes in thermo-mechanical viscoelastic properties and the corresponding morphology of biodegradable polybutylene adipate terephthalate (PBAT) and polylactic acid (PLA) blown film blends modified with four multifunctional chain-extending cross-linkers (CECL) were investigated. The introduction of CECL modified the properties of the reference PBAT/PLA blend significantly. The thermal analysis showed that the chemical reactions were incomplete after compounding, and that film blowing extended them. SEM investigations of the fracture surfaces of blown extrusion films reveal the significant effect of CECL on the morphology formed during the processing. The anisotropic morphology introduced during film blowing proved to affect the degradation processes as well. Furthermore, the reactions of CECL with PBAT/PLA induced by the processing depend on the deformation directions. The blow-up ratio parameter was altered to investigate further process-induced changes proving synergy with mechanical and morphological features. Using blown film extrusion, the elongational behavior represents a very important characteristic. However, its evaluation may be quite often problematic, but with the SER Universal Testing Platform it was possible to determine changes in the duration of time intervals corresponding to the rupture of elongated samples.

RESUME IN CZECH

V disertační práci byly zkoumány zpracováním indukované změny v termomechanických a viskoelastických vlastnostech a odpovídající morfologii vyfukovaných fólií z biodegradovatelné směsi polybutylentereftalátu adipátu (PBAT) a kyseliny polymléčné (PLA) modifikované čtyřmi typy multifunkčních CECL (chain-extending cross-linkers). Zavedení CECL významně ovlivnilo vlastnosti referenční směsi PBAT/PLA. Termální analýza ukázala, že chemické reakce nebyly dokončeny během kompoundace, ale probíhaly i během zpracování směsi vytlačovacím vyfukováním. SEM analýzy lomových povrchů vyfukovaných filmů odhalily významný vliv CECL na morfologii vytvořenou během zpracování i na degradační procesy. Navíc se reakce CECL s PBAT/PLA anizotropická - liší ve směru souběžném a kolmém na směr vyfukování. Pomocí změny rozfukovacího poměru byla zkoumána synergie mezi mechanickými a morfologickými vlastnostmi modifikovaných směsí. U vytlačovacího vyfukování je důležitá znalost elongační viskozity. Její stanovení pro PBAT/PLA je však na SER reometru značně problematické, proto byl vypracován nový postup hodnocení deformačního chování v tahu pro biodegradabilní alternativy syntetických polyolefinů.

ABSTRACT

Biopolymer-based packaging materials have become of greater interest to the world due to their biodegradability, renewability, and biocompatibility. The use of biopolymers as packaging materials is becoming a well-established trend worldwide due to their major benefits over synthetic plastics. However, the main limitation of using biopolymers in food packaging is their weak mechanical strength. Many biopolymers are combined with other biopolymers to balance the properties required for specific performance. Modifications with chain extenders cross-linkers are tested to provide optimized properties of the packaging films. This thesis addresses process-induced changes in thermo-mechanical viscoelastic properties, and the corresponding morphology of biodegradable polybutylene adipate terephthalate (PBAT) and polylactic acid (PLA) blown film blends modified with various multifunctional chain-extending cross-linkers (CECL).

The evaluation of the mechanical properties has allowed to define how the performance behaviour of the blend with the CECL occurs. Also, aging is studied to predict how this change can occur over time. With the help of thermal analysis via differential scanning calorimetry, it is shown that the CECL-modified blends do not endure thermo-oxidation. With these insights, further investigations with dynamic mechanical analysis show that certain CECLs barely affect the glass transition temperature. With the morphology analysis, by using a scanning electron microscope, the investigations have shown that the fracture surfaces of blown film extrusion reveal their significant effect during the processing. Due to the combined shear and elongation deformation during blown film spherical PLA islands were transformed into long fibrils. The same findings were seen for higher and lower blow-up ratios displaying that the PLA droplets solidify shortly after being extruded by the die and remain undeformed during blowing, while PBAT experiences all deformations. With the use of the Sentmanat Extensional Rheometer, the effect of the chosen CECL on elongation properties was determined via changes in the duration of time intervals corresponding to the rupture of elongated samples indicating the efficiency of the individual modified blends in the blown-film extrusion process. Since biodegradation is the main aspect when using biopolymers, evidence on how the disintegration can be affected by the multifunctional CECL was taken into study. The changes were significant concerning the unmodified blend where at 30 °C a pronounced annealing effect occurred proving that disintegration consists of processes that affect the amorphous and crystalline phase of PBAT in a matter that cannot be understood by a hydrolytic chain degradation only. Moreover, with the analysis of the mean masses, was observed that mass loss and structural disintegration were not due to molecular degradation but were mainly due to mechanical disintegration.

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1. STATE OF THE ART

The global plastic waste problems positively affected the development of biodegradable polymers and other sustainable materials [1-4] towards substitution of traditional packaging plastics such as polyethylene (PE) and polystyrene (PS) by biodegradable starch and biopolyesters [5-7]. The term biopolymer summarizes three classes of polymers:

- 1. Polymers that are biodegradable but do not necessarily stem from renewable resources e.g. poly(butylene adipate-co-terephthalate) (PBAT), polycaprolactone (PCL);
- 2. Polymers that stem from renewable resources directly e.g. polylactic acid (PLA), Polyhydroxyalkanoate (PHA);
- 3. Polymers that are synthesized using monomers of biological origin but are not biodegradable e.g. polyamide 6 (PA 6) from sebacic acid, PE from sugar cane.

However, for more demanding engineering applications and to accelerate the conversion to a circular economy, their mechanical properties must be enhanced without compromising their biodegradability. Many biopolymers are blended with other polymers to balance the properties required for specific applications: PLA often exhibits a brittle behavior and is inappropriate for applications requiring high deformation strains [8-9]. Therefore, blending with ductile polymers such as PBAT appears to be a reasonable approach. Often, mineral fillers, e.g. calcium carbonate or talc, are added to a blend to optimize its stiffness and permeation properties [10-12].

The compatibilization of mostly incompatible polymers is a challenge. Recent developments [13-19] showed that chain-extending cross-linkers (CECL) offer paths to improve melt strength, thermal stability and phase compatibility of polymer blends during compounding and film blowing. This may lead to tailor-made polymers for packaging applications, which is the largest market segment in the plastics industry. In Europe, 40 % of the plastics demand is used in packaging, and 50 % of all goods are packaged in plastics [20]. Thus, film blowing became the most important process in the production of polymer films [21-22]. It is important that biopolymers can be processed on PE or PP extrusion lines and adapted to any film-blowing technology for either monolayer or multi-layer films.

The environmental impacts of crude oil products and rising concerns about increasing crude oil prices make it necessary to search for ecologically more sustainable alternatives e.g. biopolymers. These biopolymers should be derived from plants or microbes and are renewable due to the primary sources. The feature of biodegradability is linked to the capability of decomposing into carbon dioxide, methane, and water by enzymatic reactions of micro-organisms [23-25].

Figure 1 depicts common types of bioplastics and how they are classified according to their biodegradability and bio-based content [26].

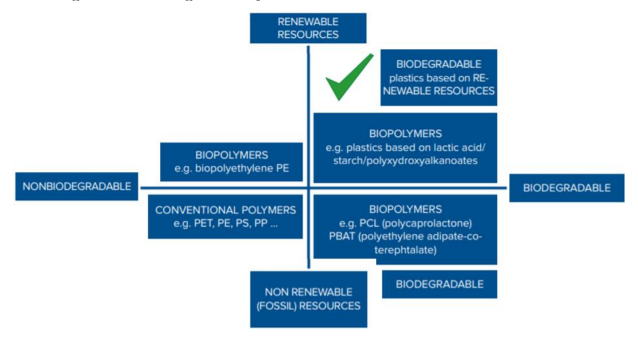


Figure 1: Classification due to the material coordinate system of bioplastics [26].

The group of bioplastics includes polymers such as starch blends made of thermoplastic-modified starch and other biodegradable polymers as well as polyesters such as PLA or PHA [26-29]. Still, this large innovative area of the plastics industry continues to grow due to the introduction of new bio-based monomers such as succinic acid, butanediol, propane diol, or fatty acid derivatives. Several materials in this group, primarily PLA, are striking a new pathway from biodegradation and towards end-of-life solutions such as recycling [8-9]. The renewable base of these materials became the focus of technical developments. Pilot projects aim to establish recycling processes and materials flows.

1.1 PBAT and PLA and their blends

1.1.1 Properties

The most promising and popular biopolymer for a wide range of applications is PBAT obtained by poly-condensation of butane diol, adipic acid, and terephthalic acid [30], Figure 2. It is synthesized by a transesterification reaction. Its biodegradability results from the butylene adipate groups and its stability and mechanical properties from the terephthalate groups [31-32].

Figure 2: Schematic diagram of synthesis of PBAT [30].

Compared to other biodegradable polyesters such as PLA and PBS (polybutylene succinate), the mechanical properties of PBAT are closer to those of LDPE, Figures 3 and 4, making PBAT a promising biodegradable polymer for many applications. PBAT is a flexible polymer with a Young's modulus (E) of 20 to 35 MPa, a tensile strength (σ) of 25 to 40 MPa, and a high elongation at break (ε) close to 1000 % exceeding the properties of many other biodegradable polyesters such as PLA and PBS [33]. The blending of PBAT with PLA increases Young's modulus slightly but decreases the tensile strength and elongation at break [33].

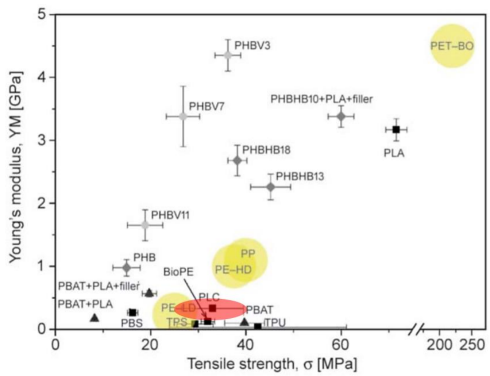


Figure 3: Plot of Young's modulus vs tensile strength for some biopolymers including PBAT (red) compared to some conventional polymers (yellow) [33].

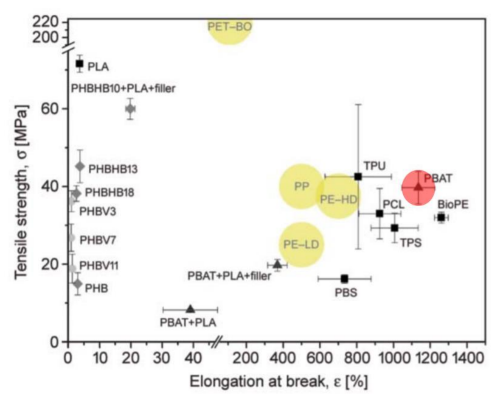


Figure 4: Plot of tensile strength vs elongation at break for some biopolymers including PBAT (red) compared to some conventional polymers (yellow) [33].

The performance of PBAT does not meet many customer requirements. To open new application fields, its performance has to be enhanced by blending, copolymerization, and adding fillers and modifiers. Furthermore, the materials and production costs have to be decreased to a competitive range. Blending with cheap polymers (e.g. starch) or reinforcing polymers (e.g. PLA) are possibilities to achieve these goals while maintaining biodegradability. PLA, Figure 5, is produced either by ring-opening polymerization of lactides or by condensation polymerization of lactic acid monomers [34-35]. These monomers are obtained by fermentation of corn, beet-sugar, cane-sugar, etc. and have the second largest production volume of all biopolymers [8-9].

Figure 5: Chemical structure of PLA [34].

There are three forms of PLA available on the market: L-PLA, D-PLA, and a racemic mixture of D- and L-PLA, Figure 6 [34-35]. Polymerization of L-lactide produces poly(L-lactide) and polymerization of D-lactide results in poly(D-lactide). Poly(L-lactide) and poly(D-lactide) have identical properties except for stereochemistry. Racemic (50 % D and 50 % L-lactide) mixtures result in amorphous poly(D,L-lactide). In addition, PLA can be produced with varying fractions of L and D lactides.

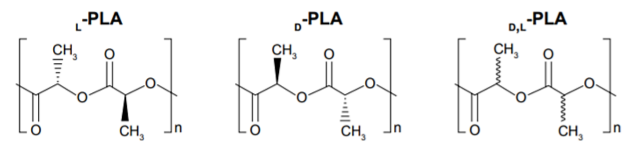


Figure 6: Structure of PLA isomers (L-PLA, D-PLA, D,L-PLA) [35].

PLA is a stiff and brittle polymer with a high Young's modulus (3.4 GPa) and low elongation at break (6%). The tensile strength (53 MPa) is similar to PET (polyethylene terephthalate) and higher compared to HIPS (high-impact polystyrene) or PP (polypropylene) [36-40]. PBAT is a candidate for toughening PLA while maintaining biodegradability, especially for extrusion film and foam applications. However, PLA and PBAT are non-compatible polymers which leads to a multiphase structure with relatively poor mechanical properties. This can be improved by introducing CECLs or compatibilizers to enhance the interface of PLA and PBAT.

Table 1 Comparison of PLA properties to some crude oil-based commodity thermoplastics [36].

	PLA	PET	PS	HIPS	PP
Glass temperature (°C)	55	75	105	-	-10
Tensile strength at break (MPa)	53	54	45	23	31
Tensile modulus (GPa)	3.4	2.8	2.9	2.1	0.9
Elongation at break (%)	6	130	7	45	120

1.1.2 The compounding process

Compounding is the process of mixing polymers, fillers, fibers, and additives to a homogeneous polymer melt that is processable and meets the requirements of the application. In the compounding of polymer blends, selected equipment, process conditions, and blend formulation have a significant impact on the final morphology. Generally, polymer blends are classified as homogeneous blends being miscible on

a molecular level like PLA and PCL [9, 40], or heterogeneous blends being immiscible like PLA and PBAT [30, 33]. Investigations of PBAT/PLA compounds revealed that the interfacial compatibility is poor and can be improved by introducing compatibilizers or reactive CECL [15-19].

Compounding is performed in several steps, Figure 7. The process starts with the intake of the components in a twin-screw extruder via the main hopper or side feeders. The elements of the screw are chosen to affirm fast and homogeneous mixing. The extruder temperature is chosen 20 to 30 K above the melting point of the matrix resin. Due to high temperatures and the shear stresses reactive substances such as oxygen, hydro-peroxides, and catalyst residues are created [41-43]. To control these reactions chemical agents can be added. The polymer melt exits the extruder in the form of strands of filaments having a diameter of 1 to 3 mm. After solidifying they are chopped into pellets for further use e.g. blown film extrusion.

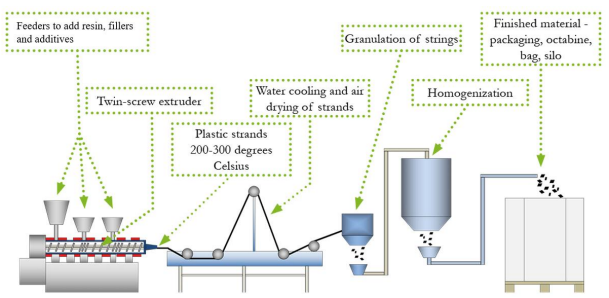


Figure 7: Scheme of the process of polymer blend compounding [41].

1.1.3 The process of blown film

Film blowing is an extrusion process to manufacture continuous polymer films mainly for packaging purposes as schematically shown in Figure 8.

The polymer melt is extruded through a die forming a tube. This tube is vertically drawn by nip rollers and simultaneously blown and cooled by injected air through the center of the die mandrel, thus forming a tubular bubble [44-48]. Finally, this tubular bubble is flattened by nip rollers and wound up.

Relevant processing quantities are:

• **blow-up ratio** (**BUR**)
$$BUR = \frac{Bubble\ diameter}{Die\ diameter}$$
 (1)

• draw ratio (DR) $DR = \frac{\text{thickness of ring}}{BUR*\text{thickness of film}}$ (2)

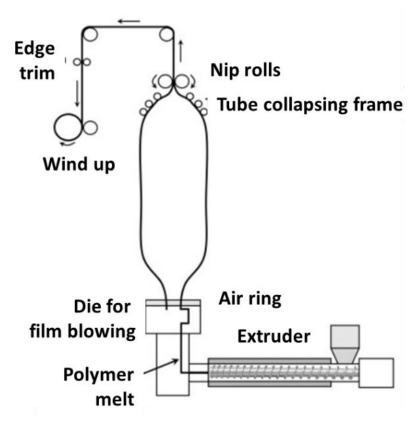


Figure 8: Scheme of a blown film process [44].

During film blowing the polymer melt is drawn both in extrusion (ED) and transverse (TD) directions. This biaxial elongation orients the polymer molecules. The degree of orientation affects the film properties concerning the corresponding directions and depends on the desired film thickness determined by the thickness of the ring, BUR, and DR as well as process parameters such as melt temperature aircooling conditions and melt viscosity [47-49]. The chosen thickness significantly affects the mechanical properties, Figures 9, 10, and 11.

The modulus of elasticity, Figure 9, in ED and TD increase with PLA content for $110 \, \mu m$ as well as for 25 $\, \mu m$ thick films [48]. The tensile strength, Figure 10, in TD of $110 \, \mu m$ films is lower than in ED. This is attributed to the process-dependent larger melt stretching in ED compared to TD leading to a higher molecule orientation in ED.

The elongation at break, Figure 11, decreases with the increasing PLA content. This occurs for 110 μ m and 25 μ m films in both directions when the PLA content increases from 10 to 40 wt. %.

In general one can say that the addition of PLA to PBAT enhances elastic moduli and tensile strengths of their blends but reduces the elongations at break.

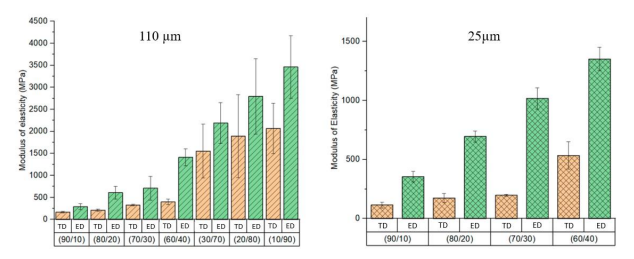


Figure 9: Elastic moduli of PBAT/PLA blends in ED and TD of 110 and 25 µm films [48].

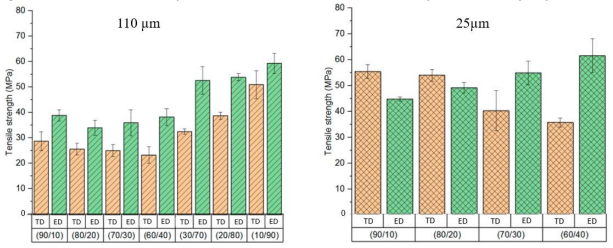


Figure 10: Tensile strengths of PBAT/PLA blends in ED and TD of 110 and 25 μ m films [48].

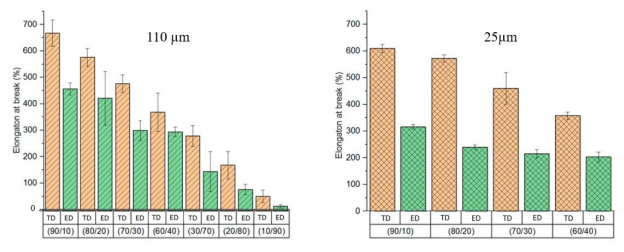


Figure 11: Elongation at break of PBAT/PLA blends in ED and TD of 110 and 25 µm films [48].

This thesis presents the innovative work of investigating the influence that blowup ratio will have on process-induced properties and morphology, with the modification with CECL that hasn't been researched.

1.2 CECL modification of PBAT/PLA blends

Concerning the polymers related to this thesis, PBAT effectively increases the melt strength of PLA during process, but the compatibility is still insufficient due to the poor interfacial adhesion leading to thermodynamically instable morphologies and relatively weak interfacial adhesion after compounding. Thus, cross-linking is considered the most appropriate approach to overcome the limitations of these biomaterials [9,10, 13-19].

Zhou et al. [42] investigated the miscibility and crystallization behavior of PBAT/PLA blends. They found that a "chain mobility promoter" can efficaciously interact with PLA chains to develop oriented crystalline architecture, and the elongational flows can deform the dispersed PBAT phase to nano-fibrils with large surface areas. They allow for more pronounced interactions between the two phases. Dil et al. [43] studied the morphology and miscibility of PBAT/PLA blends without compatibilizer in detail and found that the diameter of the dispersed PBAT phase increased significantly. A result of partial miscibility is the continuous and cocontinuous morphology development with different types of interfaces which seem to show higher interactions. These studies indicate that the level of interaction in with low interfacial tension polymer blends and miscibility/immiscibility can have a determining effect on the morphology evolution. Chain extenders cross-linkers as well as compatibilizers have been used to improve melt strength and thermal stability in polymer blends.

Al-Itry et al. [15] investigated the thermal stability of PLA and PBAT during processing and the resulting structural, rheological, mechanical and morphological properties of PBAT/PLA blends compatibilized with Joncryl® ADR 4368 (BASF). Above temperatures of 180 °C they found a decrease of molecular weight and intrinsic viscosity. The incorporation of Joncryl® increased the thermal stability, molecular weight, intrinsic viscosity, shear thinning and elasticity during processing due to the epoxy reactive functional groups.

Wang et al. [13] found that an epoxy-terminated branched polymer (ETBP) enhanced interfacial compatibility and mechanical properties. An ETBP content of 3.0 phr increased elongation at break of PBAT/PLA blends from 45 to 270 % and impact strength from 26 to 45 kJm⁻². DMA measurements revealed that the difference between the $T_{\rm g}({\rm PLA})$ and $T_{\rm g}({\rm PBAT})$ decreased with ETBP content indicating that ETBP promotes the interfacial compatibility of PLA and PBAT phases, by proposing that the epoxy groups of ETBP reacted with the terminal

carboxyl groups of PLA and/or PBAT, to enhance the interaction between the two phases.

Dong et al. [16] investigated phase morphology, mechanical properties, thermal properties and hydrolytic degradation of PLA as well as PBAT/PLA blends with and without chain extenders containing multi-epoxy groups. They improved both phase compatibility and mechanical properties. The chain-extending effect was confirmed by an increase in molecular weight. Elongations at break of PBAT/PLA blends increased from 200 % to 500 % without compromising tensile strengths.

