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(71) Applicant (for all designated States except US): BORE-ALIS TECHNOLOGY OY [FI/FI]; P.O. Box 330, FIN-06101 Porvoo (FI).

(72) Inventors; and

(75) Inventors/Applicants (for US only): GREIN, Christelle [AT/AT]; Harrachstrasse 28, A-4020 Linz (AT). WOLFS-BERGER, Anton [AT/AT]; Eschenweg 2, A-4210 Gallneukirchen (AT). NIEDERSÜSS, Peter [AT/AT]; Blindendorf 221, A-4312 Ried/Riedmark (AT).

(74) Agent: KADOR & PARTNER; Corneliusstr. 15, 80469 Munich (DE).

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(54) Title: TRANSPARENT EASY TEARABLE FILM

(57) Abstract: The present invention relates to a cast film or blown film comprising a composition comprising 80 to 95 wt% of a polypropylene matrix (A) with an MFR₂ of 1 to 30 g/10min being a homopolymer or a copolymer, which has a comonomer content of less than 3 wt%, 5 to 20 wt% of an ethylene-propylene-rubber (EPR) (B) with at least 55 wt% propylene having an intrinsic viscosity (IV) of 1 to 2.5 dl/g; and 0.0001 to 1 wt% of an α-nucleating agent.

Transparent easy tearable film

The present invention relates to a cast film or blown film having a high transparency, easy tear features and low extractables. Furthermore, the invention is related to a process for making the films as well as to the use of a heterophasic propylene composition for cast films and blown films.

In the recent years there has been an increasing demand of heat-sealed pouches, in particular in the food industry, for example for rice or fodder for dogs and cats. A very well-known problem of such filled pouches is their opening. Often, the pouches can only be opened with a high effort at a predefined indentation (or perforation or notch) without using tools like knives and scissors. Another problem is that a manual opening does not lead to a straight crack in the pouches but to a curved one increasing the risk that the content of the pouches ooze out unintended. Hence, there is a demand to develop films for pouches which can be easily opened manually at the notch leading to a straight crack in the pouch. Described with physical parameters, it means that the tear energy has to be minimized and that the crack should develop straight in a predefined direction starting from the notch and usually in parallel direction to the upper side of the pouch. The materials used for such purpose have to be tough enough to resist impact loading at room or sub-ambient temperatures. Resins like polypropylene homopolymers and ethylene-propylene random copolymers which are brittle under severe loading are therefore excluded. Impact polypropylenes which are known for their high toughness are normally hazy and tend to develop plasticity when deformed, a property which limits the propagation of a linear crack through the pouch. Thus, they are no candidates of choice. Additionally the film shall fulfill certain standards issued by the food and drug administration (FDA), i.e. inter alia having low C₆-extractables. Certainly the transparency of the film should also be good as the market favors transparent pouches.

Therefore, it is the object of the present invention to provide a cast film or blown film having a high transparency, easy tear features and low extractables.

The finding of the present invention is to provide a cast film or a blown film comprising a composition of a polypropylene matrix and an ethylene-propylene rubber (EPR) whereby the particles of the ethylene-propylene rubber (EPR) in the matrix develop a high orientation in one direction when the film is casted or blown.

Hence, the present invention is related to a cast film or blown film comprising a composition comprising

- a) 80 to 95 wt% of a polypropylene matrix (A) with an MFR₂ of 1 to 30 g/10 min being a homopolymer or a copolymer which has a comonomer content of less than 3 wt%;
- b) 5 to 20 wt% of an ethylene-propylene rubber (EPR) (B) with at least 55 wt% propylene having an intrinsic viscosity (IV) of 1 to 2.5 dl/g; and
- c) 0.0001 to 1 wt% of an alpha-nucleating agent (C).

Surprisingly, it had been found that such a cast film or blown film possesses a high transparency, low extractables and also very good tear features expressed in a low tear resistance, low tear whitening, low tear plasticity and a low tear angle.

Definition of tear features

Tear resistance, tear whitening, tear plasticity and tear angle have been used to describe the easy-tear performance of the different materials. Tear resistance (W), also called tear energy in the following, is that energy needed to propagate a crack in a precut sample at room temperature according to ISO 6383-2. It is commonly known as Elmendorf test. Tear resistance was determined at room temperature on 75*63*0.1 mm or 75*63*0.05 mm samples with a precut slit of 20 mm according to ISO 6383-2. The force required to tear the specimen completely along its length was used to calculate the tear resistance.

Tear whitening (TW), tear plasticity (TP) and tear angle (TA) have been determined post mortem on the fractured samples used to assess the tear energy within 48 hours after having performed the Elmendorf test. Values provided in the following are valid for these fractured samples and correspond to about half of what would have been measured considering the whole samples (which consist of two broken halves).

The tear whitening corresponds to the degree of whitening of the area around the fracture surface. It has been determined with a ruler and rationalized with marks: 0 correspond to no tear whitening at all, 1 to a tear whitening < 0.3 mm, 2 to a tear whitening in between 0.3-0.5 mm (0.5 mm excluded), 3 to a tear whitening in between 0.5-1.0 mm (1.0 mm excluded), 4 to a tear whitening in between 1.0-2.0 mm (2.0 mm excluded), 5 to a tear whitening above 2.0 mm (2 mm included).