Arruda et al. [17] investigated 60/40 % and 40/60 % PBAT/PLA blends after adding Joncryl® as CECL. Both PLA and PBAT showed improved thermal stability and increased complex viscosity. Furthermore, they found a preferential reactivity of the CECL with the PLA component. Pan et al. [18] studied melt compounded PBAT/PLA blends using MDI (4,4 methylene diphenyl diisocyanate) as CECL. Increasing MDI content led to higher yield and tensile strength, modulus and elongation at break associated with yielding deformation between the PLA and PBAT phase. Supplemented DSC, WAXS and SAXS studies suggested that amorphous PLA and PBAT can be well compatibilized with MDI due to an enhanced mobility of the amorphous phase. Therefore, crystallite dimensions are reduced in the entangled amorphous network.

Still, a synergy of how the compounding process with different CECLs can have on the process-induced properties of blown-film blends is an innovation presented in this thesis, that is lacking in the current literature.

1.3 Properties of modified PBAT/PLA blends after blown-film processing

The introduction of CECL to polymer blends changes their properties, which are determined by mostly standardized tests summarized in Table 2. As processes such as chain scission or cross-linking decrease and increase molecular weights of polymer chains, respectively, shifts of glass temperatures of the phases provide information in which phase the reactions took place.

Table 2: Standardized tests for polymer films.

Test procedure	Standard	Determined properties
Tensile test	ISO 527-3	Young's modulus E Tensile strength σ_{max} Strain at break ε_{break}
Tear test	ISO 6383-2	Rupture force F_{break} Tear resistance F_{tear}
Seal strength test	DIN 55529	Seal strength F_{seal}
Gel permeation chromatography (GPC)		Number molecular mass M_n , Weight molecular mass M_w
Sentmanat Extensional Rheometer (SER).		Elongational rheology
Dynamic mechanical analysis (DMA)	-	Storage modulus <i>E</i> ' Loss modulus <i>E</i> ''
Dynamic scanning calorimetry (DSC)	ISO 11357-3:2018	Glass temperature T_g Melting temperature T_m Melting enthalpy ΔH_m Crystallization temperature T_{cr} Crystallization enthalpy ΔH_{cr}
IR spectroscopy (FT-IR)	-	-
Scanning electron microscopy (SEM)	-	-

1.3.1 Tensile properties of modified PBAT/PLA blends

According to Arruda et al. [17] the mechanical behavior of neat PLA and neat PBAT is characterized by stress-strain curves and differs significantly, Figure 12. Furthermore, the effects of the introduction of 0.3 and 0.6 % Joncryl® can be seen. In particular, the elongation at break increased from around 400 % to 900 % for PBAT, whereas it decreased for PLA [17].

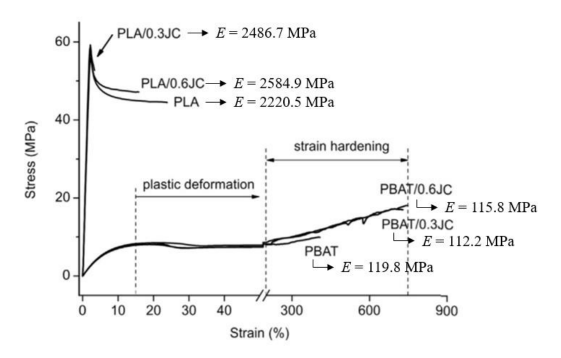


Figure 12: Stress-strain curves of neat PLA, neat PBAT and PBAT/PLA blend with different Joncryl® concentrations [17] with corresponding Young's modulus (E) values.

1.3.2 Tear properties of PBAT/PLA blends

The tear strength of blown films is a measure of their resistance against tearing forces [50-52]. Kim et al. [53] reported that blown films made by a 65/35 % PBAT/PLA blend with a thickness of 30 µm achieved an anisotropic tear resistance of 4.6 N mm⁻¹ in TD and 8.7 N mm⁻¹ in ED. The mixture ratio of PBAT/PLA blends has a significant effect as tear strengths increase with PBAT content, thus demonstrating that PBAT functions as an impact modifier for PLA.

Su et al. [48] have studied un-compatibilized PBAT/PLA blends with different contents of both polymers concerning tear strength properties, Table 3. The tear strength increases with PBAT content, showing that PBAT functions as an impact modifier for PLA.

Table 3: Tear propagation resistance of blown films in ED and TD directions [48].

PBAT/PLA	Tear strength ED (N/mm)	Tear strength TD (N/mm)
90/10	50.6 ± 0.9	Not possible to measure
80/20	48.5 ± 1.1	45.6 ± 2.1
70/30	27.6 ± 0.8	11.5 ± 0.6
60/40	8.8 ± 0.5	5.9 ± 0.3

Very little is known how this impact property is affected with different CECLs and the influence of different blown-film parameters to determine the most sustainable parameters for blown-film production.

1.3.3 Seal strength of PBAT/PLA blends

Polymer films are often sealed or welded if used for packaging. Therefore, the seal strength is an important property and quality assurance feature that affirms package integrity during shelf life. The seal strength test is performed using a tensile testing machine that measures a force-displacement curve. The seal force is given by the plateau force, Figure 13.

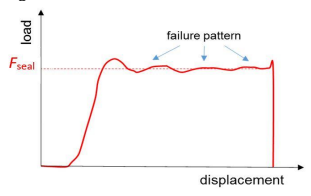


Figure 13: Scheme of load displacement curve showing the determination of the seal strength.

The force oscillations of the plateau contain information on the mechanisms determining seal failure.

In the sealing process, two hot jaws heat the two films to a partially molten state and press them to one another for a given time until the seal is formed [54-55]. Relevant process parameters are sealing temperature, dwell time, and pressure that affect the achievable seal strength [56]. Amorphous PLA exhibits good sealing performance with a high seal strength of 10 N as it exhibits sufficient chain mobility at the surface leading to a better inter-diffusion of molecules and adhesion under standard sealing temperatures of 140 °C. Semi-crystalline PLA is not sealable under standard sealing temperatures because this requires some melting of crystallites [57]. For economic reasons high packaging speeds are desired which require short sealing times. Therefore, it is an issue how the PBAT/PLA ratio and CECL affect the seal quality.

1.3.4 Rheological properties of modified PBAT/PLA blends

For film blowing, the polymer melt requires high melt strength [47-48]. Compared to PE, PLA and PBAT are linear polymers with rather low molecular weight, and thus, low concentration of entanglements being responsible for melt strength under elongational flow.

Chain extending and crosslinking evoke a complex rheological behavior such as the appearance of strain hardening caused by more entanglements and the formation of covalent bonds between PLA and PBAT [58]. The application of chain extenders may result in the formation of long chain branching [59].

Reactive CECL may enhance extensional viscosities and may cause strain hardening [60-63]. Al-Itry et al. [63] investigated the structure modification of physical properties and morphology of PBAT/PLA blends with CECL Joncryl® including their behavior under elongational flow. They determined the elongational viscosity η_E using the Cogswell method exploiting the relationship between elongational viscosity and die pressure [62]. Elongational and shear viscosities increase with CECL content, Figure 13 indicating that reactions of CECL increased the mean molecular weight, and thus, the concentration of entanglements. Therefore, the improvement of the blown film extrusion capability can be expected.

Both PLA and PBAT exhibit no strain-hardening for strain rates less than $1~\rm s^{-1}$. This corresponds to the strain-softening phenomena of linear polymers discussed e.g. by Liu et al. [60]. It has been seen that the presence of free volume within polymer chains may hinder the formation of entanglements through reduced contact surface and ineffective overlapping. Linear chains are also free of branching and can easily slip from one another. The absence of strain hardening of both PLA and PBAT are the reasons why they are not favorable polymers for blown films.

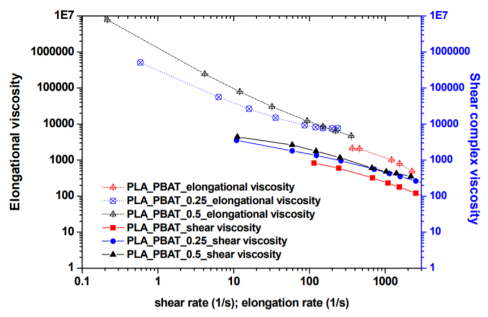


Figure 13: Effect of CECL content on rate dependent elongational and shear viscosities of a non-compatibilized and compatibilized PBAT/PLA blend at 180 °C [63].

Chain extending and crosslinking evoke a complex rheological behavior such as the appearance of strain hardening caused by more entanglements and the formation of covalent bonds between PLA and PBAT [63]. The application of chain extenders may result in the formation of long chain branching [65-68] Pure linear PLA naturally does not exhibit strain hardening [69], the PLA/PBAT blends lack this phenomenon for extension rates lower than 1 s⁻¹ where the process of disentanglement prevails [67-69].

As evidenced, the measurement of elongational viscosity does not range to trivial operations. Ingenious devices proposed by Meissner [70-71] and Münstedt [72] in the 1980s are no longer in use. At present the field is dominated by the Sentmanat Extensional Rheometer Universal Testing Platform (SER) [71-73]. The big advantage of this apparatus is its small size enabling its housing practically in any standard rotational rheometer, where its easy manipulation makes its usage comfortable. This evaluation is made on this thesis proving insight of the synergy of new CECLs with the blown-film processing.

1.3.5 Dynamic mechanical analysis of modified PBAT/PLA blends

The DMA allows for precise measuring of glass transition temperatures $T_{\rm g}$, in particular, it provides two different $T_{\rm g}$ due to the maximum of E" (= static $T_{\rm g}$) and the maximum of $tan \delta$ (= dynamic $T_{\rm g}$). The characterization of thermal mechanical properties in terms of temperature dependent storage modulus E' and loss modulus E'' is crucial to understand the effects due to blending, mixture ratio and CECL. In Figure 14, storage moduli curves of neat PLA, neat PBAT and a 50/50 % PBAT/PLA blend are shown. Furthermore, the effects of four CECL concentrations on storage moduli of the blend are shown for MDI (4,4 methylene diphenyl diisocyanate) [18].

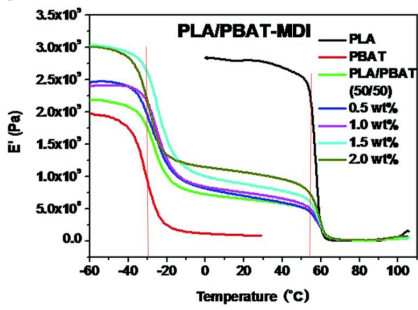


Figure 14: Temperature dependent storage moduli (E') of PBAT, PLA and a 50/50 PBAT/PLA blend modified with CECL [18].

Dynamic mechanical analysis measurements indicate enhancement of compatibility of the PBAT/PLA-MDI blends. The pure PLA displays a high E' at about 3.0 GPa, which remains in the same order up to 55 °C, while neat PBAT shows a smoother drop in E' at about -30 °C. For the PBAT/PLA-MDI blends, E' drops at about -30 °C, when the fraction of MDI becomes higher. This is also seen on the 50/50 % blends with the increasing of MDI percentage; the higher the amount of modifier, the higher is the shift in $T_{\rm g}$, and also higher is the E'. The varied morphology, coupled with chemical interaction between the functional groups of the two polyesters, may affect mobility of the phases, and result in a shift of glass transition temperature with respect to the expected values [58].

1.3.6 Thermal properties of PBAT/PLA blends

As both PBAT and PLA are in principle semi-crystalline polymers, it is interesting how reactive processing affects the phase structure and, thus, the transition temperatures and corresponding enthalpies. Figure 15 shows the effects content ratio on DSC traces of PBAT/PLA blends with indication of glass and melt temperatures.

The $T_{\rm g}$ of neat PLA is around 60 °C. The $T_{\rm g}$ of PBAT is around -30 °C. The melt peak temperatures of PLA and PBAT are around 150 °C and 123 °C, respectively. $T_{\rm g}$ and $T_{\rm m}$ of PBAT/PLA blends remain at values corresponding to pure PBAT and PLA indicating a two-phase structure [59-60]. PLA shows no melting peak, and 10/90 % PBAT/PLA has pronounced crystallization and melting peaks meaning that PBAT acts as a nucleating agent. Increasing PLA content reduces the peak areas more than expected with respect to the content. At 50/50 % PBAT/PLA, crystallinity has almost vanished.

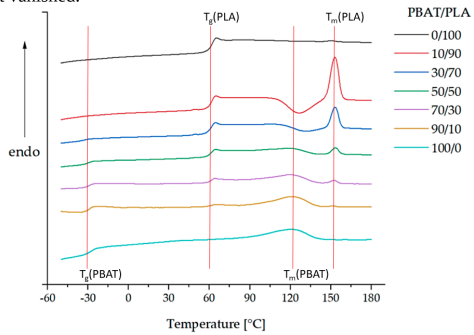


Figure 15: DSC Analysis of neat PBAT and neat PLA films [48].

Abdelwahab et al. [76] reported changes in T_g in PBAT/PLA blends with 20 wt.% organosolv lignin (OL) and different Joncryl[®] ADR loading.

The $T_{\rm g}$ of PLA in the blend remained at 59 to 60 °C indicating that there is very low interaction between the two polymers, because there are always some interactions. The addition of OL shifts $T_{\rm g}$ of PLA to 56.7 °C.

The addition of CECL slightly increased the $T_{\rm g}$ of the PLA blends (as shown on Figure 16) indicating the decrease of the chain mobility due to the increase of the molecular weight by branching or crosslinking and possibly due to enhancement of the interface.

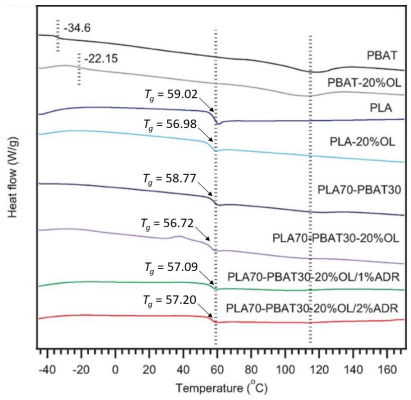


Figure 16: DSC traces of PLA, PBAT and 30/70 PBAT/PLA blend without and with 20 wt.% organosolv lignin (OL) containing 0, 1 and 2 % CECL (Joncryl® ADR) [76].

IR spectroscopy may provide information of structural changes due to chemical reaction of CECL. Wu et al. [77] have investigated how Joncryl® (ADR) and dicumyl peroxide (DCP) modify the spectra of an 80/20 % PBAT/PLA blends, Figure 17, and found the characteristic bands at 1714 cm⁻¹ for PBAT and 1752 cm⁻¹ for PLA due to the –C=O stretching vibration in PBAT and PLA. The bands at 849, 908 and 1252 cm⁻¹ were assigned to the –C–O stretching vibration of the ADR epoxy group.

In addition, there were no epoxy group characteristic bands in the PBAT/PLA/ADR (in one step mixing of all components) and (PBAT/ADR)/PLA

(first melt blending of PBAT and ADR for 4 min followed by mixing of PLA for another 4 min) spectra which indicate terminal carboxyl and hydroxyl groups of PLA and PBAT having reacted with the epoxy groups [15]. FTIR spectra confirmed that the epoxy group of ADR can react with the hydroxyl or carboxyl group of PLA and PBAT in reactive blending, because there were no similar epoxy group characteristic peaks in the PBAT/PLA/ADR and (PBAT/ADR)/PLA spectra which indicated the terminal carboxyl and hydroxyl groups of PLA and PBAT could react with the epoxy groups. The spectra of PBAT/PLA/DCP (in one step mixing of all components) and (PBAT/DCP)/PLA (first melt blending of PBAT and ADR for 4 min followed by mixing of PLA for another 4 min) are similar to those of PBAT/PLA blends.

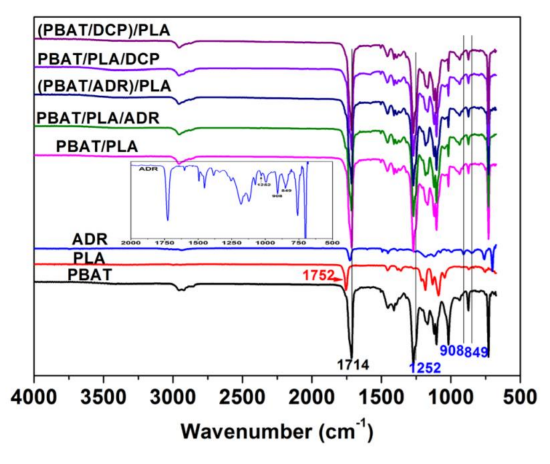


Figure 17: FTIR spectra of PBAT, PLA, ADR and both unmodified and modified 80/20 % PBAT/PLA blends [77].

1.3.7 Morphological investigations of PBAT/PLA blends

1.3.7.1. Structure formation of blends

The properties of immiscible blends depend on mixture ratio of blend components and processing induced morphology. With increasing PBAT content, the equilibrium morphology of PBAT/PLA blends is changed, Figure 18 [78].

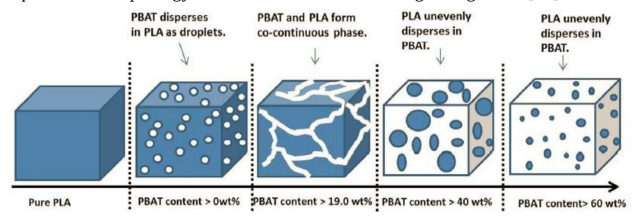


Figure 18: Schematic diagram of phase structure as a function of composition [78].

Under elongation and/or shear flow spherical droplets may deform to fibrils being solidified in that shape if rapidly cooled. The surface is more irregular with a two-phase morphology of PLA and PBAT as hard and soft phase, respectively. In liquids consisting of immiscible components the difference of surface tensions tries to minimize the surface energy of each phase. This mechanism leads to a phase structure showing spherical inclusions. Compatibilizer reduces the difference of surface tensions, and for miscible components only the external surface is left.

The fracture surfaces of the PBAT/PLA blends exhibit a brittle failure character as fracturing took place at -196 °C, Figure 19. The disperse phase is observed as spherical particles or pores on the fracture surface, Figure 21, and the arrows represent the main morphological changes. Furthermore, the mean diameters of the disperse phase depend on the PBAT/PLA ratio as they increase from 30 to 50 % PBAT content [79-82], Figure 19 c-e.

PBAT has longer repeating units and more flexibility compared to PLA, thus, low interfacial adhesion and phase separation is observed [83-84], because only van der Walls forces are responsible for adhesion and no covalent bonds in form of tie molecules are present. The additions of CECL are to generate such tie molecules bonding.

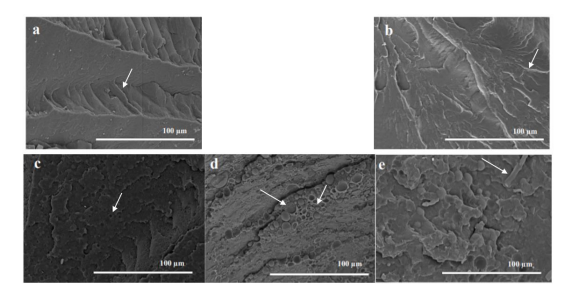


Figure 19: SEM micrographs of cryogenic fracture surfaces of PBAT/PLA: (a) PLA 100 %, (b) PBAT 100 %, (c) PBAT/PLA 30/70 % (d) PBAT/PLA 40/60 %, (e) PBAT/PLA 50/50 % [79].

1.3.7.1. Processing effects on structure and properties

A few investigations deal with CECL in PBAT/PLA blends [13-17]. Fracture surfaces of films in ED and TD of 60/40 and 40/60 PBAT/PLA blends show the effects of film blowing and addition of CECL (0.3 and 0.6 % Joncryl®, respectively) [17], Figures 20 and 21. All blends exhibit a skin core structure with smaller dimensions of the dispersed phase closer to skin than in core. This is attributed to the fact that close to the skin the dispersed phase is stretched to longer and thinner fibrils, which then decay to smaller droplets. Furthermore, the polymer melt is colder than in the core, thus, having a higher melt viscosity avoiding relaxation to coarser structures of the dispersed phase.

The fracture surfaces of the 40/60 % PBAT/PLA blends, Figure 20, show that the dispersed phase is elongated to fibrils with aspect ratios exceeding 5 in ED due to a high DR, whereas it is deformed to ellipsoids in TD with aspect ratios between 2 and 4. With increasing CECL content the aspect ratios become smaller indicating that the reactions of CECL have increased the melt viscosity. The dispersed phase cannot be oriented furthermore. Without CECL, the fracture surfaces exhibit a brittle character that seems to become slightly more ductile by adding CECL. Both the reduced aspect ratios of the dispersed phases, and the slightly increased ductility show that CECL additions below 1 % are a tool to adjust properties of PBAT/PLA blends. The fracture surfaces of the 60/40 PBAT/PLA films differ significantly in ED and TD, Figure 21. In ED, a ribbon structure occurs indicating that the dispersed phase has been deformed. This ribbon structure is less pronounced if CECL is added.

This also indicates that CECL addition increases the melt viscosity. These compositions present PBAT as the dispersed phase. The characteristic morphology in these blends is ribbon- or sheet-like aligned to drawing direction. A structure similar to the skin-core type can be noticed, in which the dispersed phase is coarse in the middle of the thickness (core region) of the film but, as the dispersed phase gets closer to the film surface (skin region), the morphology presents a finer, fibrillar structure. In TD the aspect ratio of the disperse phases exceeds that of the 60/40 % PBAT/PLA films, in particular in the core where it reaches aspect ratios around 10. Furthermore, the interface adhesion seems to be similar for all CECL contents. The cross-sectional change of the dimensions can be accounted for the shear rate gradients occurring in the die. In the high shear rate areas close to the skin, the spherical inclusions of the dispersed phase are reduced in diameter, whereas they remain coarse in the core. After exiting the die for film blowing, the flows become elongational both in ED and TD due to *DR* and *BUR*.

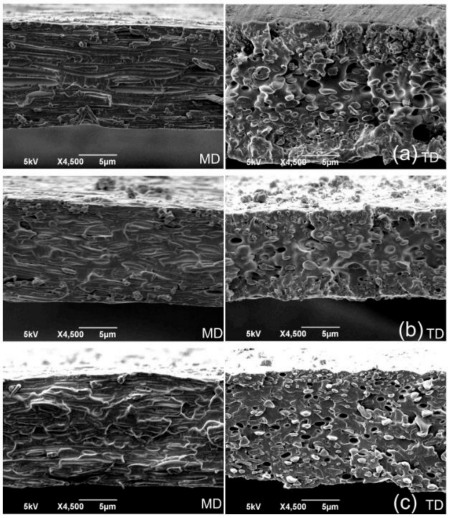


Figure 20: Fracture surface of a PBAT/PLA 60/40 % blend in SEM in ED and TD - (a) without CECL, (b) with CECL 0.3 % Joncryl[®], (c) with CECL 0.6 % Joncryl[®] [17].

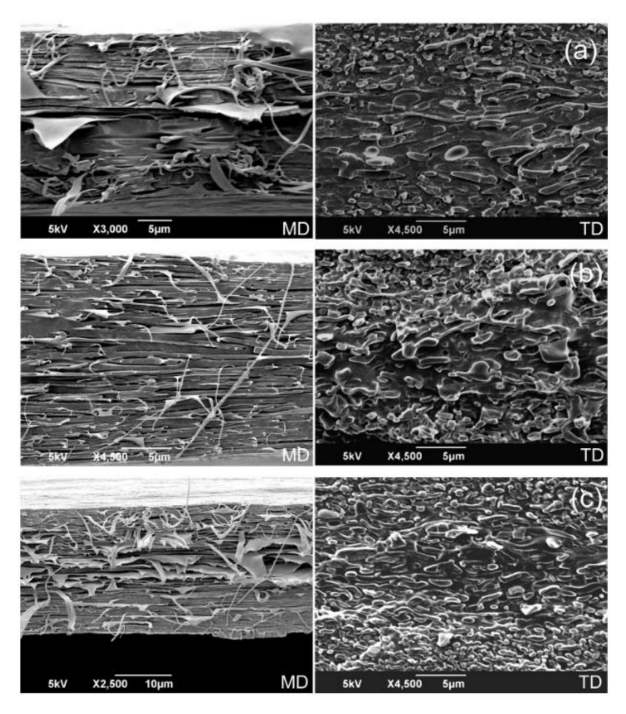


Figure 21: Fracture surface of a PBAT/PLA 40/60 % blend in SEM in ED and TD - (a) without CECL, (b) with CECL 0.3 % Joncryl[®], (c) with CECL 0.6 % Joncryl[®] [17].

1.4 Disintegration process of PBAT/PLA blends

The advantage of biodegradable polymers is that they "vanish" if they are exposed to environmental conditions after becoming waste. One distinguishes different environmental conditions (compost, soil, water, different temperatures and under aerobic or anaerobic influences) under which biopolymers may degrade. According to the IUPAC¹ definitions, biodegradation means degradation caused by enzymatic processes resulting from the action of living cells [85-87]. Lucas et al. [85] classified the process in three main steps:

- **biodeterioration** is materials fragmentation in tiny pieces due to decomposers such as microorganisms and/or abiotic factors;
- **depolymerization and biofragmentation** is molecular mass reduction due to chain cleavages;
- mineralization and biomass production mean uptake of small molecules into the living cells and therein conversion to biomass, storage vesicles and metabolites. This finally leads to total mineralization, which is the formation of simple molecules such as H₂O and CO₂.

The disintegration of polymers requires chain cleavage which can be initiated by hydrolysis, thermal or mechanical activation as well as oxidation processes, and enzymatic processes. For polyesters, such as PLA, PBAT and their blends, hydrolysis either purely chemical or enzymatic mediated is a dominantly occurring process [85-88]. Tolga et al. [88] have studied the time dependent disintegration processes of three biopolymer blends under industrial composting conditions. They found that degradation started after an incubation time of 5 to 6 weeks, whereas the two PBS/PLA blends started after 4 weeks.

After an incubation period in which PLA is subjected to hydrolytic degradation until the polymer segment have reached a length enabling microorganisms to attach. This step is characterized by a slight weight increase due to water absorption. In the next step, microorganisms will deteriorate the segments and produce biomass, water and CO₂. Once started, hydrolytic degradation in PLA proceeds faster in the inner molecular structure than on the outer surface due to the fact that water penetrates the material and generates chain cleavages in amorphous regions at the ester bonds. Once initiated, the process accelerates as a consequence of a high amount of carboxylic end groups being formed, which decreases the pH in the inner core of the PLA material. Additionally, an erosion process on the specimen's surfaces in the present biotic environment accelerated by enzymatic action is also to be expected as being already shown in the literature [89-90].

¹ IUPAC - International Union of Pure and Applied Chemistry

2. AIM OF THE WORK

The global plastic waste problems positively affected the development of biopolymers and other sustainable materials leading to the substitution of traditional packaging plastics with biodegradable polymers. However, for more demanding engineering applications, their mechanical properties must be enhanced without compromising their biodegradability. Many biopolymers are combined with other polymers to balance the properties required for specific applications. Compounding is used to make a biopolymer fit for specific applications by blending it with other polyesters and adding chain extenders cross-linkers to provide optimized properties.

Currently little is known about how multi-functional chain extenders cross-linkers (CECL) affect the properties and morphologies of PBAT-PLA blends. Therefore, the objective of this study is to investigate the effects of four CECLs on the morphology of PBAT-PLA compounds to optimize processability and usage performance.

The largest market in the plastics industry is the packaging segment with more than 40 % of plastics demand in Europe. Blown film extrusion is the most important industrial manufacturing process of polymeric films.

As for all polymer processing techniques, blown film processing parameters e.g. extrusion temperature, blow-up ratio, and draw ratio affect the thermo-viscoelastic properties and the morphological structure of the film, have been collected. Within this work, the process-induced thermo-viscoelastic and mechanical performance is to be investigated using dynamic mechanical analysis, tensile properties, and differential scanning calorimetry. To further obtain a better insight, the morphological structure will be examined by scanning electron microscopy and elongational rheology. Moreover, the impact of the biodegradation (disintegration effect) is also studied to see which effect CECL can have on the rate of disintegration in blown films.

The morphologies are interpreted concerning structure formation during film blowing with a blow-up ratio and a draw ratio, where the temperature and CECL are dependent on melt viscosities of blends of PBAT and PLA. After extrusion, elongation flows stretch and orient the melt, at the beginning in the transverse direction, due to blowing, and subsequently in the extrusion direction, due to the drawing. Rate-dependent shear and elongation viscosity measurements would provide a deeper understanding of the effects that the introduced CECL has on melt viscosities and the process-generated film morphologies.

A better understanding of the introduction of the CECLs in PBAT PLA blends provided in the thesis, Figure 22, will lead to predictions of the CECL effects on blown film extrusion and film properties.

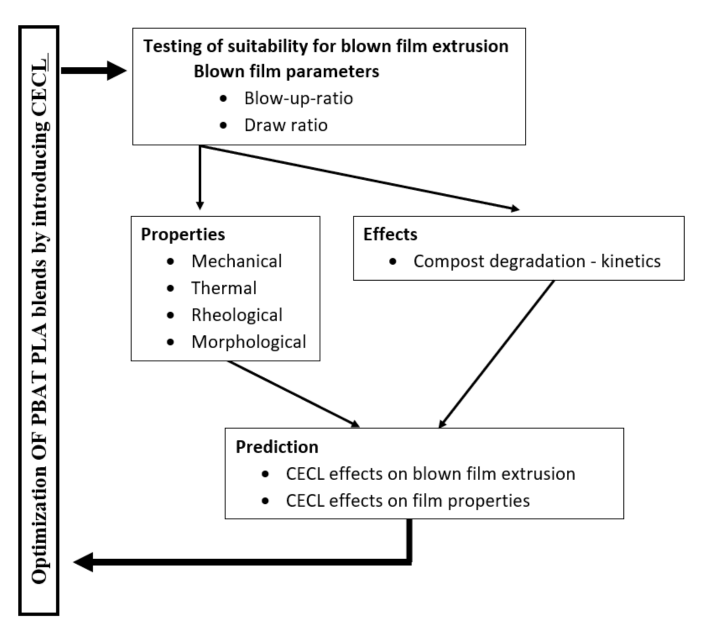


Figure 22: Flow chart of the methodology and techniques employed in this PhD thesis.

3. MATERIALS AND METHODS

Four CECL (1 wt. %) were compounded into the reference PBAT-PLA (REF) compound named M·VERA® B5029 [91] from BIO-FED, a branch of AKRO-PLASTIC GmbH, Germany developed before this dissertation, and which is now used for packaging (fruit and vegetable bags, labels) and agricultural applications:

Table 4 Information about the CECL used with the introduction of chemical name, trade name and manufacturer.

	Chemical name	Trade name	Manufacturer
V1	tris(2,4-di-tert-	Songnox TM	Songwon Industrial
V 1	butylphenyl)phosphite	1680	Co [92]
V2	1.2 mbanylanahisayagalina	1,3-PBO	Examile [02]
V Z	1,3-phenylenebisoxazoline	powder	Evonik [93]
V3	aromatic polycarbodiimide	Stabaxol® P110	Lanxess [94]
T 7.4	poly(4,4-	Carbodilite TM	Nicolaimha [05]
V4	poly(4,4-dicyclohexylmethanecarbodiimide)	HMV-15CA	Nisshinbo [95]

All ingredients were uniformly mixed using a Mixaco CM 150-D (Mixaco Maschinenbau, Germany) and compounded in a twin-screw extruder (FEL 26 MTS, Feddem GmbH, Germany) with a diameter of 32 mm and a L/D of 26, screw speed of 260 rpm and output rate of 20 kg h⁻¹. The films were blown using an LF-400 (Labtech Engineering Company, Thailand) machine with a single screw having a diameter of 25 mm and an L/D of 30, extrusion temperature of 165 °C, and blow-up-ratio (BUR) of 1:2.5 for both 25 μm and 100 μm films. From an extrusion gap of 0.8 mm, the draw ratio was estimated from 12 to 14 for 25 μm films and 3 to 4 for 100 μm films. The extrusion pressures were 240 bar for the REF blend, 290 bar (V1), 159 bar (V2), 230 bar (V4) and 313 bar (V4). Before testing, all films were stored for 24 h at 23 °C/50 % r.h. Further films with different BUR were made and on Table 5 is presented the thickness with standard deviation (STD) and draw ratio (DR).

Table 5 Mean thicknesses of films of REF, V1 to V4 with the following standard deviation (STD), blow-up ratio (BUR) and draw-ratio (DR).

	BUI	BUR 2.5			BUR 3.5				
	Thickness (µm)	STD	DR	Thickness (µm)	STD	DR	Thickness (µm)	STD	DR
REF	26.0	1.0	20.5	25.8	0.7	12.4	26.0	1.0	8.8
V1	27.0	1.0	19.8	30.5	0.5	10.5	26.4	0.5	8.7

V2	26.6	0.9	20.1	26.3	0.5	12.2	26.0	0.7	8.8
V 3	27.4	0.5	19.5	27.3	0.6	11.7	26.6	0.5	8.6
V4	25.2	0.4	21.2	29.9	0.8	10.7	26.2	0.4	8.7

Tensile properties

Young's modulus, tensile strength, and fracture strain were determined according to ISO 527-3 using a tensile testing machine (2.5 kN Zwicki, Zwick Roell, Germany) at 23 °C/50 % r.h. with a crosshead speed of 200 mm min⁻¹. The specimens having 170 mm length and 15 mm width were examined at least 10 times each in extrusion direction (ED) and transverse direction (TD).

Tear properties

The tear strength of blown films was determined according to ISO 6383-2. Samples were cut from a blown film both in ED and TD with $L = 63 \pm 0.5$ mm, $W = 76 \pm 0.5$ mm, placed in the test device, and a cut of defined length was set by a knife. Then the pendulum is released, and a tear is generated in the film between the mobile and stationary clamp, as demonstrated in Figure 23.

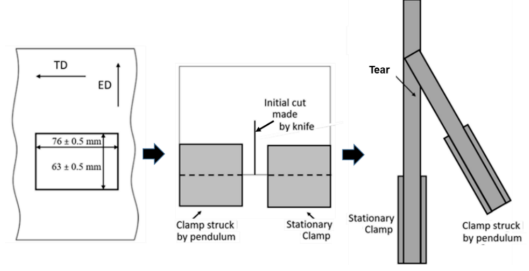


Figure 23: Scheme of Elmendorf tear test [51].

Seal strength

The seal strength tests were performed according to DIN 55529 using a tensile testing machine (2.5 kN Zwicki, Zwick Roell, Germany) with a clamp distance of (40 ± 5) mm and a trigger angle of 90°, as seen in Figure 24. The seal strength is given by the plateau force F_{seal} . Samples ($200 \times 200 \text{ mm}$) of 25 µm films were sealed at 90° concerning ED using a sealer (SGPE 3000, Willi Kopp e.K., Germany) at 140 °C and 130 N for 3 s with a seal width of 10 mm. Then, stripes having a width of 15 mm were cut out, and stored for 24 h at 23 °C/50 % r.h.

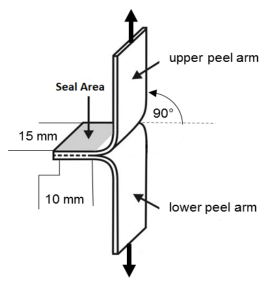


Figure 24: T-test sample to measure seal strength acc. to [DIN 55529].

Dynamic Mechanical Analysis (DMA)

Storage modulus E' and loss modulus E'' were measured in tension mode (displacement controlled) using a DMA 242 E *Artemis* (Netzsch, Germany). The DMA tensile mode was calibrated in accordance with the manufacturer's procedure. Samples (7×5.9) mm were punched out of $100 \, \mu m$ thick films in ED and TD directions. 25 μm films could not be measured due to low forces. Measurements were performed at a frequency of 1 Hz, a heating rate of 2 K min⁻¹, a deformation amplitude of $20 \, \mu m$, and a preload of $0.2 \, N$ over a temperature range of $-70 \, ^{\circ}C$ to $80 \, ^{\circ}C$.

Sentmanat Extensional Rheometer (SER)

Measurements were carried out using an SER Universal Testing Platform (model SER-HV-P01) that was applied with the Anton Paar MCR501 rotational rheometer host system (Graz, Austria) equipped with the convection heated measuring chamber CTD 450 (Graz, Austria).

Four extensional rates 0.0316, 0.1, 0.316, and 1 s⁻¹ were adjusted via the rotational speed of the drums at temperatures ranging from 135 to 170 °C (given by the melting temperatures of PBAT and PLA, respectively). The measurements were carried out at least in triplicate. Rectangular samples of length 18 mm and widths of 9 and 12 mm, respectively were taken out of blown films having thicknesses of approximately 0.1 mm, Table 6, both in extrusion direction (ED, 90°) and in transverse direction (TD, 0°), see Figure 25. Two widths were chosen to check the homogeneity of the blown films. Before testing, the films were conditioned for 24 h at 23 °C/50 % relative humidity.

Table 6 Mean thicknesses of films of REF, V1 to V4 with the following standard deviation (STD), blow-up-ratio (BUR) and draw-ratio (DR).

	Thickness (µm)	STD	DR
REF	98.8	1.3	3.24
V1	101.0	1.6	3.17
V2	100.2	1.3	3.19
V 3	100.4	1.7	2.19
V4	101.4	1.7	2.16

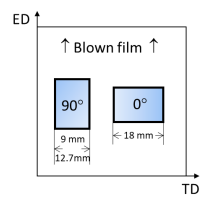


Figure 25. Sample taking and notation.

Prior to the measurements it was necessary to determine the temperature under which the SER device is possible to use. Based on the range between 135 °C and 170 °C it was found that the optimal temperature with respect to sagging is 150 °C. Only in the case of REF, the temperature was lowered to 140 °C, due to intensive sagging at 150 °C. The dimensions of the identical samples were thickness - 0.1 mm, length - 18 mm, width - 12.7 and 9 mm. No sagging corrections were considered.

Fourier transform infrared spectroscopy (FTIR)

Fourier Transform Infrared Spectroscopy (FTIR) was used to identify structural changes due to chemical reactions of the CECL with PBAT and PLA. IR-spectra were recorded in the wavenumber ranging from 2000 to 600 cm⁻¹ using an FTIR Microscope System (Perkin Elmer Spectrum Spotlight 200, USA) with Attenuated Total Reflectance (ATR) in continuous scan mode, with a spectral resolution of 16 cm⁻¹ and 15 scans averages per spectrum.

Gel permeation chromatography (GPC)

The granules of pure PBAT and PLA as well as film samples of REF, V1 to V4 were dissolved in THF and filtered using PTFE filters with a mesh of 0.45 μm into

vials of 2 ml. All samples showed turbidity due to a filler content of 11 %. The sample concentration in each vial was 2.5 g/L. The GPC measurements were conducted by a Waters HPLC system equipped with a Waters model e2695 and Waters model 2414 differential refractometer (Waters Corporation, Massachusetts United States of America). The GPC system was calibrated with polystyrene standards having molecular masses of 580; 10,440; 38,640; 132,900; 492,500 and 990,500 g/mol.

The measuring conditions were:

- Series of gel-mixed bed columns: PL gel MIXED-A (300 x 7.5 mm, 20 μ m) + PL gel MIXED-B (300 x 7.5 mm, 10 μ m) + PL gel MIXED-D (300 x 75 mm, 5 μ m)
- Mobile phase: tetrahydrofuran (THF) stabilized with butylated hydroxytoluene
 - Temperature: 40 °C
 - Injection volume: 100 μL
 - Flow rate: 1 ml/min
 - Detector: refractive index detector (RI)

All data processing was carried out using Empower 3 software (Version FR 3).

Differential scanning calorimetry (DSC)

DSC experiments were performed using a DSC Diamond (Mettler Toledo, Switzerland) for granules and a DSC 214 Polyma (Netzsch Gerätebau GmbH, Germany) for films in standard Al pans in three steps -1^{st} heating, cooling, 2^{nd} heating. The measuring conditions were: sample weight (m_s) (6 ± 1) mg, starting temperature (T_{start}) 0 °C, end temperature (T_{end}) 200 °C, heating/cooling rate 10 K min⁻¹, and repetition (n) was ≥ 2 .

The thermal quantities as glass temperature of hard segments $T_{g,hs}$, melting temperature of PBAT T_{ml} , melting temperature of PLA T_{m2} , melting enthalpy of PBAT ΔH_{ml} , melting enthalpy of PLA ΔH_{m2} , crystallization temperature T_{cr} and crystallization enthalpy ΔH_{cr} were evaluated from DSC curves according to ISO 11357-3:2018.

Scanning electron microscopy (SEM)

Films with 25 µm thickness were fractured under cryogenic conditions in ED and TD directions using liquid nitrogen and subsequently sputtered with gold at 20 mA for 3x30 s. The fracture surfaces of all samples were investigated using a field-emission Scanning Electron Microscope SEM (JSM-7200F, Jeol, Japan) at acceleration voltage 5.0 kV and amplifications of 3.000 and 10.000, both in ED and TD, Figure 26.

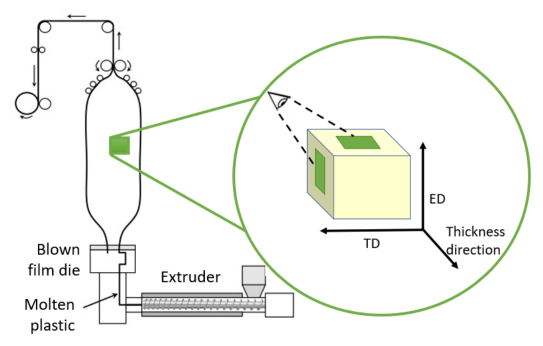


Figure 26. Sample orientation with respect to ED, TD and thickness directions; ED shows the TD-thickness direction-plain and TD shows the ED-thickness direction-plaine.

Disintegration according to ISO 16929:2018

The samples were stabilized at 23 °C for 24 hours after extrusion. Films were cut to tensile test stripes having the dimensions 170 x 15 mm and weighed. Trays with at least 3 liters of volume were filled with compost (Plantiflor Pro Natur torffrei, Dortmund, Germany) with 3 subsequent layers: a bottom layer of compost (thickness: 5 cm), a layer of individualized test specimens, and a top layer of compost (thickness > 5 cm). To prevent cross-contamination, differently colored trays were used for REF, V1 to V4 and stored in an oven at 30 °C or 60 °C, respectively, with 60 % relative humidity to be close to industrial composting conditions described by ISO 16926:2018. The storage times at 30 °C were 168, 336, 569, 672, and 1344 hours; at 60 °C they were 24, 48, 72, 96, 120, 144, and 168 hours. As V3 and V4 hardly showed visual disintegration they were also subjected to compost storage at 60 °C for 336 and 672 hours. To follow ISO 16926:2018 pH values were determined when the samples were removed and cleaned.

Determination of mass change after disintegration in compost

The sample masses after compost exposure were measured using the balance Sartorius BP221S (Sartorius AG, Göttingen, Germany) having a measuring accuracy of 0.1 mg. The determined masses showed scatter in the range of 3 to 7 % due to sticking compost particles, adsorbed and absorbed disintegration substances, and loss of microplastic particles during cleaning in cases of severe film disintegration.

For a better comparison of mass changes of REF, V1 to V4, the data are converted to relative masses (please see Appendix A for detailed formulation).

Melt Volume Rate (MVR)

MVR was measured using a melt flow indexer (MI-3, Göttfert, Germany) according to ISO 1133 at 190 °C with a weight of 2.16 kg. Prior to the measurements, the granules were dried in an IR-oven (HB43-S, Mettler Toledo, Switzerland) for 20 minutes at 110 °C.

Thermal aging

As cross-linking or chain reactions can occur at elevated temperatures, to investigate the influence of CECL reactions, films were conditioned at 50 $^{\circ}$ C for 1 and 2 months. After this thermal aging, films were again stabilized for 24 h at 23 $^{\circ}$ C/50 % r.h. Subsequently, mechanical testing was performed.

4. Discussion of the Results

The presented results are also summarized in the following publications:

I The Effects of Chain-Extending Cross-Linkers on the Mechanical and Thermal Properties of Poly(butylene adipate terephthalate)/Poly(lactic acid) Blown Films.

J. V. C. Azevedo, E. R. Dorp, B. Hausnerová, B. Möginger; *Polymers*, 2021, Volume 13, 3092.

Paper I

II Process-induced morphology of poly(butylene adipate terephthalate)/poly(lactic acid) blown extrusion films modified with chain-extending cross-linkers.

J. V. C. Azevedo, E. R. Dorp, R. Grimmig, B. Hausnerová, B. Möginger; *Polymers*, 2022, Volume 14, 1939.

Paper II

III Effect of Chain Extending Cross-Linkers on the Disintegration Behavior of Composted PBAT/PLA Blown Films.

J. V. C. Azevedo, B. Hausnerová, B. Möginger, T. Šopík; International Journal of Molecular Sciences, 2023, Volume 25(5), 4525. Paper III

IV Improvement of elongational behavior of poly(lactic acid)/poly(butylene adipate terephthalate) blends in blown film extrusion using chain extending cross-linkers.

P. Filip, B. Hausnerová, B. Möginger, J. V. C. Azevedo; Submitted to *International Journal of Biological Macromolecules*, November 2023

Paper IV

V Blow molding-induced structure and performance of renewable PLA/PBAT packaging films modified with chain-extending cross-linkers.

J. V. C. Azevedo, E. R. Dorp, C. Kössler, C. Thomser, T. Šopík, B. Möginger B. Hausnerová;

Submitted to *Journal of Industrial and Engineering Chemistry*, November 2023

Paper V

4.1 Behavior under deformation

4.1.1 Mechanical properties

Processing influences material structure and properties. Due to the blown film processing, the mechanical properties are anisotropic. In order to evaluate the effect of chain-extenders cross-linkers (CECL) on the process-induced mechanical properties relationship, Young's modulus E, tensile strength σ_{max} , elongation at break $\varepsilon_{\text{break}}$, tear resistance F_{tear} in extrusion direction (ED) and transverse direction (TD) and seal strength F_{seal} as well as the melt volume rate (MVR) were determined. Tests were performed on the unmodified blend (REF) after conditioning for 24 hours for stabilizing (initial) and after thermal aging for 1 and 2 months at 50 °C, Table 7.

Initial E and F_{tear} in ED clearly show higher values than in TD, whereas the elongation at break coincides with standard deviations (STD). This can be explained by the molecular orientation in ED caused by film blowing resulting in a flow-induced stiffening [87]. Furthermore, melt volume rate (MVR) increases by 20 % and 40 % after 1 and 2 months, respectively. This reduction of the mechanical properties and the increase in flowability after thermal aging indicates chain scission or degradation of polymer chains.

Table 7 Mechanical properties and MVR of unmodified REF blend in extrusion direction (ED) and transverse direction (TD): initial and aged for 1 and 2 months at 50 °C. (Paper I)

Property	Initial	1 month	2 months
$E_{\rm ED}$ (MPa)	457±36	437±37	363±48
$E_{\mathrm{TD}}\left(\mathrm{MPa}\right)$	181±22	238±10	197±12
$\sigma_{ m max,ED}(m MPa)$	27.8±3.1	20.4±2.6	21.2±1.5
$\sigma_{ m max,TD}(m MPa)$	28.8 ± 3.1	21.8 ± 2.2	26.1±1.8
Ebreak, ED (%)	400±48	310±39	410±26
$\mathcal{E}_{\mathrm{break,TD}}\left(\% ight)$	450 ± 27	390±20	500±20
$F_{\text{tear, ED}}$ (N mm ⁻¹)	107±3	104±33	80±25
$F_{\text{tear, TD}}$ (N mm ⁻¹)	54±17	57±18	51±16
$F_{\rm seal}$ (N mm ⁻¹)	7.0 ± 0.3	5.9 ± 0.4	4.8 ± 0.6
MVR (cm ³ 10 min ⁻¹)	4.0±0.1	4.7±0.1	5.5±0.2

Table 8 shows the properties of introducing the CECL V1 to V4. While E of V1 to V3 show only slight differences compared to the initial, V4 shows a clear increase of up to 40 % in both ED and TD. Additionally, σ_{max} , ED and TD, for V1 and V4 show only a slight increase, and σ_{max} for V2 and V3 show only a slight decrease,

whereas the $\varepsilon_{\text{break},ED}$ reveals a clear decrease in ED with similar values in TD. This can be explained by the combination of the effect of various CECLs and the increasing orientation causing a stiffening effect during tensile testing.

The tear resistance only shows a minor increase in ED and a little decrease in TD. Furthermore, the seal strength for V1 decreases dramatically, whereas for V2 to V4, it depicts a relevant increase. These are important for packaging industries' important mechanical properties, which show the process-induced orientation in ED and TD proved to provide mechanical failure in the films such as cracks which are to be initiated in ED where crystals are preferably aligned [96].

Thermal aging reveals an increase of up to 25 % for E for V4 in ED after 2 months, while for the rest of the blends, it remains constant. σ_{max} increases for V2 in ED and TD after 1 month. Furthermore, ε_{break} in ED decreased up to 40 % for V3 and V4 after 1 month and up to 50 % for V1 to V4 after 2 months. This is explained by the fact that thermal aging causes hydrolysis of the macromolecules prompting a pronounced effect of further degradation of the polymer chains in the blends.

This is also in line with the decreasing F_{tear} in the TD, especially after 2 months. The seal strengths decreased for V1, V3, and V4 to close to "zero", indicating a complete loss of the sealing capability after 2 months. Only V2 showed an increase in the seal strength in the range of 10 to 25 %. The importance of seal strength is of huge interest to the packaging industry since the content needs to be restrained inside the bag. Here the stretching and deformation caused by the application load is distributed directly on the seal area and therefore needs to be minimized. Therefore, a decrease of F_{seal} after aging has a negative impact on packaging.

MVR measurements show a clear increase for V1 and V2, which demonstrates chain scission or degradation of polymer chains. On the contrary, the MVR of V3 and V4 decreases up to 80 %, indicating a cross-linking effect.

Table 8 Mechanical properties and MVR of modified compounds V1 to V4 in extrusion direction (ED) and transverse direction (TD) aged for 1 and 2 months at 50 °C. (Paper I)

			Initial		
Prop.	REF	V1	V2	V3	V4
$E_{\rm ED}$ (MPa)	457±36	399±42	501±31	407±66	627±56
$E_{\mathrm{TD}}(\mathrm{MPa})$	181 ± 22	177 ± 22	162 ± 45	147 ± 5	380±66
$\sigma_{ m max,ED}(m MPa)$	27.8±3.1	31±3	24±2	22±3	33±3
$\sigma_{ m max,TD}(m MPa)$	28.8 ± 3.1	33±3	24 ± 2	25 ± 2	35±3
Ebreak, ED (%)	400±48	210±10	250±31	250±31	240±14
$\mathcal{E}_{\text{break, TD}}(\%)$	450±27	450±30	450±30	460±16	440 ± 27

F _{tear, ED} (N mm ⁻¹)	107±3	123±38	110±34	118±38	112±35
$F_{\text{tear, TD}}$ (N mm ⁻¹)	54±17	41±13	35±11	40±13	40±12
$F_{\rm seal}$ (N mm ⁻¹)	7.0 ± 0.3	0.4 ± 0.2	8.5 ± 0.3	8 ± 0.6	10 ± 2
MVR (cm ³ 10 min ⁻¹)	4.0±0.1	5.0±0.1	5.8±0.2	0.6±0.1	1.0±0.1
		Aging 1	l month at	50 °C	
Prop.	REF	V1	V2	V3	V4
$E_{\rm ED}$ (MPa)	437±37	379±21	422±20	453±16	454±38
$E_{\mathrm{TD}}(\mathrm{MPa})$	238 ± 10	183±10	182 ± 18	206±11	265±33
$\sigma_{ m max, ED}$ (MPa)	20.4±2.6	20±1	22±2	31±1	30±1
$\sigma_{ m max,TD} ({ m MPa})$	21.8 ± 2.2	22 ± 2	23±2	33 ± 2	31±1
Ebreak, ED (%)	310±39	270±17	260±17	200±16	220±11
Ebreak, TD (%)	390±20	450±23	480±15	450±32	420±19
F _{tear, ED} (N mm ⁻¹)	104±33	139±43	142±44	132±41	111±35
$F_{\text{tear, TD}}$ (N mm ⁻¹)	57±18	35±11	31±10	30±10	29 ± 9
F_{seal} (N mm ⁻¹)	5.9 ± 0.4	0.4 ± 0.2	6.5 ± 0.3	0.6 ± 0.1	0.9 ± 0.1
MVR (cm ³ 10 min ⁻¹)	4.7 ± 0.1	5.5 ± 0.2	6.3 ± 0.2	0.4 ± 0.1	1.4±0.1
		Aging 2	2 months a	t 50 °C	
Prop.	REF	V1	V2	V3	V4
$E_{\rm ED}$ (MPa)	363±48	400±11	436±26	436±37	460±27
$E_{\mathrm{TD}}\left(\mathrm{MPa}\right)$	197±12	187 ± 23	207 ± 20	207 ± 14	236±25
$\sigma_{ m max, ED} (m MPa)$	21.2±1.5	20±2	32±2	32±3	30±2
$\sigma_{ m max,TD}(m MPa)$	26.5 ± 1.8	21±1	36 ± 2	36±3	29 ± 3
Ebreak, ED (%)	410±26	250±24	200±20	200±32	210±10
$\mathcal{E}_{\mathrm{break, TD}}(\%)$	500 ± 20	430±14	480±39	480 ± 29	430±33
$F_{\text{tear, ED}}$ (N mm ⁻¹)	80±25	138±43	130±30	112±35	109±34
$F_{\text{tear, TD}}$ (N mm ⁻¹)	51±16	31±13	32±11	30±10	43±14
$F_{\rm seal}$ (N mm ⁻¹)	4.8 ± 0.6	0.6 ± 0.3	5.5 ± 0.9	0.5 ± 0.1	
MVR (cm ³ 10 min ⁻¹)	5.5±0.2	5.9±0.2	7.9±0.3	0.35±0.1	1.2±0.1

These results show that the considered CECLs influence on both mechanical properties and flowability, also affected by aging time and temperature. From these results, it is possible to deduce that the introduction of the CECL V1 and V2 leads to chain scission or polymer chain degradation, whereas V3 and V4 indicate a cross-linking effect.

Gel permeation chromatography (GPC) was performed to evaluate if molecular degradation has occurred, Table 9. The results confirm the earlier findings of MVR measurements that the CECL causes cross-linking in V3 and V4 as their molecular masses ($M_n \approx 75,000 \text{ g/mol}$) are roughly doubled compared to REF. REF, V1 and V2 with molecular masses around 44,000 g/mol showed that the CECL did not significantly change the molecular masses and as a consequence the macromolecular structure. Most CECL molecules have to remain as low molecular components in the polymer chain scission. The significantly higher MVR values of V1 and V2 compared to REF were explained by chain scission. However, according to the GPC results, Table 9, this cannot be the reason for the higher MVR values. They have to be explained by an enhancement of the melt flowability provided the remaining unreacted CECL as low molecular components.

Table 9 Determination of the mean molecular masses due to GPC measurements with number-average molecular mass M_n , weight-average molecular mass M_w , and polydispersity index of the polymer PDI; accuracy of measurements is 2,000 g/mol for M_n and 3,000 g/mol for M_w . (Paper III)

Compound	$M_{ m n}$	$M_{ m w}$	PDI						
	g/mol	g/mol	-						
	Initial state								
PBAT	43,000	107,000	2.45						
REF	44,000	83,000	1.89						
V1	42,000	81,000	1.92						
V2	46,000	92,000	1.98						
V3	75,000	149,000	1.99						
V4	72,000	141, 000	1.95						
PLA	218,000	309,000	1.42						

The film properties are influenced by e.g. the base polymer's intrinsic properties such as molecular weight, molecular weight distribution, density, long and short branches, equipment and processing conditions which result in different film properties. This means that orientation, crystallinity and morphology have a pronounced effect on the final film's mechanical and thermal properties. Due to the film-blowing process with different BUR and DR in ED and TD, anisotropic mechanical behavior and corresponding differences in a morphological structure can be expected.

Figure 27 a-d shows Young's modulus E for the various BUR (1.5, 2.5 and 3.5), the tensile strength σ_{max} , the elongation at break ε_{break} and the tear resistance F_{tear} in the ED and TD.

The different BUR and the film morphology strongly influenced the mechanical properties. For a BUR of 1:2.5, the introduction of CECL modified the properties of the REF.

The *E*, Figure 27a, with increasing BUR, *E* decreases in ED. As the film is stretched in ED the structure deforms into an iso-stress pattern, where *E* is dominated by the amorphous phase. In TD however, *E* increases for V1 to V4 and slightly decreases for REF, showing that when stretching takes place in TD, the structure deforms in an iso-strain pattern and the *E* is dominated by the crystalline phase.

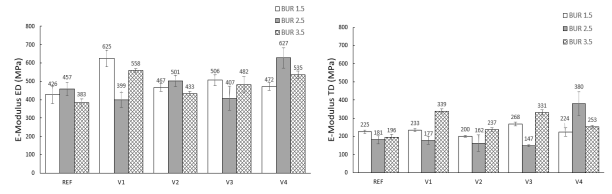


Figure 27a. Young's modulus E for REF and V1 to V2 with the different BUR: 1.5, 2.5 and 3.5, in ED and TD directions. (Paper V)

Continuing the comparison BUR 1.5 to 3.5, the results show that with increasing BUR the σ_{max} in ED slightly decreases, for REF, V2 and V3, but for V1 a more distinct decrease was measured, Figure 27b. In TD however, σ_{max} increases with increasing BUR (1.5 vs 3.5) for all blends. Increasing the BUR causes an increase of molecular orientation in TD and a decrease in orientation in ED due to a higher feed rate to maintain constant thickness.

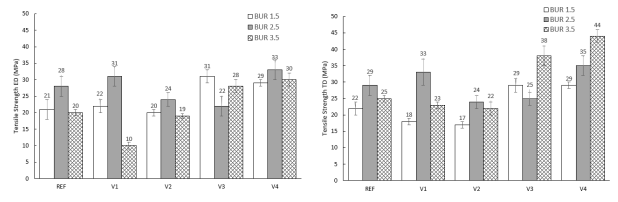


Figure 27b. Tensile strength σ_{max} for REF and V1 to V2 with the different BUR: 1.5, 2.5 and 3.5, in ED and TD directions. (Paper V)

Concerning $\varepsilon_{\text{break}}$, Figure 27c, in ED the values increase from lower BUR to higher BUR, and in TD it decreases from BUR 1.5 to 3.5. The higher surface area to cool the bubble at higher BUR could be a major factor in the lower crystallinity, resulting

in a higher TD elongation. Thus, the non-uniform stretching behavior reflects the anisotropic orientation of the blends.

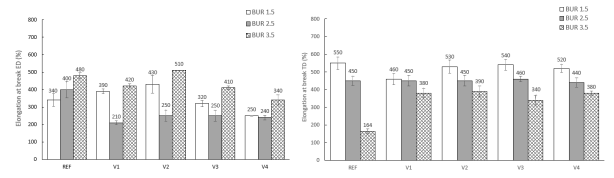


Figure 27c. Elongation at break ε_{break} for REF and V1 to V2 with the different BUR: 1.5, 2.5 and 3.5, in ED and TD directions. (Paper V)

Regarding the tear resistance, Figure 27d, in ED a decrease from BUR 1.5 to 3.5 for REF, V1 and V4 while for V2 and V3 an increase was measured. First, the decrease can be explained by the nucleated structure where a high degree of chain orientation along ED leads to poor tear strength in that direction. This is due to the fact that the tear propagation parallel to the backbone chain requires less energy than propagation perpendicular to the backbone chain. For the increase in V2 and V3, as evidenced already, the cross-linking effect could lead to higher energies being promoted for the propagation into the backbone chain. In TD, the decrease reaches more than 50 % from BUR 1.5 to 3.5, where films are unable to resist tear propagation along TD for BUR 3.5.

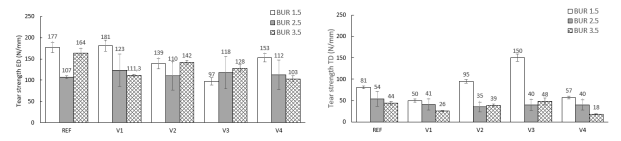


Figure 27d. Tear resistance F_{tear} for REF and V1 to V2 with the different BUR: 1.5, 2.5 and 3.5, in ED and TD directions. (Paper V)

Seal strength behavior, Figure 28, shows relatively constant values for the various BURs. Except for V2 and V4, BUR 2.5 a clear increase of 25 % was measured compared to REF, indicating that independently of the BUR, V2 and V4 characteristics of cross-linking are shown as reported above.

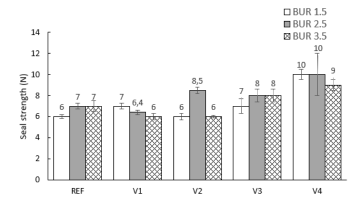


Figure 28. Seal strength for REF and V1 to V2 with the different BUR: 1.5, 2.5 and 3.5, in ED direction. (Paper V)

To understand these mechanical findings, a model was developed that describes the BUR-dependent lamellae formation during crystallization. In blown film extrusion an originally almost isotropic melt volume V_0 is drawn in ED and blown up in TD. To achieve the desired film thicknesses, DR is adjusted due to the chosen BUR, Table 5. This generates a macroscopic geometry change of V_0 , Figure 29, and introduces corresponding anisotropic orientation states to the macromolecules being smaller than the macroscopic deformation because of relaxation processes. Furthermore, the molecular orientations are governed by melt viscosities of PBAT and PLA and local cooling rates.

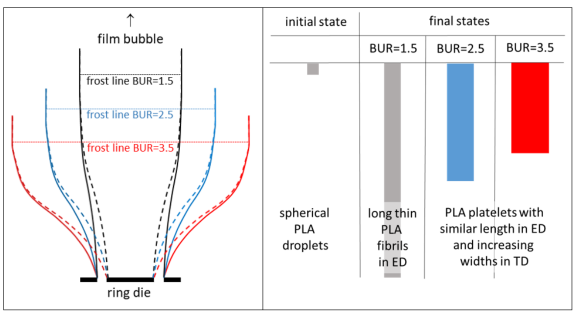


Figure 29: Visualization of the BUR dependent film bubble geometries with corresponding frost lines (left) and macroscopic deformation states of blown films and expected geometries of the PLA phase if deformed during film blown extrusion (right).

Le Marec et al [97] report zero shear viscosities of 8,000 MPa s for M_W^{PLA} =111 kg/mol and 600 MPa s for M_W^{PLA} =54 kg/mol at 166 °C. Deng et al [98] modeled the melt viscosities and found 14,000 Pa s for PLA with M_W^{PLA} =94 kg/mol and 1,200 Pa s for PBAT at 175 °C. This shows that one deals with a viscosity ratio of PLA to PBAT of 4 to 10. The consequence is that PLA droplets are significantly less deformed than the PBAT matrix for a given stress state. Due to the PLA melting temperature of 160 °C and a roughly 4 times larger mean molecular mass compared to PBAT (Paper III), PLA crystallizes prior to PBAT and much closer to the die at still small melt speeds and stresses.

After compounding the spherical PLA droplets have diameters of 200 to 1,000 nm confirming results of other researchers that provide diameters of 200 to 1,500 nm depending on the processing conditions. Assuming that the molecular orientations relax similarly in ED and TD with a polymer characteristic behavior, most of the orientation is introduced to the PBAT matrix and only a little to the dispersed PLA phase. At BUR=1.5 representing an almost uniaxial deformation, one can expect that the PBAT macromolecules are extremely oriented in ED and only little in TD, and that the dispersed spherical PLA droplets are partly deformed to fibrils by elongational flows to elongations being smaller than the DR. With increasing BUR, the deformation gets a more biaxial character, and one may expect that the PLA spheres are deformed to platelets of similar lengths and increasing widths according to DR and BUR.

The molecular masses of the base polymers were determined to M_N^{PBAT} =43,000 g/mol and M_N^{PLA} =218,000 g/mol, and M_W^{PBAT} =107,000 g/mol and M_W^{PLA} =284,000 g/mol, respectively (Paper III) Taking this into account, the melt viscosity of PLA has to be significantly higher (around a factor 10 due to Fox [99]) compared to PBAT. Furthermore, PLA crystallizes between 110 and 120 °C to crystallinities of around 30 % whereas PBAT crystallizes between 60 and 80 °C to crystallinities of around 15 %. This has the consequence that there is hardly any deformability of the PLA droplets in the still soft PBAT matrix during blown-film extrusion shown by the large number of small and relatively undeformed PLA droplets in the fracture surfaces. This needs to estimate the time and distance from the die at which PLA starts to crystallize.

The melt leaves the extrusion die with a thickness of 0.8 mm and a temperature of 165 °C. At the frost line, the blown film has a thickness of 25 μ m and a temperature of 50 to 70 °C being below T_{cr} . The cooling process is characterized by the cooling time $\tau_{cooling}$, the time in which the temperature difference between melt and film decreased by 63 %. It depends strongly on the film thickness d

$$\tau_{cooling} = \frac{1}{4\pi^2} * \frac{c_p \rho}{\lambda} * d^2 \tag{1}$$

with specific heat capacity c_p , density ρ and heat conductivity λ . The time to ambient temperature $t_{ambient}$ – here approximately 40 °C – is given by

$$t_{ambient} = 3 * \tau_{cooling} \tag{2}$$

Introducing polymer typical material constants provides cooling times of 5 to 6 s for a thickness of 0.8 mm and 5 to 6 µs for a thickness of 25 µm, respectively. As the roll-up speed is around 20 mm/s and the melt output is around 300 mm³/s, the polymer is drawn within 10 to 12 s from the die to the frost line. Then the melt speed close to the die is around 1.5 mm/s. Thus, the melt temperature decreased under the crystallization temperature of PLA after 3 s. Within these 3 s, the extruded tube has been drawn away 6 to 8 mm from the die indicating small elongations. Thus, the elongational stresses in the melt remain also small, and one can conclude that the PLA droplets have crystallized before massive drawing occurs. This explains why the melting temperature hardly varies with BUR. Another consequence is that the deformation during blown film extrusion takes place only in the PBAT matrix after 5 s.

At small BUR corresponding to large DR, the macromolecules of the melt are highly oriented in ED easing crystallization in relatively thick lamellae that grow in TD. These lamellae may fix the highly oriented amorphous regions, Figure 30 left. Thus, the macromolecules of the rubbery amorphous PBAT phase keep their orientation states and increase stiffnesses in ED and decrease elongations at break in ED compared to TD. As the crystallinity of PBAT is around 15 %, the lamellae represent "solid platelets" in between oriented macro-molecules of the amorphous phase are sheared. This explains the roughly 50 % reduction of stiffnesses in TD compared to ED and the larger elongations at break.

With increasing BUR, the orientation of the macromolecules is increased in TD and decreased in ED causing inclined preferential orientations with respect to ED. Because of the biaxial orientation of the macromolecules, thinner crystalline lamellae grow with an inclination preferentially in TD and between them, thin and short cross-lamellae grow preferentially in ED, Figure 30 right. The orientation in ED still exceeds that in TD significantly even at BUR=3.5. Thus, one expects increasing elongations at break in ED – true if one compares BUR=1.5 and BUR=3.5 but not for V1 to V4 at BUR=2.5 – and decreasing elongations at break in TD – true for REF, V1 to V4 at all BUR, Figure 27c.

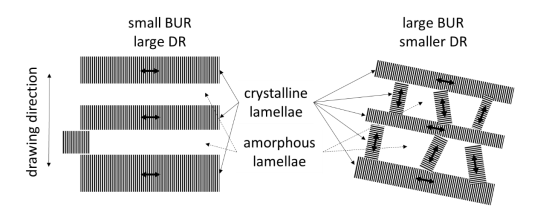


Figure 30: Scheme of BUR dependent lamellae formation in the PBAT phase.

With the BUR evaluation, the tensile strengths do not depend systematically on BUR. This concludes that the different processing induced BUR dependent lamellae structures have only minor effects on tensile strengths and that the tensile strengths are mainly determined by the entanglement density in the PBAT phase. This explains the higher tensile strengths V3 and V4 both in ED and TD with double molecular masses compared to REF, V1 and V2.

4.1.2 Viscoelastic properties

Both CECL compounding and processing affect temperature-dependent storage moduli in ED and TD in terms of $T_{\rm g}$ shifts and changes in stiffness. Viscoelastic properties were measured on 100 μ m films because the resolution of the DMA was insufficient for the 25 μ m films.

Figure 31 shows the comparison of temperature-dependent logarithmic storage moduli of REF and V1 to V4 in ED and TD at 1 Hz. For both ED and TD, $T_{\rm g}$ of the soft segment ($T_{\rm g,ss}$) can be detected around -40 °C and for the hard segment ($T_{\rm g,hs}$) around 50 °C. E' between $T_{\rm g,ss}$ and $T_{\rm g,hs}$ increase in ED for V1 and V4 compared to REF. Although, V2 and V3 show lower E' values than REF at lower temperatures, but don't differ between $T_{\rm g,ss}$ and $T_{\rm g,hs}$. Above $T_{\rm g,hs}$ only V1 shows similar values as REF, while V2 to V4 present lower stiffness. In TD, V1 exhibits an increase in storage modulus, whereas only slight increases are observed for V2, V3 and V4 compared to the REF. Analogous to ED, above $T_{\rm g,hs}$ only V1 depicts higher stiffness and V2 to V4 clearly shows lower stiffness. Overall, the moduli increase is more pronounced in ED compared to TD, which presents anisotropic behaviour. This shows that CECL reactions are leading to cross-linking for V1 and V4, whereas the shift of the $T_{\rm g,hs}$ to lower temperatures indicates chain scission or degradation. Meanwhile, V2 and V3 hardly show any differences in stiffness but above $T_{\rm g,hs}$ both depict lower values which indicates chain scission or degradation.

Figure 32 shows the comparison of temperature dependent loss factor ($tan \delta$) of REF and V1 to V4 in ED and TD at 1 Hz.

 $Tan \ \delta$ at $T_{g,ss}$ shows a decrease of values in ED for V1 to V4 compared to REF. In contrast, at $T_{g,hs} \ tan \ \delta$ shows a shift to lower temperatures. Also, the exact same effect is also seen in TD. This indicates that CECL reactions preferably take place in the soft segment phase.

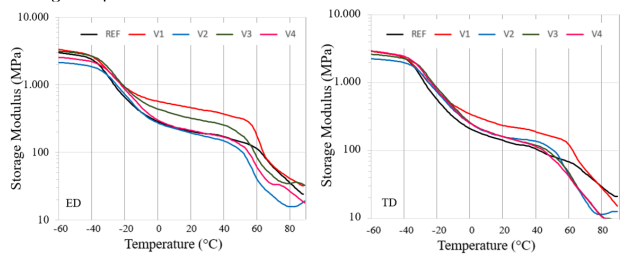


Figure 31. Comparison of temperature dependent logarithmic storage moduli [log(E'(T, ω =1Hz)] of REF, V1 to V4 in ED (left) and TD (right). (Paper II)

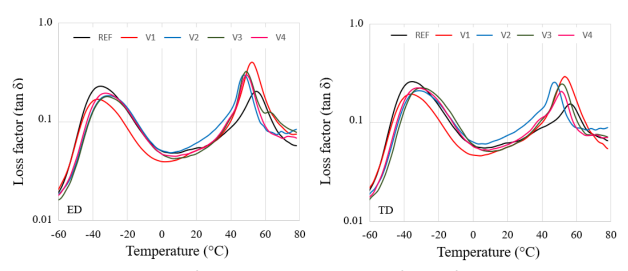


Figure 32. Comparison of temperature dependent loss factor of REF, V1 to V4 in ED (left) and TD (right) at 1 Hz. (Paper II)

Evaluating the DMA measurements in detail, Table 10, for REF and V1 the $T_{\rm g,ss}$ (E''_{max}) coincide, whereas they are shifted for V2 to V4 by 3 to 5 K towards higher temperatures indicating cross-linking in soft segment phases. Interestingly, $T_{\rm g,hs}$ of V1 to V4 are decreased by 2 to 4 K, respectively, compared to REF. This is in contrast to the DSC data, where $T_{\rm g,hs}$ of V1 to V4 increased by 7 to 8 K, Table 10.

This can be explained by the fact that films shrink freely in the DSC experiment reducing the free volume and shifting $T_{g,hs}$ to higher temperatures. Shrinkage is prevented in the DMA experiment due to the sample clamps and the free volume is maintained at least at the beginning of softening.

Table 10 Storage moduli E' and loss moduli E'', glass temperatures of soft segments $T_{g,ss}$ and glass temperatures of hard segments $T_{g,hs}$ determined using maxima of E'' and $tan \delta$ of REF and V1 to V4 compounds evaluated at 1 Hz. (Paper I)

Sam	ple	E'	<i>E</i> ''	$T_{\mathrm{g,ss}}(E^{\prime\prime})$	$T_{\mathrm{g,hs}}(E^{\prime\prime})$	$T_{\rm g,ss}$ (tan δ)	$T_{\rm g,hs}$ (tan δ)
		MPa	MPa	°C	°C	°C	°C
REF	ED	177±33	10.3±1.2	-41.7±0.2	52.4 ± 0.8	-35.5±0.1	54.8 ± 0.7
KLT	TD	124±13	8.0±1	-41.9±0.1	55.0±0.4	-35.2±0.1	56.7±0.3
V1	ED	387±38	21.0±1.4	-41.7±0.6	50.6±1.2	-36.8±0.9	53.3±1.2
V 1	TD	322±17	20.0±1.6	-41.9±0.1	51.0±0.1	-36.1±0.3	53.6±0.1
V2	ED	210±33	13.0±1.4	-38.8±0.7	50.0±4.2	-32.7±0.5	51.9±4.5
V 2	TD	151±12	11.0±0.8	-38.9±0.9	50.0±1.3	-32.2±0.9	52.1±1.1
V3	ED	184±15	9.3±1.2	-36.8±0.7	50.5±5.8	-31.0±0.5	52.5±6.2
V 3	TD	126±5	7.3±0.5	-37.2±0.5	53.6±3.8	-30.6±0.5	56.1±4.0
V4	ED	270±24	15.0±1.9	-38,7±1.3	50.0±4.3	-32.9±1.9	52.0±4.4
V4	TD	133±6	8.7±0.9	-38.0±0.4	49.6±5.7	-31.8±0.8	51.8±6.1

According to DMA data, V1 hardly affected the $T_{\rm g}$ values of both segments, whereas other CECLs caused the $T_{\rm g}$ to increase in the soft segment phase and a decrease in the hard segment phase. This means that cross-linking mainly occurred in the soft segment phase. In conclusion, cross-linking in the hard segment phase was found for V3 and V4 after processing.

4.1.3 Elongational properties

The replacement of packaging polymers as low-density polyethylene, having well-defined strain hardening, with biodegradable materials brings the necessity of reliable evaluation of their resistance to elongational deformation. An evaluation using a Sentmanat Extensional Rheometer (SER) was performed (Paper IV).

Prior to measurements of elongational viscosities using the SER device, two principal conditions should be checked: mild intensity of sample sagging during the process of temperature setting and an acceptably small deviation from exponential decrease of sample cross-sections. These evaluations are presented in Appendix B.

Due to sagging, the precise of this quantity shortcomings cannot be determined. However, if we want to compare two materials – regardless of their geometrical form

– using the SER device the term 'elongation ratio at film rupture' (ER_{FR}) seems to be adequate and relates the time spans t_{rupt} that elapsed between a start of drum revolution with a fixed sample and a final film rupture for two samples 1 and 2

$$ER_{FR} = \frac{t_{\text{rupt,1}}}{t_{\text{rupt,2}}} \tag{3}$$

First, it was necessary to determine the temperatures under which the SER device could be used for the investigated five PLA/PBAT blends. Based on the range between 135 °C and 170 °C it was found that the optimal common temperature interval is 150 ± 5 °C for the CECL modified blends V1 to V4, and for the material V4 even at higher temperatures. In this interval, all PLA/PBAT blends exhibited optimal balanced fluid-like behaviors with acceptable sagging. Nevertheless, the remaining deviations of individual optimal temperatures may worsen comparisons among sample characterizations using ER_{FR} a little bit. Only in the case of REF, the temperature had to be lowered to 140 °C due to intensive sagging at 150 °C.

The thermo-mechanical properties of blown films of REF, V1 to V4 are anisotropic (Paper I) because of orientation effects of dispersed droplets in the melt [100] and polymer chains both introduced during blown film extrusion [101].

The mean times to film rupture scattered less than 5 % for each extensional rate, and the TD values always exceeded those in ED, as seen on Figure 33, showing that the morphology formation differs with respect to ED and TD during the blown film process.

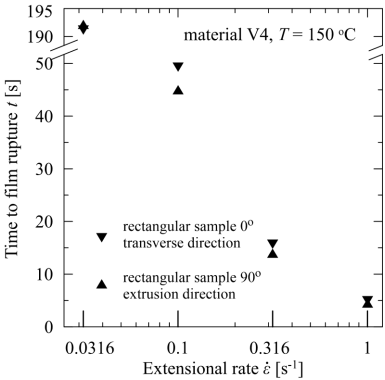


Figure 33: Times to film rupture of V4 - PLA/PBAT modified with V4 in extrusion and transverse directions. (Paper IV)

Not surprisingly the time values in both directions for the lowest extensional rate $(0.0316~{\rm s}^{\text{-1}})$ are practically identical (differs by 0.5~%) as for such low extensional rates and temperature of $150~^{\circ}\text{C}$ the process of disentanglement effectively contributes to morphological structural change. At these rates the macromolecules have enough time to slip one out of the other and the initial differences generated by the anisotropic film blowing conditions vanish completely. The values for a slant angle 45° were also measured and are located between the two limiting (0° and 90°) values, for the sake of lucidity not depicted in Figure 33.

The differences for other rates expressed by the anisotropy ratio R_{TD-ED}

$$R_{\rm TD-ED} = \frac{t_{\rm rupt,TD} - t_{\rm rupt,ED}}{t_{\rm rupt,TD}} \times 100 \%$$
 (4)

attain approximately 10 %, as seen on Figure 34.

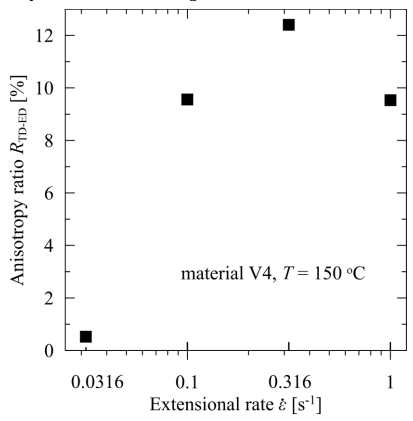


Figure 34: Anisotropy ratio of times to film rupture for V4 - PLA/PBAT modified with V4 in extrusion and transverse directions. (Paper IV)

For REF and V1 to V4, there was measured a time interval that elapsed between the start of SER drums revolution and material rupture. All measurements were carried out for two widths of rectangular samples (transverse direction, 0°): 12.7 mm (maximum possible width in the SER device) and 9 mm. Each measurement was conducted at least in triplicate and the results as shown on Figure 35.

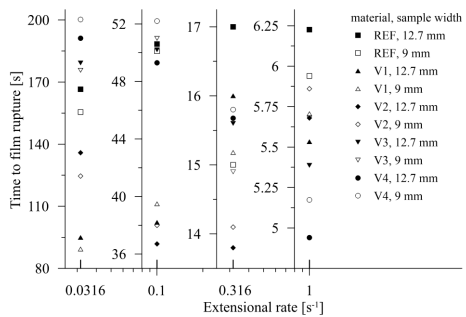


Figure 35: Elongational rate dependent times to film rupture in TD, individual charts. (Paper IV)

It is apparent that the initial macromolecular interplay for low extension rates favors disentanglement. It is replaced by consecutive passage to strain hardening where the rupture time intervals for pure PLA/PBAT blend significantly dominate. It is necessary to remind that REF was measured at 140°°C, i.e. by 10 °C lower than the modified samples V1-V4. This difference is suppressed for lower extension rates (as for 0.0316 s⁻¹) due to the process of disentanglement as discussed above. Dominance of this process is documented by intermixing REF values with the modified samples regardless of the temperature under which the experiments were carried out.

With increasing extensional rates, strain hardening starts to compete and successively replaces the process of disentanglement. This can be demonstrated for REF at an extensional rate of 1 s⁻¹ at 140 °C having the highest values of time to film rupture. However, at 150 °C REF exhibits a pronounced liquid-like behavior preventing elongation measurement. On the contrary, decreasing temperature by 10 °C reduces the liquid-like behavior of modified samples, which would reflect in apparent prolongation of breakage time (for instance by 17 % for V1, sample width 12.7 mm, extensional rate 1.0 s⁻¹).

The practical coincidence of the results expressed through a ratio relating the times to film fracture for both widths

$$R_{\text{width}} = \frac{t_{\text{rupt,12.7 mm}}}{t_{\text{rupt,9 mm}}} \tag{5}$$

documents an excellent homogeneity of the prepared materials without air bubbles and with initial uniform thickness due to good mixing during melt compounding. The dispersion of the ratio R_{width} is limited to the interval between 0.95 and 1.1, as seen on Figure 36, which corresponds to the experimental errors.

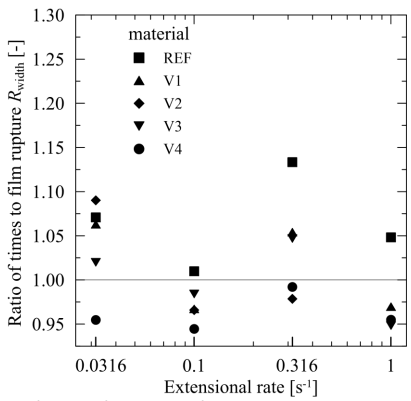


Figure 36: Ratios of times to film rupture of width 12.7 mm to width 9 mm. (Paper IV)

The result obtained for the modified blends corresponds to the molecular mass exhibited by the results from GPC (Paper III). The results showed significant increases in the molecular masses of V3 and V4 due to cross-linking reactions which correspond to findings based on the MVR measurements (Paper I). For V3 and V4 the number average molar mass nearly doubled that of REF (meaning that a single CECL molecule joins two polymer chains on average), whereas V1 and V2 have more or less identical molecular masses as REF. This indicates that in the case of V3 and V4 the CECL molecules react chemically with the polymer chains mainly forming a link between two macromolecules of PBAT on average. In the case of V1 and V2 the mean molecular masses are identical to REF. This may mean that CECL molecules hardly react with the PBAT molecules but cause chain scission of PLA molecules. Furthermore, a certain amount of the CECL may remain unreacted in V1 and V2. This explains the higher MVR values of V1 and V2 and does not exclude grafting reactions of CECL to macromolecules.

The obtained results justify the introduction of the quantity 'elongation ratio at film rupture ER_{FR} ', as presented in (3), comparing elongational attributes of the

modified blends. For the interpretation of the measurements using the SER device, one has to keep in mind that V1 and V2 may contain unreacted rests of their CECL easing chain slipping of macromolecules due to their softening effects and that V3 and V4 consist of longer macromolecules, thus, having higher entanglement or cross-linking densities with a more pronounced strain hardening behavior.

4.2 Thermal properties

CECL process-induced structural changes were investigated with DSC.

Figure 37 shows 1st heating run of the granules after compounding of REF with the CECL's but before film blowing. Around 50 °C a glass transition of the hard segment $T_{\rm g,hs}$ occurs followed by an evaporation of tetrahydrofuran (THF) at 62 °C indicating chemical reactions of PBAT and PLA during compounding [96]. Furthermore, broad melting peaks for PBAT around 115 °C and for PLA around 150 °C are measured.

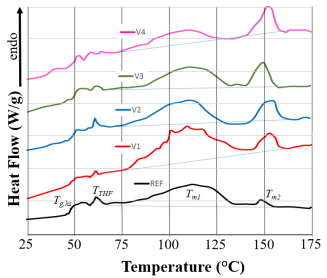


Figure 37. DSC traces of REF, V1 to V4, I^{st} runs (0.2 W/g) of granules with glass temperature $T_{g,hs}$, THF boiling temperature T_{THF} , melting temperatures T_{m1} and T_{m2} , and corresponding melting and crystallization peaks. (Paper I).

Evaluating the DSC traces in detail of the 1st, cooling and 2nd heating runs of REF and V1 to V4 in granules, Table 11, it can be seen that $T_{g,hs}$ of V1 to V4 decrease in relation to REF in the 1st run, meaning that the hard segments have higher mobility due to CECL induced reactions which increase distances between chains or leave unreacted CECL molecules acting as softeners [102]. During the 2nd heating run, the $T_{g,hs}$ increased compared to the 1st heating run indicating further chemical reactions in the compound representing the extrusion process before film blowing. Concerning REF the $T_{g,hs}$ (onset) value of V1 is reduced from 55.9 °C to 51.1 °C indicating some chain scission or degradation in the hard segment phase. $T_{g,hs}$ 2

(onset) for V2 is unaltered indicating no degradation, whereas $T_{\rm g,hs}$ (onset) for V3 and V4 has increased from 55.9 °C to 58.4 °C and from 55.9 °C to 59.2 °C, respectively, indicating cross-linking.

The melting temperatures $T_{\rm ml}$ of V1 to V4 show lower values than $T_{\rm ml}$ (REF) of 113 °C indicating that CECL hinders the formation of thicker lamellae in PBAT. In the 2nd heating run, all compounds have the first melting temperatures $T_{\rm ml}$ around 116 °C. The second melting temperature $T_{\rm m2}$ of REF and V1 to V4 range between 150 °C and 153 °C indicating similar structures of the PLA phase in all compounds. Nevertheless, on the second melting temperature $T_{\rm m2}$ peaks are only found for REF and V2 signalling a suppressed PLA crystallization in V1, V3 and V4. The heats of fusion of V3 and V4 are in both heating runs smaller than in the REF showing that cross-linking reduces crystallinity. The crystallization enthalpy in the cooling runs exceeds the total heats of fusion (sum of both peaks) by 20 to 50 % which is surprising as they also exceed crystallization enthalpies. This can be explained by the superposition of the crystallization and glass transition of the hard segment.

Table 11 Glass temperature of hard segments $T_{\rm g,hs}$, crystallization temperature $T_{\rm cr}$, melting temperatures $T_{\rm m1}$ of PBAT and $T_{\rm m2}$ of PLA with corresponding heats of fusion $\Delta H_{\rm m1}$ and $\Delta H_{\rm m2}$ determined by 1st run, cooling run and 2nd run of DSC measurements of granules of REF and V1 to V4 compounds. (Paper I)

			1	` 1 /	
Prop.	REF	V1	V2	V3	V4
$T_{g,hs}1 (^{\circ}C)$	50±0.03	43.7±2.02	46.7±0.08	47.8±0.27	46.2±0.07
$T_{\rm m1}$ _1 (°C)	81.2±0.5	84.8 ± 0.5	77.8±1.2	94.9±1.5	81.1±0.1
$T_{\text{m2}}I$ (°C)	143.2±0.8	144.2±0.2	143.4±0.7	142.7±1.5	143.2±0.6
$T_{\rm cr}(^{\circ}{ m C})$	75.4±0.3	83.4±0.02	83.0±0.43	87.3±0.23	87.2±0.21
$T_{\rm g,hs}$ _2 (°C)	55.9±0.28	51.1±2.1	54.9±0.86	58.4±0.46	59.2±1.5
T_{m1}_{2} (°C)	88.4±0.35	89.1±0	88.9±0.35	83.6±0.24	83.8±0.12
T_{m2} 2 (°C)	143.8±0.3	-	144.0±0.12	-	-
$\Delta H_{\rm ml} l (J g^{-1})$	8.5±0.17	9.4±0.6	6.9±0.6	5.1±0.15	5.2±0.75
$\Delta H_{\text{m2}} 1 \text{ (J g}^{-1})$	0.3 ± 0.08	1.1±0.18	1.9±0.04	1.1±0.5	2.8±0.12
$\Delta H_{\rm cr}({ m J g^{-1}})$	11.0±0.3	11.6±0.13	10.9±0.2	9.1±0.12	10.3±0.09
$\Delta H_{\rm m1}$ _2(J g ⁻¹)	7.3±0.2	7.3±0.16	5.8±0.14	5.8±0.28	5.8±0.05
ΔH_{m2} 2 (J g ⁻¹)	0	-	0.8±0.05	-	-

Table 12 shows a detailed evaluation of the DSC traces for 25 μ m films. Results for films of V1 to V4 exhibit similar behaviour as the granules: in the 1st heating run V3 and V4 have lower $T_{g,hs}$ than V1 and V2, whereas in the 2nd heating run it is vice versa. This indicates that the amorphous hard segment phases of V3 and V4 are more disordered after film blowing probably due to incomplete chemical reactions.

The crystallization enthalpies during the cooling run of REF and V1 to V4 are higher than those of the granules, showing that the remaining preferential chain orientations due to the film blowing even in the melt after the first heating run easing and enabling subsequent crystallization [88, 103].

Table 12 Glass temperature of hard segments $T_{\rm g,hs}$, crystallization temperature $T_{\rm cr}$, melting temperatures $T_{\rm m1}$ of PBAT and $T_{\rm m2}$ of PLA with corresponding heats of fusion $\Delta H_{\rm m1}$ and $\Delta H_{\rm m2}$ determined by 1st run, cooling run and 2nd run of DSC measurements of 25 µm films of REF and V1 to V4 compounds. (Paper I)

Property	REF	V1	V2	V3	V4
$T_{g,hs}_1(^{\circ}C)$	43.0±0.7	44.9±0.6	44.4±1.7	41.7±0.4	43.3±0.9
$T_{\rm m1}$ _1 (°C)	121.2±0.1	121.4±0.1	117.1±0	106±2.0	118±0.2
$T_{\rm ml}$ _2 (°C)	151.0±0.1	151.1±0.1	152 ± 0.2	148±0.2	149.2±0
$T_{\mathrm{THF}}(^{\circ}\mathrm{C})$	60.5±0.4	61.3±0.8	63.9±0.1	63.8±0.7	63.1±0.1
$T_{\rm cr}(^{\circ}{\rm C})$	78.2 ± 0.2	77.9 ± 0.1	72.3±0	74.3 ± 0.5	75.4±0.3
$T_{\rm g,hs}$ _2 (°C)	50.0±1.1	53.5±1.6	51.8±1.5	55.1±1.8	53.2±3.5
$T_{\rm ml}$ _2 (°C)	151.0±0.1	151.1±0.1	153.0 ± 0.2	148.0±0.2	149.2±0
T_{m2} 2 (°C)	-	-	152.9±0	-	149.3±0
$\Delta H_{\mathrm{THF}}(\mathrm{J}\;\mathrm{g}^{-1})$	0.2 ± 0.1	0.3 ± 0.1	0.5 ± 0.0	0.4 ± 0.0	0.3±0.1
$\Delta H_{\rm ml} l (J g^{-1})$	5.4±0.4	5.4±0.5	5.1±0.2	6.4±0.1	3.7±0.0
$\Delta H_{\text{m2}} 1 \text{ (J g}^{-1})$	1.1±0.1	1.2±0.1	2.9 ± 0.1	1.2±0.0	3.5±0.1
$\Delta H_{\rm cr}({\rm J}~{\rm g}^{-1})$	15.2±0.9	15.1±0.1	14.7 ± 0.1	11.7±0.3	12.7±0.6
$\Delta H_{\rm ml}$ _2 (J g ⁻¹)	5.4±0.4	9.9±0.4	5.1±0.2	6.4 ± 0.1	3.7±0
ΔH_{m2} 2 (J g ⁻¹)	1.1±0.1	1.2±0.1	2.7±0.1	1.2±0	3.5±0.1

The effects on thermal properties of REF, V1 to V4 with the variation of BUR were also evaluated, as seen in Table 13. The glass transition temperatures of the hard segments $T_{g,hs}$ decrease with BUR indicating higher molecular mobilities of the amorphous phases due to slightly larger mean chain distances caused by grafted CECL molecules. Only at BUR=3.5, V3 and V4 exhibit higher $T_{g,hs}$. Possibly polymer chains can better align with each other under a more pronounced biaxial orientation because of higher melt viscosities.

The PBAT melting peaks have widths of more than 30 °C indicating wide distributions of lamellae thicknesses. In that respect, the temperatures of the melting peaks T_{ml} provide insights into how the choice of BUR affects the lamellae growth during the crystallization of blown films and the achievable crystallinity. For all BUR, T_{ml} of V3 and V4 are lower than for REF, V1 and V2 indicating that higher molecular weights and melt viscosities shift the peaks to lower lamellae thicknesses. Furthermore, a strong dependency of T_{ml} is observed for BUR=2.5 with 5 to 15 °C decreased T_{ml} , and for BUR=3.5 with 1 to 5 °C increased T_{ml} compared to BUR=1.5. Interestingly no systematic change is observed for heats of fusions ΔH_{ml} indicating that the interplay of nucleation, crystal growth and film cooling is complex. This is stated by the crystallization temperatures T_{cr} and crystallization enthalpies ΔH_{cr} which both show no systematic BUR dependency.

Table 13 BUR dependency of thermal properties of REF, V1 to V4 in terms of glass temperature of hard segments $T_{g,hs}$, melting temperatures T_{ml} of PBAT and T_{m2} of PLA with corresponding heats of fusion ΔH_{ml} and ΔH_{m2} determined from 1st heating runs, crystallization temperature T_{cr} , and crystallization enthalpy ΔH_{cr} determined from cooling runs; temperature accuracy ΔT is \pm 1 °C and enthalpy accuracy ΔH is 1 J/g. (Paper V)

	$T_{ m g,hs}$	T_{m1}	$\Delta H_{\mathrm{m}1}$	$T_{ m m2}$	ΔH_{m2}	$T_{ m cr}$	$\Delta H_{ m cr}$
	°C	°C	J/g	°C		°C	J/g
	midpoint	PBAT	PBAT	PLA	PLA	peak	
			BUR 1	5			
REF	53	117	4.6	147	0.5	73	10.0
V1	57	120	3.9	151	1.7	83	11.9
V2	50	119	6.5	150	1.4	83	13.3
V3	49	113	5.9	149	1.0	71	12.3
V4	51	116	7.5	150	0.2	80	13.0
			BUR 3	3.5			
REF	48	122	4.2	149	0.5	87	13.6
V1	49	120	3.4	151	1.9	93	11.4
V2	48	125	8.5	149	1.9	93	13.7
V3	51	112	4.1	151	1.1	71	12.2
V4	50	119	5.6	151	0.7	79	12.9

These DSC results for the granules and films showed that the chemical reactions were incomplete after compounding and the film blowing intensified them, even for 25 µm films were being rapidly cooled. This behavior is explained by the CECL

molecules being linked with one reactive site of polymer chains during compounding, while other reactive sites only show reactive possibilities if a chain slip occurs, such as in elongational flow during film blowing. For the evaluation with BUR, it is possible to conclude that the PLA phases crystallize during blown film extrusion under similar conditions for all BUR.

To confirm the interactions interpreted from DSC and mechanical analysis, FTIR-ATR was performed on the 25 μ m blown films, Figure 38. FTIR-ATR from the granules are not shown here, because no differences between granules and blown films were detected. According to Standau et al. [103] and Yuniarto et al. [104], the PLA band around 752 cm⁻¹ together with the vibration of α -methyl band around 864 cm⁻¹ are associated with ester (O-CH-CH₃). Many weaker peaks in the range of 1250-1050 cm⁻¹ are assigned to C-O from carboxyl groups and C-O-C stretching vibrations, and the peak at 1748 cm⁻¹ is associated with the carbonyl C=O stretching vibration.

The functional groups of PBAT are described as peak at around 1710 cm⁻¹ represents carbonyl groups (C=O) in the ester linkage, at 1265 cm⁻¹ a peak intercepts C-O in the ester linkage, and at around 725 cm⁻¹ a peak stand for methylene (-CH₂-) groups, Figure 38. Bending peaks of the benzene substitutes are located at wavenumbers between 700 and 900 cm⁻¹.

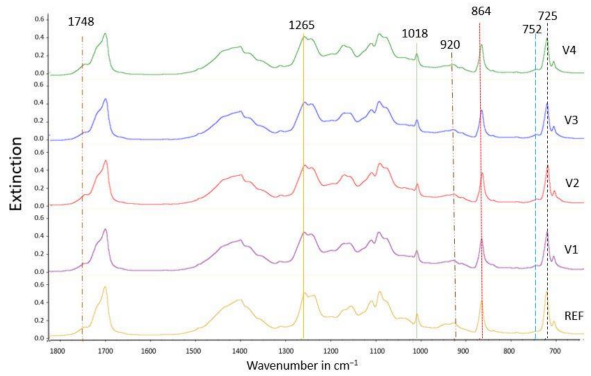


Figure 38: FTIR spectra of the 25 µm blown films of unmodified PBAT/PLA (REF) and the CECL modified samples (V1 to V4). (Paper III)

After CECL modification of the REF blend, the PLA band at 864 cm⁻¹ is slightly shifted for V2, V3 and V4 to smaller wavenumbers up to 850 cm⁻¹ indicating reactions, which enhance vibrations of the α-methyl group. A weak band occurring at 920 cm⁻¹ is characteristic of unsaturated vinyl groups.

Overall, the spectra of V1 to V4 do not differ significantly from REF suggesting only small changes in the chemical structure of PBAT/PLA within the sensitivity limit of FTIR. Furthermore, FTIR performed on the granules does not show significant differences from those obtained on blown films. Thus, regardless of its frequent usage, FTIR may not be an efficient tool to detect the CECL-attributed chemical interactions possibly due to the strong self-interactions of the PBAT and PLA [104].

4.3 Process-induced morphology

In order to study the process-induced morphology, SEM of fracture surfaces of blown extrusion films were made. Figure 39 shows PBAT as the matrix and PLA as the dispersed phase with calcium carbonate being completely dispersed in the PBAT matrix. It can be seen that the dimensions of the dispersed PLA phase differ for ED and TD showing a clear anisotropy.

REF exhibits a clear dispersion of PLA in the PBAT matrix with "sea-island" morphologies < 500 nm, Figure 39 – arrows 1 and 6, confirming nonhomogeneous dispersion. Such morphology is associated with poor mechanical properties due to a weak interfacial adhesion between PBAT and PLA as well as internal stresses at an interface. Furthermore, the skin-core structure is found for REF consisting of coarsely dispersed PLA in the core region of the sample and fine PLA fibrils in the skin regions, Figure 39 – arrows 4 and 5 in TD.

The comparison of fracture surfaces of V1 to V4 with respect to REF shows that the CECLs change the fracture appearance: V1 fails brittle, V2 slightly more ductile, and V3 to V4 significantly more ductile and tougher, Figures 39 – arrow 5. REF and V1 to V3 exhibit fibrils, Figure 39 – arrows 2, 8, 12 and 23, indicating the tough and ductile behavior of the PBAT matrix.

The introduction of CECL modifies the morphology of the PBAT/PLA and the structure of the fracture surfaces with respect to the dimension and geometry of the PLA islands. However, there are morphological similarities in all compounds as the structure of the dispersed PLA is circular or spherical, Figure 39 – arrows 1, 6, 7, 11, 13, 18, 19, 24, 28 and 30. This and the coarser islands support the process of chain scission for V1 and V2.

The calcium carbonate particles having diameters of 2 to 3 μ m (D50 of $1.2 \pm 0.2 \,\mu$ m) are well visible on the fracture surfaces and show a good adhesion with the PBAT matrix, Figure 39 – arrows 3, 5, 21 and 22, indicating a good matrix wetting especially for V4.

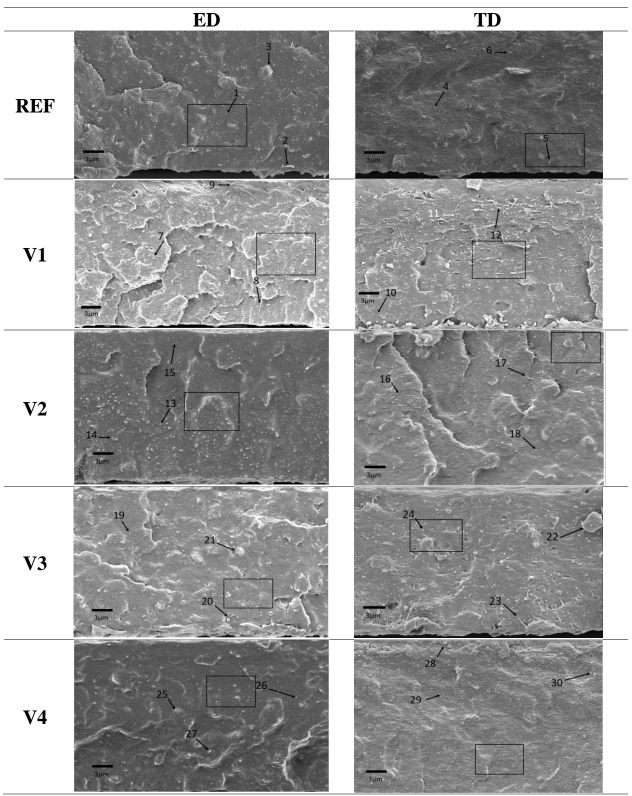


Figure 39: SEM of fracture surfaces of unmodified PBAT/PLA (REF) and CECL modified (V1 to V4) films in ED and TD. The rectangles represent the magnified areas displayed in Figure 40. (Paper II)

The process-induced differences in ED and TD show that PLA fibrils are found only in TD, Figure 39 – arrows 5, 12, 17, 23 and 28, but not in ED. In the ED, the fibrils seem to be decayed in spherical or ellipsoidal PLA islands whereas in TD both spherical PLA islands and fibrils occur. The exception is V4 modified blend, which does not exhibit PLA fibrils in TD which suggests a better interface adhesion, where the fibrils are completely embedded in the matrix.

The morphology of the modified blends has to be interpreted concerning the structure formation during film blowing with BUR of 1:2.5 and drawing ratio (DR) of 12 to 14. During the blowing phase, the elongation flow stretches and orients the melt at the beginning in TD due to blowing, and subsequently in ED due to drawing. Simultaneously, the melt cools down faster as the film is stretched, and therefore the melt viscosity increases. The occurrence of the fibrils suggests that cooling and freezing exceed the orientation relaxation.

At temperatures below 130 °C, PLA starts to crystallize causing freezing in the sea-islands in the morphology. Due to DR >> BUR, the initially spherical PLA particles in the melt are more stretched in ED than in TD explaining the occurrence of PLA fibrils only in TD. After the solidification of PLA, only PBAT matrix can be further stretched until it has been cooled below T_g of PLA at around 60 °C. During this stretching, the calcium carbonate particles orient perpendicular in TD.

At larger magnifications, the fibrils due to fracturing during sample preparation of the PBAT matrix are well visible, Figure 40 – arrows 31, 35, 39, 40, and 46. This can be explained by the lower stiffness and yield stress of PBAT ($E \approx 400$ MPa, $\sigma_{max} \approx 35$ MPa) [105] compared to PLA (E = 3500 MPa, $\sigma_{max} \approx 60$ MPa) [9] leaving the PLA in almost nondeformed state during the fracture. Furthermore, it also enhances the fracture propagation along the PBAT/PLA interface. Rather poor adhesion is indicated with many small PLA particles (D < 200 nm), Figure 40 – arrows 32, 34, 37, 41, 44, 48, 51, 52, 56 and 58.

Besides processing-induced PLA fibrils, one also finds fracture-induced fibrils of the PBAT matrix having 5 to 10 times smaller diameters, Figure 40 – arrows 31, 35, 39, 40, and 46.

The cracks appearing on the fracture surfaces of V2 to V4, Figure 40 – arrows 45, 49, 55 and 59, are presumably not a result of the CECL modification, but of electron beam heating of the surface causing the relaxation of internal stresses and void formation. In V1 and V3, the dispersed PLA is not covered by the PBAT matrix indicating that the CECL hardly affects compatibilization. The increased elliptic shape of the PLA islands could arise from the fact that the process-induced PLA fibrils decay in ellipsoidal droplets, which are frozen in and appear in the morphology.

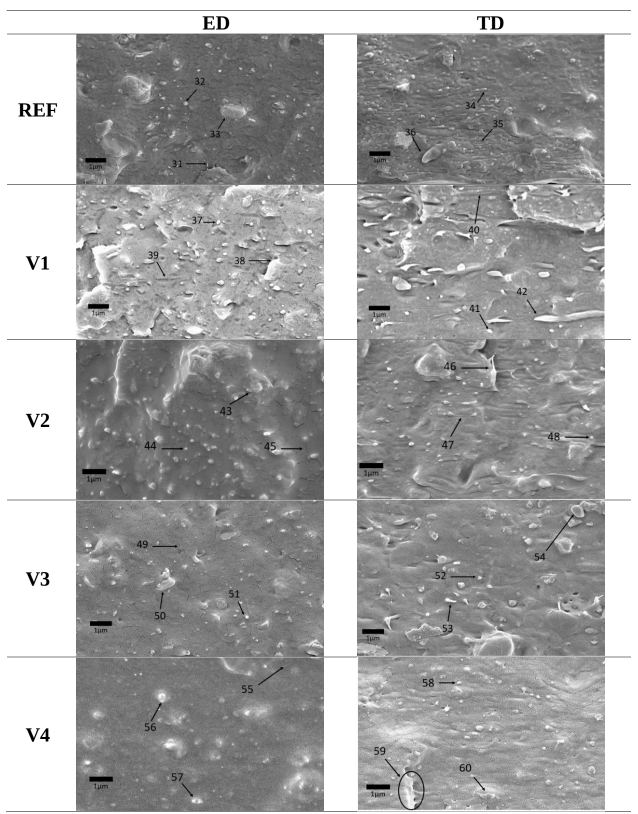


Figure 40: Detailed (magnification of 10,000) SEM of fracture surfaces of unmodified PBAT/PLA (REF) and CECL modified (V1 to V4) films in ED and TD blown film extrusion directions.

The morphological analysis has shown V1 and V3 with a dispersed PLA phase that was not covered by the PBAT matrix, indicating that these two CECLs do not provide compatibilization, whereas V4 showed the dispersed PLA partially covered by the PBAT matrix. The most synergetic effect was obtained for V2 where the PLA phase was well embedded in the PBAT matrix, indicating adhesion and improved compatibilization.

4.3.1 Influence of blow-up ratio dynamic

SEM micrographs (Figure 39 to 42) show cryogenically fractured surfaces of blown films made of PBAT/PLA blends with CECLs, for the two different BUR, 1.5 and 3.5.

Figure 39 presents SEM of fracture surfaces of REF and V1 to V4 films in ED and TD for a BUR of 1.5. It exhibits REF with a clear lamellar distribution in ED and a clear dispersion of PLA in the PBAT matrix with the so-called "sea-island morphologies", see arrows 32, 33 and 34. The same sea-islands are seen in TD, but no lamellar and filament effect as in ED. This comes to confirm the nonhomogeneous dispersion of PLA in PBAT as reported in the literature [30]. The lamellar distribution is attributed to the orientation with a low BUR revealing a high degree of orientation, as reported above by the mechanical properties analysis.

In some cases, voids are noticeable at the interface, indicating low interfacial adhesion between the PBAT and PLA phases associated with poor mechanical properties, which arise from weak interfacial adhesion between the phases as well as stress at the interface boundaries. It seems that the fracture went preferentially through the interface between spherical droplets and matrix. The typical sea-islands morphology in SEM depicts poor miscibility between PBAT and PLA, which is consistent with the observation in other studies [30]. The introduction of CECLs modified the morphology of the PBAT/PLA as the structure of the fracture surfaces changed with respect to the dimension and geometry of the PLA islands, and the lamellar/filamentary structure.

The differences in ED and TD were to be expected, as the orientation level is different in each direction. In ED, a lamellar effect is seen as it seems that orientation is made in layers while in TD, sea-islands of PLA are clearly seen in the matrix in a spherical or ellipsoidal form with the occurrence of fibrils in V2, Figure 48 - see arrows 39, 40 and 41, and V4, Figure 39 - see arrows 47 and 50. For V1 - see arrows 35, 36, 37 and 38, and V3, see arrows 43, 44 and 45, that do not present any fibrils in TD, suggest a better interface adhesion, where the fibrils were completely embedded in the matrix.

Interpreting the morphology of the modified blends with respect of the structure formation during film blowing with a BUR of 1.5, during the blow phase, the elongation flow stretched and oriented the melt at the beginning in TD due to

blowing and subsequently in ED due to drawing. Simultaneously, the melt cooled down the faster the film was stretched and melt viscosity increased. The occurrence of the fibrils suggests that cooling and freezing exceeded the orientation relaxation.

At larger magnifications, Figure 40, the sea-islands morphology and the fibrils due to fracturing of the matrix were very well visible. This can be explained by the lower stiffness and yield stress of PBAT, as mentioned for BUR 2.5, compared to PLA, leaving the PLA in an almost nondeformed state during the fracture. Furthermore, it also enhances the fracture propagation along the PBAT/PLA interface.

In V1, Figure 40 – see arrows 55 and 58, and V3 – see arrows 64, 65 and 66, the dispersed PLA was not covered by the PBAT matrix, indicating that the CECLs hardly affected compatibilization. The more elliptic shape of the PLA islands could have arisen due to the fact that the processing-induced liquid PLA fibrils to decay in ellipsoidal droplets, which were frozen in the current shape.

Further making the analysis with a higher BUR, 3.5, Figure 43 presents SEM of fracture surfaces of REF and V1 to V4 films in ED and TD for a BUR of 3.5.

The analysis on the morphology with different BUR shows that is clear the influence of process-induced changes, and it greatly changes with the different directions. In ED, samples present an elongated ribbon-like PLA phase in the PBAT matrix towards the melt stretching direction (ED), due to the orientation of the polymer chains.

REF presents a lamellar structure in ED, see Figure 43 – arrow 72, with fibrils being extended in this structure. Also, the clear dispersed phase of PLA being embedded in the matrix phase. Fibrils reach to a length extension of 1 µm. In TD, REF presents both spherical PLA islands, see Figure 43 – arrow 73, and fibrils with length higher than 1 µm, see Figure 43 – arrow 74. The morphology of the modified blends has to be interpreted also with respect to structure formation during film blowing. Two steps may take place process-induced morphology during the film extrusion process: first, orientation-induced structure (nuclei) in the melt at the precrystallization stage; and second, the subsequent morphological development based on the first stage nuclei. The final morphology should be mainly dominated by the structure formed in the melt at the initial stages of crystallization [52].

There are morphological similarities in all compounds as the structure of the dispersed phase (PLA) is circular or spherical. V1 presents similar morphology as REF, both in ED and TD, with a lamellar structure in ED with fibrils, see Figure 43 – arrow 76, being extended in this structure, and in TD the dispersed phase of PLA, see Figure 43 – arrow 78, being embedded in the matrix phase and the fibrils reach to a length extension of 1 μ m.

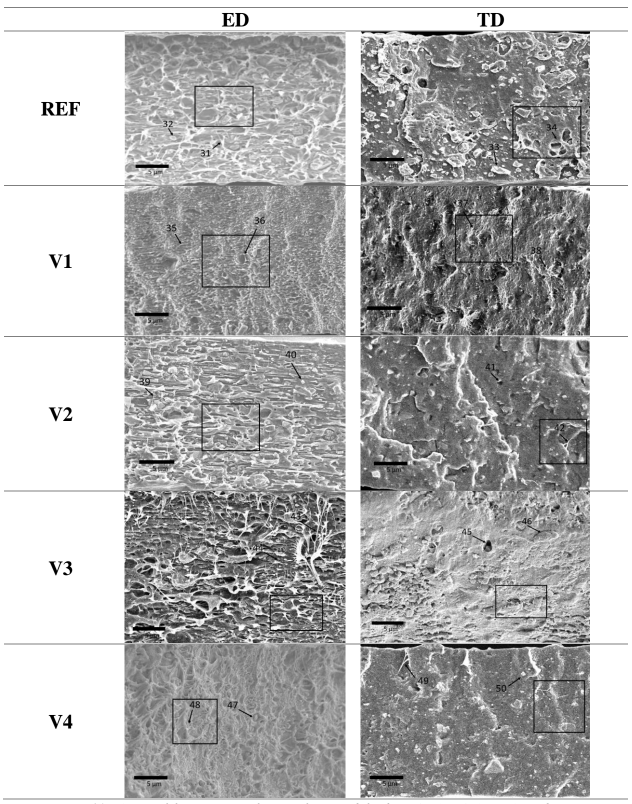


Figure 41: SEM of fracture surfaces of unmodified PBAT/PLA (REF) and CECL modified (V1 to V4) films in ED and TD for a BUR of 1.5. The rectangles represent the magnified areas displayed in Figure 42.

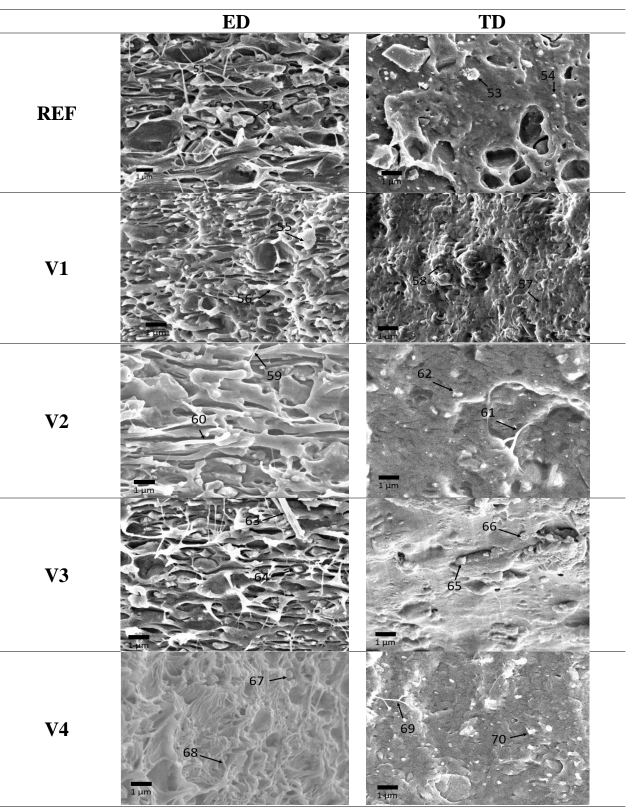


Figure 42: Detailed (magnification of 10,000) SEM of fracture surfaces of unmodified PBAT/PLA (REF) and CECL modified (V1 to V4) films in ED and TD blown film extrusion directions for a BUR of 1.5.

For V2, no fibrils, both in ED and TD can be seen, only the coarser dispersed PLA phase embedded completely on the PBAT matrix, see Figure 43 – arrows 79 to 83. In the case of V3, in ED presents the same morphology as V2 but in TD is possible to see some fibrils formed with a length higher than 1 µm, see Figure 43 – arrow 85. For V4, a very lamellar structure in ED is seen while in TD fibrils are seen and the PLA "sea-islands" are seen dispersed in the matrix phase, see Figure 43 – arrow 89. During the blow phase, the elongation flow stretched and oriented the melt at the beginning in TD due to blowing and subsequently in ED due to drawing. The more the melt cooled down the faster the film was stretched and melt viscosity increased. The occurrence of the fibrils suggests that cooling and freezing exceeded the orientation relaxation.

Low BUR tends to offer greater resistance to tear propagation along the ED and TD. As the polymer molecules exit the film die, the extensional forces applied will tend to orient them along the stresses. Relaxation of the extended conformation of these chains will simultaneously compete with their orientation prior to the crystallization process. Consequently, at high BUR a greater number of molecules will be oriented along the ED before the onset of crystallization. The stronger tensile strength in MD over TD is attributed to the fibrillar structure, which serves as nuclei.

The evaluation with different BUR along ED and TD, have shown a further insight in blown film extrusion of PBAT/PLA blends was that the PLA droplets solidify shortly after being extruded by the die and remain undeformed during blowing. This means that the PLA droplets represent thermoplastic filler particles in the blown films and that the PBAT matrix experiences all deformations.

All BUR undeformed spherical PLA droplets are well visible in ED, Figure 39 to 44 e.g. arrow 32, 33, 37, 44, 51, 56, 57, 64, 67, 92, 95 and 107 and in TD, Figure 39 to 44 arrow 35, 48, 52, 58, 98, 102 and 110 for REF, V1 to V4. Interestingly V4 exhibits in ED at BUR=3.5 a highly ductile fracture surface looking like a ruptured layer structure with embedded PLA droplets.

This ductile fracture behavior in ED can be explained by the high DR of 18 to 19. The macromolecules in the melt are highly oriented in ED but hardly in TD what favors the growth of crystalline lamellae in TD. This may create a pronounced shish-kebab morphology with the succession of crystalline and amorphous lamellae in ED. As at BUR=1.5, the polymer molecules are only little oriented in TD, crystal lamellae growth in ED hardly occurs. The amorphous lamellae can provide a lot of ductility as they are not stiffened by crystalline cross-lamellae.

GPC was also performed to evaluate the molecular mass changes and it showed that the molecular masses of the films of REF, V1 to V4 were not altered due the choice of BUR within the accuracy of measurements. This means that choosing different BUR has negligible effects on the molecular masses of REF, V1 to V4.

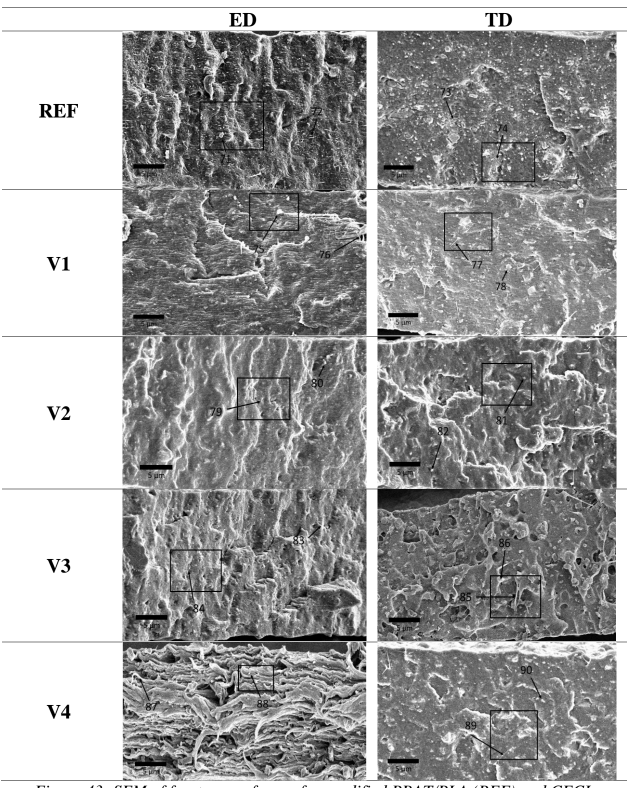


Figure 43: SEM of fracture surfaces of unmodified PBAT/PLA (REF) and CECL modified (V1 to V4) films in ED and TD for a BUR of 3.5. The rectangles represent the magnified areas displayed in Figure 44.

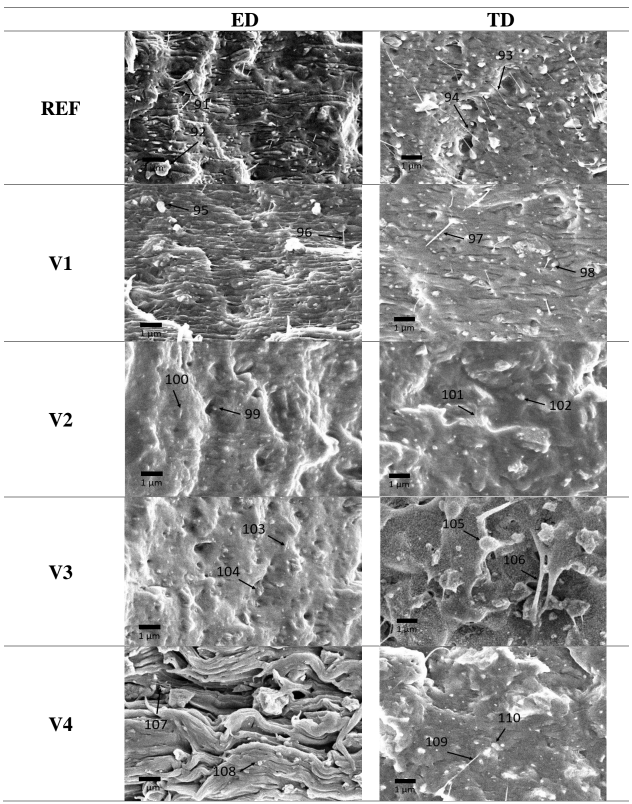


Figure 44: Detailed (magnification of 10,000) SEM of fracture surfaces of unmodified PBAT/PLA (REF) and CECL modified (V1 to V4) films in ED and TD blown film extrusion directions for a BUR of 3.5.

For all BUR, clear "sea island morphologies" –PBAT matrix with embedded PLA droplets – are seen on the fracture surfaces in TD for REF, V1 to V4 what confirms results of other researchers who found inhomogeneously dispersed PLA droplets in the PBAT matrix. The SEM pictures show that the visual impression of the fracture surfaces, and thus the failure behavior during fracturing, changes with BUR significantly and in a characteristic manner for each compound.

4.4 Disintegration in compost and their performance effect

4.4.1 Kinetics of disintegration behavior

The results of the disintegration trials will be presented and discussed in comparison between the different samples and their behavior under composting conditions in each trial series. The measurement of disintegration by mass loss has to be regarded as a rather qualitative assessment.

The visual inspection of films shows, Figure 45, that temperature has a significant effect on the disintegration behavior of the PBAT/PLA compounds modified by the CECLs (Paper III). At 30 °C (Figure 45a), REF, V1 and V2 exhibit small stains on the surface which indicate a starting disintegration after 2 months. At 30 °C PLA is in a glassy state (glass temperature of 60 °C) preventing its degradation.

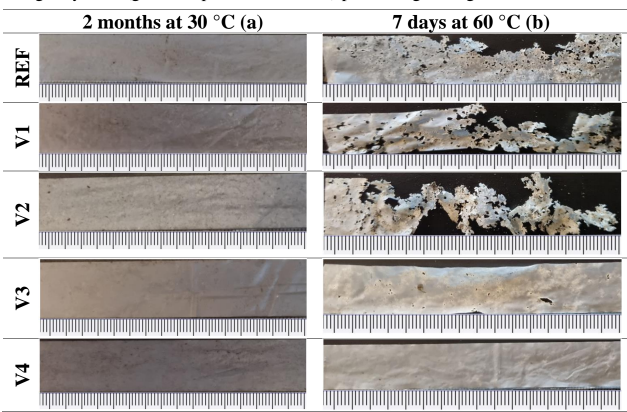


Figure 45. Compost storage for 2 months at 30 °C (a) and 7 days at 60 °C (b) on 25 µm thick films of PBAT/PLA compounds; scaling in mm.

At 60 °C (Figure 45b) REF exhibits a mean disintegration after 7 days, V1 and V2 a severe one, whereas V3 shows starting disintegration, and V4 still seems to be unaffected. The states of disintegration are in line with the increased melt flow ratios due to chain scission of V1 and V2 compared to REF, and the decreased melt flow ratios of V3 and V4 showing cross-linking (Paper I).

By evaluating the relative mass at 30 °C for 8 weeks for REF, V1 to V4, was possible to see that it is relatively unchanged in the range of the scatter of the mass determination, as seen in Figure 46, indicating that the disintegration processes have not led to mass decreases yet. The mass disturbances can be attributed to processes, that occur within the initiation time, e.g. mass increase due to chemical reactions, humidity uptake and remaining compost particles or mass decrease due to the release of degradation reaction or loss of microplastics during cleaning.

At 60 °C the relative mass remains relatively unchanged only for 4 days, Figure 46. Afterwards, for V1 and V2 a pronounced mass decrease is observed. REF exhibits a less pronounced mass decrease between days 5 and 6. V3 shows a tendency of decreasing mass, whereas V4 seems not to disintegrate even after 7 days. This shows that the initiation time is significantly affected by the chosen CECL, and the chemical changes induced with CECL.

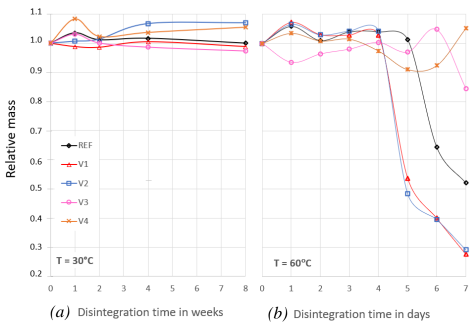


Figure 46. Relative mass change of 25 μ m films of REF, V1 to V4 over 8 weeks at 30 °C (a) and 7 days at 60 °C (b) of disintegration in compost; standard deviations (STD) are between 0.03 and 0.07, the lines serve for better visualization. (Paper III)

Because of the scatter of the mass measurements, the disintegration becomes confirmed only if the masses have decreased to less than 90 % of the initial mass. Thus, the time until the significant mass decrease in Figure 45 represents the upper

limit of the initiation time for these 25 μ m films. It is defined as $t_{0.8}$ being the time when masses reach 80 % of the initial mean mass. In order to determine $t_{0.8}$, the mean masses were calculated in the time range in which the mass remains between 90 and 110 %. From this mean initial mass, the 80 % value was determined and the time $t_{0.8}$ was calculated by linear interpolation between two neighboring mass measurements (Paper III).

After 8 weeks at 30 °C, the REF lost 4 %, and 7 % after 7 days at 60 °C (Paper III). The decrease of V1 and V2 exceeds that of REF showing that the biodegradation of V1 and V2 was more proceeded. For V3 and V4 the decrease is less than for REF showing that crosslinking hinders biodegradation and decelerates disintegration. However, the decreases are still within the scatter of the relative masses during the initiation times. This indicates further that the mass may increase at first due to the uptake and generation of low molecular weight components during the initiation times of storage in compost in addition to adherent compost particles.

For all compounds the DSC traces of the 1st runs differ significantly from the 2nd runs, Figure 47, as well as the enthalpies between 20 and 170 °C. These differences can be attributed to the processing history (which is similar for all films) and the state of biodegradation.

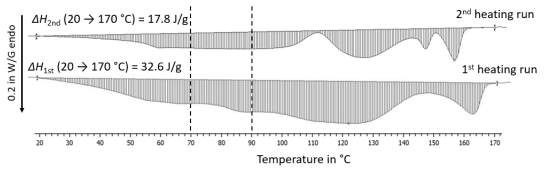


Figure 47. 1st and 2nd heating run of REF after compost storage for 7 days at 60 °C showing the enthalpies 20 and 170 °C. The dashed lines mark the range 70 to 90 °C in which most of the water desorption is expected.

The films stored in compost at 60 °C, Figure 48, were further investigated with respect to their disintegration kinetics by determining the cross-sectional areas of the holes in the films after storage in compost to elucidate the kinetic parameters initiation time t_{init} and disintegration time t_{disint} of REF, V1 to V4. The experimental data can be fitted by equation (7b Appendix A), with reasonable R^2 , Table 14, although the fitting assumes only a single dominating disintegration process. This assumption seems not to be fully applicable for REF, V1 and V2 as the time-dependent increases of the degrees of disintegration x_{disint} is initially smaller than predicted by equation (7b Appendix A). However, this behavior can be explained by the fact that this sudden increase in mass loss is caused by the loss of micro-plastic

particles during sample cleaning if disintegration processes have reached a certain stage.

The times to the mass decrease of 80 % $t_{0.8}$ exceed the initiation times t_{init} by a factor 3 to 4, Table 14. This indicates that the initiation time of the disintegration process depends on the mass and that the holes in the films may occur before the significant mass changes. In the case of the increasing cross-sectional areas of the holes during the storage in compost, several processes might be involved (an uptake of humidity, settlement of micro-organisms on a film surface, chemical modification of polymers, and chain scissions) before the metabolizing of the polymer segments really begins.

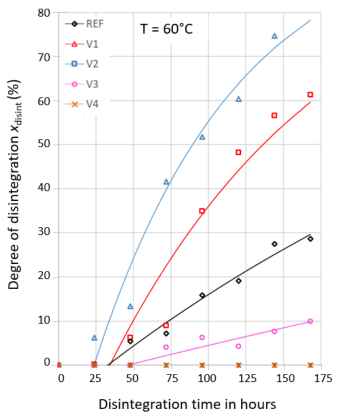


Figure 48. Time dependent degree of disintegration at 60 °C of REF, V1 to V4; comparison of experimental data (symbols) to fit according to equation (7b); STD from 0.03 to 0.05. (Paper III)

Clearly, some degree of disintegration happened before the occurrence of the first holes. Thus, the initiation time t_{init} has to depend on the film thickness. Assuming that the first appearance of holes depends linearly on a film thickness, the initiation time of V1 (30 µm instead of 25 µm) is roughly 20 % too long – leading to a corrected $t_{init} \approx 27$ h. V4 did not show any holes within 7 days indicating that the network structure generated by V4 prevented significant disintegration. This brings the initiation times in line with the times to 80 % of the initial mass. Both effects can

be explained by the fact that the CECL in V1 and V2 cause chain scission, and in V3 and V4 cross-linking, as already investigated above.

The disintegration times τ_{disint} show that the CECL significantly affects the disintegration processes. The disintegration rate of V1 and V2 exceeds that of REF by factors 2.5 and 4, respectively, whereas it is decreased to 0.25 for V3. Regarding V4, as mentioned above, it did not show any holes within 7 days, so it was not evaluated, Table 14.

Table 14 Kinetic parameters of REF and V1 to V3 compounds over 7 days at 60 °C of disintegration in compost.

Kinetic Parameters Compound					
	Unit	REF	V1	V2	V3
Time to 80 % of initial mass $t_{0.8}$	h	132.7	105.6	105.8	174.1
Initiation time t_{init}	h	32.8	34.1	23.7	45.8
Disintegration time $ au_{disint}$	h	385.1	147.3	94.8	1179.5
R ² of fits		0.96	0.95	0.97	0.83

It's possible to conclude that even small amounts of the added CECL significantly alter the disintegration and degradation behavior of REF depending on whether it causes chain scission or crosslinking. Polymer chains have to be segmented to metabolizable lengths by e.g. hydrolytic degradation before biodegradation can start. This corresponds to the initiation time, which is the characteristic quantity for a given blend composition. In the case of crosslinking, V3 and V4, it is increased because the mobility of the polymer chains and the diffusion of the low molecular weight components from the compost to the blown films are reduced due to a denser network.

4.4.2 Disintegration effect on mechanical kinetics

Evaluation was carried out on the film samples during the storage of REF, V1 to V4 in compost for 8 weeks at 30 °C, Figure 49 (Paper III). At first, it led to an increase in Young's modulus before they slowly decreased again. This increase can be explained by annealing and/or post-crystallization effects in the amorphous phase of PLA. As PLA is the disperse phase here, it is protected by the PBAT matrix against degradation at the beginning of compost storage at 30 °C. Therefore, the increases of PBAT moduli may occur already at 30 °C as humidity leads to more mobile polymer chains, and thus enabling a slow post-crystallization. After 2 weeks, *E* was at highest in ED, while in 4 weeks the highest was achieved in TD indicating that the degradation processes during the initiation time affect *E* in ED and TD in a different manner. This means that the orientation of polymer chains and the sites,

that enable hydrolytic and microbial degradation, differ in ED and TD. Thus, during the starting phase, one direction is more affected by biodegradation than the other for the given number of degradation reactions.

Regarding the tensile strengths, for REF, V1 and V2 a continuous decrease in ED was found, whereas V3 and V4 exhibited a plateau of tensile strength for 4 weeks before a decrease of 60 to 70 %. The plateau behavior of tensile strengths for 4 weeks was observed for all compounds in TD combined with a slightly more pronounced subsequent strength decrease in the order of 10 %. Similar behavior was also found for the elongations at break. This concludes that tensile strength and elongation at break are very sensitive to structural changes prior to visible mechanical disintegration. This is confirmed by the fact that the V3 and V4 were cross-linked by the CECLs and exhibited reduced tensile strengths and elongations at break and after compost storage for 4 to 8 weeks at 30 °C, despite no visible disintegration. For V1 and V2, the effect of chain scission is confirmed.

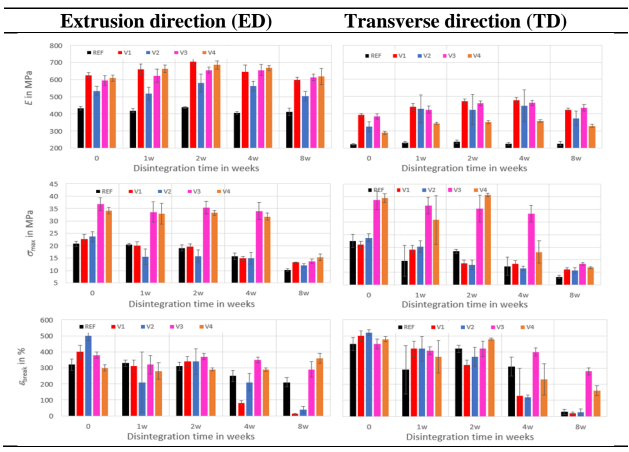


Figure 49. Time dependent effects of disintegration on mechanical properties (Young's modulus E, tensile strength σ_{max} , elongation at break ε_{break}) in extrusion direction (ED) and transverse direction (TD) of films after storage in compost at 30 °C. (Paper III)

With compost storage at 60 °C, Figure 50, has a significantly more pronounced impact on mechanical properties. However, annealing effects cannot occur anymore

as the amorphous phase of PLA is now in the liquid state. The *E* of V1 and V2 in ED and TD were decreased respectively, after 7 days, and close to zero after 14 days, whereas Young's modulus plateaus were observed for REF (3 days), V1 and V2 (14 days) followed by a decrease of 60 % (REF) after 7 days, 70 % (V3) and 20 % (V4) after 28 days. Young's modulus plateaus were found for REF (3 days) with a fast decrease close to zero after 7 days, V3 (14 days) with a decrease of 60 % after 28 days, and V4 (28 days) with no decrease in TD.

The tensile strengths in ED and TD of REF, V1 and V2 were continuously decreased after 7 days. The decreases for V3 and V4 were 70 and 60 %, respectively, after 28 days. Only V4 exhibited an initial strength plateau during 2 to 3 days. The elongations at break in ED and TD of V1 and V2 decreased continuously close to zero after 7 and 3 days, respectively, REF and V3 exhibited a plateau for 1 day before decreasing close to zero after 7 days and 28 days, respectively. Only V4 showed a plateau for 3 days before decreasing by 80 % in ED and close to zero in TD.

The changes of tensile strengths and elongation at break indicate that these properties are more sensitive with respect to degradation and disintegration processes.

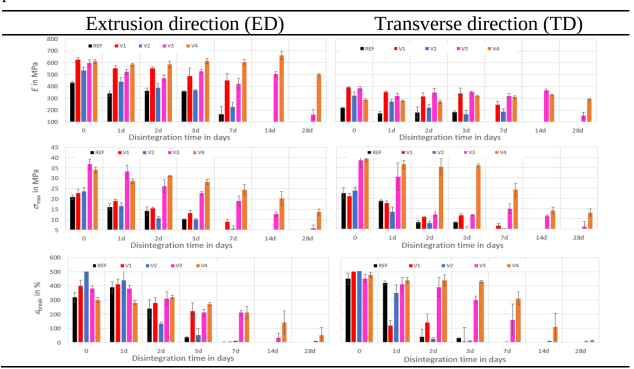


Figure 50. Time dependent effects of disintegration on mechanical properties (Young's modulus E, tensile strength σ_{max} , elongation at break ε_{break}) in extrusion direction (ED) and transverse direction (TD) of films after storage in compost at 60 °C. (Paper III)

It's possible to conclude that REF, V1 to V4 have been mechanically degraded already during compost storage at 30 °C, but visibly these processes were still on an

initiation level with negligible effects on film masses. At 60 °C. the degradation processes were significantly accelerated as REF, V1 and V2 were already severely disintegrated after 7 days (Paper III) and both tensile strengths and elongations at break dropped to low values. In general, the degradation and disintegration behavior are in line with the fact that V1 and V2 experienced chain scission during compounding and processing, and V3 and V4 underwent cross-linking, which decelerates polymer degradation due to fewer attack possibilities at the chains. Thus, shorter and more mobile polymer chains are more prone to microbial degradation.

4.4.3 Disintegration effect on mean molecular masses

Further GPC was used to check if molecular degradation has occurred during compost storage, Table 15. The results confirm the earlier findings of MVR measurements that the CECL cause cross-linking in V3 and V4 as their molecular masses ($M_n \approx 75,000 \text{ g/mol}$) are roughly doubled compared to REF. REF, V1 and V2 with molecular masses around 44,000 g/mol showed that the CECL did not significantly change the molecular masses and as a consequence the macromolecular structure. Most of CECL molecules have to be remained as low molecular components in the polymer. chain scission. The significantly higher MVR values of V1 and V2 compared to REF in Paper I were explained by chain scission. However, according to the GPC results, Table 15, this cannot be the reason for the higher MVR values. They have to be explained by an enhancement of the melt flowability provided the remaining unreacted CECL as low molecular components [106]. However, the GPC curves of REF, V1 to V4 showed single peak chromatograms with PBAT/PLA-ratio dependent peak heights, Figure 51, in contrast to double peak chromatograms observed by Fu et al [105] which were independent of the PBAT/PLA-ratio.

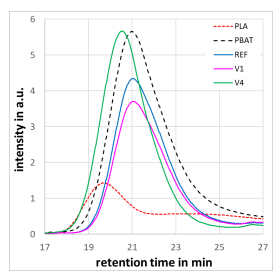


Figure 51. GPC curves of pure PBAT, REF, V1, V4 and pure PLA.

The single peak chromatograms can be explained by fact that the height of the pure PBAT peak exceeds 5 times that of pure PLA. Thus, in REF there remains only a small PLA peak overlaid by the PBAT peak (knowing that REF contains a ratio PBAT/PLA of 76/24 %).

Compost storage at 30 °C did not change the molecular mass of REF, V1, V3 and V4 with respect to the accuracy of the measurement. Only V2 exhibited further cross-linking with increased molecular masses in the order of 10 %. Compost storage at 60 °C led to molecular degradation of REF and V1 after 7 days, and also to further cross-linking of V2 in the order of 10 %. For V3 and V4 molecular degradation was not detected either at 30 °C nor at 60 °C.

Table 15 Determination of the mean molecular masses due to GPC measurements with number-average molecular mass M_n , weight-average molecular mass M_w , and polydispersity index of the polymer PDI; accuracy of measurements is 2,000 g/mol for M_n and 3,000 g/mol for M_w .

Compound	$M_{\rm n}$	$M_{ m w}$	PDI	
	g/mol	g/mol	-	
	Initial state			
PBAT	43,000	107,000	2.45	
REF	44,000	83,000	1.89	
V1	42,000	81,000	1.92	
V2	46,000	92,000	1.98	
V3	75,000	149,000	1.99	
V4	72,000	141, 000	1.95	
PLA	218,000	309, 000	1.42	
After comp	ost storage	for 2 weeks	at 30 °C	
REF	46,000	86,000	1.89	
V1	47,000	87,000	1.88	
V2	52,000	105,000	1.99	
V3	75,000	160,000	2.12	
V4	75,000	148,000	1.98	
After comp	ost storage	for 8 weeks	at 30 °C	
REF	43,000	82,000	1.94	
V1	43,000	83,000	1.92	
V2	51,000	103,000	2.00	
V3	78,000	162,000	2.07	
V4	74,000	148,000	2.01	
After compost storage for 3 days at 60 °C				
REF	41,000	80,000	1.95	

V1	42,000	79,000	1.90
V2	52,000	99,000	1.91
V3	72,000	154,000	2.14
V4	71,000	141,000	1.98
After compost storage for 7 days at 60 °C			
REF	34,000	67,000	1.97
V1	37,000	71,000	1.93
V2	49,000	96,000	1.97
V3	72,000	154,000	2.13
V4	72,000	143,000	1.99

The GPC results clearly show that under the given composting conditions molecular degradation started only after 3 days at 60 °C for REF and V1. This means that the determined mass losses shown in Figure 45 and the growth of the hole area have to be caused by mechanical disintegration towards micro-plasticization.

5. CONCLUSION

In this thesis, the process-induced changes in thermo-mechanical viscoelastic properties and the corresponding morphology of biodegradable polybutylene adipate terephthalate (PBAT) and polylactic acid (PLA) blown film blends modified with various multifunctional chain-extending cross-linkers (CECL) was investigated.

The modification of the PBAT/PLA blend (REF) with the four different CECLs has modified the properties significantly. Tris(2,4-di-tert-butylphenyl)phosphite and 1,3-phenylenebisoxazoline act rather as flow enhancers without changing the molecular masses, evidenced by the gel permeation chromatography (GPC), while aromatic polycarbodiimide and poly(4,4-dicyclohexylmethanecarbodiimide) led to cross-linking. Further, process-induced changes are different for the extrusion direction (ED) and the direction transverse to extrusion (TD). This has been reflected in the elongation at break in ED and tear resistance in TD decreased, whereas elongation at break in the TD and tear resistance in the ED remain unaltered.

Thermal evaluation (with differential scanning calorimetry - DSC) for the granules and films exhibited that the chemical reactions were incomplete after compounding and that film blowing intensifies them, even for 25 μ m films being rapidly cooled. This behavior is explainable by the molecules being linked with one reactive site of polymer chains during compounding, while other reactive sites only show reaction possibilities if a chain slip occurs, such as in elongational flow during film blowing.

According to DMA data, tris(2,4-di-tert-butylphenyl)phosphite hardly affected the glass transition temperature (T_g) values of either segment, whereas other CECLs caused increases in the soft segment phase and decreases in the hard segment phase. This meant that cross-linking occurred in the soft phase, whereas CECL reactions occurred in the hard phase only with increased free volume or CECL.

With the Fourier-transform infrared spectroscopy (FTIR), the PLA band (864 cm⁻¹) was shifted for 1,3-phenylenebisoxazoline, aromatic polycarbodiimide and poly(4,4-dicyclohexylmethanecarbodiimide) to a smaller wavenumber length indicating interactions between PBAT/PLA and these CECLs during the compounding process. From the scanning electron microscopy (SEM) of the unmodified and modified blown extrusion films, the circular/spherical PLA dispersed phase was observed. When PLA starts to crystallize at temperatures below 130 °C, then thus freezes in the current geometry. The spherical PLA particles in the melt are not more stretched in ED than in TD direction due to the substantial large draw ratio than blow-up ratio (BUR), creating the occurrence of PLA fibrils happening only in the TD direction. The investigation with 2 different BURs (1.5 and 3.5) has shown that the PLA droplets with diameters of 200-500 nm solidify shortly after being extruded from the die and remain undeformed during blowing.

This means that the PLA droplets represent thermoplastic filler particles in the blown films and that the PBAT matrix experiences all deformations. This relates to the mechanical performance showing the elongation at break and tear strength in TD to decrease systematically with BUR, whereas an arbitrary dependence was obtained for other properties such as Young's modulus, tensile strength, and seal strength. The anisotropic morphology introduced during film blowing has presented its effects during the degradation processes. Their disintegration behavior was significantly altered concerning the unmodified reference blend. Finally, in the process of blown film extrusion, the elongational behavior represents a very important characteristic. However, its evaluation may be quite often problematic, and its accuracy dramatically differs from that of shear viscosity. By using an SER Universal Testing Platform, results have shown that for very low extensional rates, an elongational process is dominated by disentanglement of the studied materials preventing strain hardening. By comparing the rupture time intervals, it is possible to evaluate mutually the chosen materials. It was confirmed that out of four chain-extending cross-linkers, the best contribution to blown film processing was exhibited by poly(4,4-dicyclohexylmethanecarbodiimide).

To conclude, this work has brought more and better insight into how process-induced properties of PBAT/PLA blends function with modified CECLs and the synergy of it with blow-film processing by evaluating the main characteristics to improvement for the packaging industry.

6. CONTRIBUTION TO SCIENCE AND PRACTICE

Bioplastics make it possible to develop innovative, alternative solutions compared to conventional plastics. Biodegradable plastics allow enhanced end-of-life scenarios for disposal and recycling. The packaging industry is currently the largest user of bioplastics, but there are many other suitable applications possible, and consumers are increasingly interested in alternative bio-solutions. To reach this goal, materials have to be improved and optimized in terms of processing-induced properties, to achieve the right balance of performance. A better understanding of the processing-induced properties will lead to optimized blend films.

This PhD thesis is focused on the question of how process-induced properties are affected by the introduction of four different CECLs. A comprehensive study on the influence of reactive extrusion (compounding) aligned with different blow-up ratios on the blown film extrusion was performed. To interpret the results, several models/ideas are presented because direction comparison to the results of other researchers could not be made due to the lack of literature correlating specific behaviors and their conjugated effects of CECL influenced with processing (anisotropy in extrusion/transversal directions). The following topics of this PhD thesis are considered the most important contribution to both science and practice:

- 1. The investigation of the effects of four multifunctional chain-extending cross-linkers on the processability, mechanical performance, and structure of PBAT and PLA blends to provide an insight into the reactive extrusion of different functional groups and their effect on mechanical and thermal performance and morphological effects.
- 2. The insight of the different blow-up ratio and the comparison with the mechanical and morphological performance to determine the correct set-up to provide the best suitable end-use.
- 3. The study of extension characteristics by using an SER Universal Testing Platform provides good insight that elongation behavior represents an important characteristic in the blown film extrusion process, thus leading to a better understanding of extensional rates and their correlation to the strain hardening behavior.

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ABBREVIATIONS

AA	Adipic acid
ATR	Attenuated Total Reflectance
BDO	Butane diol
BUR	Blow-up ratio
CECL	Chain extending cross-linkers
CO_2	Carbon dioxide
CTE	Coefficient of thermal expansion
DIM	Deutsches Institut für Normung (in English, German Institute for
DIN	Standardisation)
DCP	Dicumyl peroxide
DMA	Dynamic mechanical analysis
D-PLA	Poly(D-lactic acid)
DR	Draw ratio
DSC	Differential scanning calorimetry
ED	Extrusion direction
ETBP	Epoxy terminated branched polymer
FTIR	Fourier-transform infrared spectroscopy
GPC	Gel permeation chromatography
H ₂ O	Water
HDPE	High density polyethylene
HIPS	High impact polystyrene
IR	Infrared
ISO	International Standard Organization
IUPAC	International Union of Pure and Applied Chemistry
LDPE	Low density polyethylene
L-PLA	Poly(L-lactic acid)
MDI	4,4 methylene diphenyl diisocyanate
MVR	Melt Volume Rate
OL	Organosolv lignin
PA 6	Polyamide 6
PBAT	Polybutylene adipate terephthalate
PDI	Polydispersity index

PBS	Polybutylene succinate
PCL	Polycaprolactone
PDLA	Poly(D,L-lactic acid)
PE	Polyethylene
PHA	Polyhydroxyalkanoate
PHB	Polyhydroxybutyrate
PHV	Polyhydroxyvalerate
PLA	Polylactic acid
PP	Polypropylene
PS	Polystyrene
PTA	Terephthalic acid
SAXS	Small-angle X-ray scattering
SEM	Scanning electron microscopy
SER	Sentmanat Extensional Rheometer
TD	Transverse direction
THF	Tetrahydrofuran
WAXS	Wide-angle X-ray scattering
wt.%	Weight percentage

SYMBOLS

c_p	Heat capacity
E	Young's modulus
E'	Storage modulus
E''	Loss modulus
\mathcal{E}_{break}	Strain at break
F_{break}	Rupture force
F_{seal}	Seal strength
F_{tear}	Tear resistance
L	Length
M_n	Number molecular mass
M_w	Weight molecular mass
T	Temperature
$tan\delta$	Loss factor
T_{cr}	Crystallization temperature
T_{end}	End temperature
T_g	Glass transition temperature
$T_{g,hs}$	Glass temperature of hard segments
$T_{ m g,ss}$	Glass temperature of soft segments
T_m	Melting temperature
T_{ml}	Melting temperature of PBAT
T_{m2}	Melting temperature of PLA
T _{start}	Starting temperature
W	Width
ΔH_{cr}	Crystallization enthalpy
ΔH_m	Melting enthalpy
ΔH_{ml}	Melting enthalpy of PBAT
ΔH_{m2}	Melting enthalpy of PLA
η_E	Elongational viscosity

$\sigma_{\!f}$	Fracture stress
$\sigma_{\!f}$	Fracture stress
σ_{max}	Tensile strength
ρ	Density
λ	Heat conductivity

PUBLICATIONS

Publications in context of this doctoral work:

- J. V. C. Azevedo, E. R. Dorp, B. Hausnerová, B. Möginger, The Effects of Chain-Extending Cross-Linkers on the Mechanical and Thermal Properties of Poly(butylene adipate terephthalate)/Poly(lactic acid) Blown Films. *Polymers* 13 (2021) 3092.
- J. V. C. Azevedo, E. R. Dorp, R. Grimmig, B. Hausnerová, B. Möginger, Process-Induced Morphology of Poly(Butylene Adipate Terephthalate) /Poly(Lactic Acid) Blown Extrusion Films Modified with Chain-Extending Cross-Linkers. *Polymers* 14 (2022) 1939.
- J. V. C. Azevedo, B. Hausnerová, B. Möginger, T. Šopík, Effect of chain extending crosslinkers on disintegration behavior of composted PBAT/PLA blown films. *Int. J. Mol. Sci.* 24 (2023) 4525.
- P. Filip, B. Hausnerová, B. Möginger, J. V. C. Azevedo, Improvement of elongational behavior of poly(lactic acid)/poly(butylene adipate terephthalate) blends in blown film extrusion using chain extending cross-linkers. Submitted to *Int. J. Biol. Macromol.*, November 2023, November 2023.
- J. V. C. Azevedo, E. R. Dorp, C. Kössler, C. Thomser, T. Šopík, B. Möginger B. Hausnerová, Blow molding-induced structure and performance of renewable PLA/PBAT packaging films modified with chain-extending cross-linkers. Submitted to *J. Ind. Eng. Chem.*, November 2023.

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APPENDIX A

For better comparison of mass changes of REF, V1 to V4, the data are converted to relative masses

$$m_{rel}(t) = \frac{m(t)}{m_0} \tag{1}$$

with mass $m_{(t)}$ after a given time of compost storage and initial mass m_0 .

Kinetics of the process of disintegration in compost due to the decrease of film cross-section

After the given disintegration times the samples were taken out of the compost, cleaned with a soft paint brush to remove compost particles, and subsequently stored at 23 °C/50 % r.h to establish a constant water content. Then the pictures of the sample films were taken with a camera Sony Cyber-shot DSC-HX60V.

The sample films appear in greyish colors, whereas holes appear rather in black. Then the pictures were discriminated to black and white pixels and the corresponding pixel areas were calculated using the public domain image analysis software Fiji [38]. The degree of disintegration x_{disint} was determined by

$$x_{disint}(t) = \frac{A_{black}(t)}{A_{white}(t) + A_{black}(t)}$$
 (2)

with the black pixel area A_{black} and white pixel area A_{white} .

The quantitative kinetic approach assumes that the micro-organisms from the compost attack the film at the contact areas and start disintegration processes from the surface. As the compost has a coarse and crumble structure, the disintegration happens not uniformly on the surface but locally at the certain points. After an initiation time t_{init} , first holes appear in the films which grow in size and number, thus, decreasing the remaining film cross-section. If one assumes that the time dependent change of film cross-section $\frac{dA^{film}}{dt}$ depends on the available film cross-section A^{film} one gets

$$\frac{dA^{film}}{dt} \sim A^{film}(t) = A_0 - A^{hole}(t) \tag{3}$$

with hole area A^{hole} .

After dividing by the initial cross-section A_0 one gets the normalized rate equation with the temperature dependent disintegration constant k_{disint}

$$\frac{dA_{rel}^{film}}{dt} = -k_{disint}(T) A_{rel}^{film}(t)$$
(4)

Solving by separation of variables and subsequent integration over equivalent limits leads to

$$\ln A_{rel}^{film}(t) = -k_{disint}(T) t + C \tag{5}$$

with the integration constant C=0. Solving for $A_{rel}^{film}(t)$ and substitution by $x_{disint}(t)$ yields

$$x_{disint}(t) = 1 - e^{-k_{disint}(T)t}. ag{6}$$

To take into account the *initiation time* t_{init} of the disintegration process, one has to modify equation (6) by

$$x_{disint}(t) = 1 - e^{-k_{disint}(T)(t - t_{init})}.$$
or
(7a)

$$x_{disint}(t) = 1 - e^{-\frac{(t - t_{init})}{\tau_{disint}}}.$$
 (7b)

with the disintegration time τ_{disint} being a measure how fast the disintegration process proceeds.

A least square fitting procedure of equation (7b) with the excel solver was used to determine t_{init} and τ_{disint} to quantify the disintegration kinetics.

APPENDIX B

Before measurements of elongational viscosities using the SER device, two principal conditions should be checked: mild intensity of sample sagging during the process of temperature setting and an acceptably small deviation from exponential decrease of sample cross-sections. For REF and V1 to V4, both conditions are strongly violated as documented in Figure 1.





Figure 1. Deformation of REF rectangular samples: at 140 °C (left), at 150 °C (right).

Other restriction of the usage of SER Universal Testing Platform can be caused if the sample does not rupture prior to the completion of the first revolution of the SER drums. However, this restriction can be ignored for thin samples with good adherence to the drums. In this case, a sample does not have to overcome an abrupt increase of drum diameter caused by clamps when starting the second revolution. It experiences only a negligible diameter increase if the sample is adequately fixed at its ends (an increase from zero to the thickness of the lateral ends). In our case the stretched film samples have to overcome only a gradual gentle slope, increasing the drum diameter by approximately 2 %.

The sagging behavior is governed by sample temperatures and the corresponding solid-like and liquid-like behavior. If the temperature interval between these two states is narrow as in the case of PLA/PBAT compounds, then the SER device can hardly be applied for measurements of elongational viscosities. This results from the fact that the measuring temperature should be established without temperature overshoots to maintain the morphological structure of the films. This requires long temperature setting times (the final temperature achieved by a series of temperature settings from below) that may lead to unacceptable sagging. Moreover, not only sagging limits the application of the SER device but also the overhanging of upper film parts leading to thickening along the upper film edge. A combination of these two effects prevents the proper determination of elongational viscosities. These phenomena diverge from the principal assumptions under which the SER device can be applied: exponentially decaying uniform sample thickness across the whole active narrowing rectangular area.