The tear plasticity corresponds to the size of the damaged area around the fracture surface. This damaged zone is often called plastic zone. Depending on the polymer composition it will or won't exhibit whitening. It has been determined with a ruler and rationalized with marks: 0 correspond to no plasticity at all, 1 to a localized damage near the crack which size is < 1 mm, 2 to a damage localized in a zone of 1-2 mm around the main crack (2 mm excluded), 3 to a damage localized in a zone of 2-5 mm around the main crack (5 mm excluded), 4 to a damage localized in a zone of 5-10 mm around the main crack (10 mm excluded), 5 to a damage localized in a zone of $\ge 10 \text{ mm}$ around the main crack.

The tear angle corresponds to the deviation of the crack towards a straight propagation. It has been measured with a protractor.

All the above mentioned parameters have to be fulfilled to consider a material as easy tearable:

- \Rightarrow W_{MD} < 20 N/mm, the tear resistance in the machine direction;
- \Rightarrow W_{TD} /W_{MD} > 2, where W_{TD} is the tear resistance in the transverse direction;
- \Rightarrow TW \leq 2 in both machine and transverse directions where TW is the tear whitening;

 \Rightarrow TP \leq 2 in both machine and transverse directions where TP is the tear plasticity;

 \Rightarrow TA \leq 2.5° in machine direction (TA_{MD}) and TA \leq 4.5° in transverse direction (TA_{TD}) where TA is the tear angle.

Usually the notch (precut) is made in the machine direction, since the energy needed to propagate the crack is there the lowest. However, if $W_{TD} < 20 \text{ N/mm}$ (preferably < 15 N/mm) and $TA_{TD} < 2.5^{\circ}$, the transverse direction is also perfectly adapted as tear direction.

To achieve these very good tear features, it is in particular necessary that the ethylene-propylene rubber (EPR) particles are expanded in one direction, i.e. that they are oriented in the direction in which the crack develops. To achieve this aim, a composition is needed as defined above. Hence the synergistic effect of all features contributes to a deformation of the ethylene-propylene rubber (EPR) particles in the propylene matrix leading to the orientation effect, which is mainly responsible for the good tear features but has also a positive influence on the transparency and low extractable properties.

An important requirement in this context is the intrinsic viscosity of the ethylene-propylene rubber (EPR).

The intrinsic viscosity (IV) is a measure of the capability of a polymer in solution to enhance the viscosity of the solution. Intrinsic viscosity (IV) is correlated positively with the polymer molecular weight. The intrinsic viscosity (IV) is defined as the limiting value of the specific viscosity/concentration ratio at zero concentration. It thus becomes necessary to find the viscosity at different concentrations and then extrapolate to zero concentration. The variation of the viscosity number with concentration depends on the type of molecules as well as on the solvent. In general, the intrinsic viscosity of linear macromolecular substances is related to the molecular weight or degree of polymerization. With linear macromolecules, viscosity number measurements can provide a method for the rapid determination of molecular weight when the relationship between viscosity and molecular weight has been established.

The intrinsic viscosity (IV) in this application is measured according to DIN ISO 1628-1 (October 1999).

Accordingly, if the intrinsic viscosity (IV) of the ethylene-propylene rubber (EPR) is too high, i.e. higher than 2.5 dl/g, then the film will have a too high impact strength, which influences the tear resistance negatively, i.e. the tear energy is rather high. Moreover a high intrinsic viscosity (IV) is also correlated with rather big particles (with a diameter above 500 nm as observed on electron microscopy micrographs). Big particles, however, diminish the transparency of the film. On the other hand, if the intrinsic viscosity (IV) is too low, i.e. lower than 1.0 dl/g, then a production of the composition is rather difficult.

It is in particular preferred, that the intrinsic viscosity (IV) is not higher than 2.3 dl/g, more preferably not higher than 2.2 dl/g and most preferably not higher than 2.1 dl/g. In turn, the lower limit of the intrinsic viscosity (IV) is preferably not lower than 1.2 dl/g, more preferably not lower than 1.4 dl/g. Preferred ranges are 1.0 to 2.3 dl/g, more preferably 1.2 to 2.3 dl/g, still more preferably 1.2 to 2.2 dl/g and most preferably 1.2 to 2.1 dl/g.

Preferably, not only the intrinsic viscosity (IV) of the ethylene-propylene rubber (EPR) alone should be considered, but also the polypropylene matrix used. It should be in particular avoided that the ethylene-propylene rubber (EPR) has a similar intrinsic viscosity (IV) to the polypropylene matrix. A good elongation of the ethylene-propylene rubber (EPR) is especially achieved if the intrinsic viscosities of the ethylene-propylene rubber (EPR) particles and the polypropylene matrix differ to some extent. Therefore to further improve the orientation effect, it is preferred that the polypropylene matrix has a higher intrinsic viscosity (IV) than the ethylene-propylene rubber (EPR). This proportion of the polypropylene matrix and the ethylene-propylene rubber (EPR) to each other enables a specifically good elongation of the ethylene-propylene rubber particles in the polypropylene matrix. It has been, in particular, found out that it is especially preferred that the ratio of the intrinsic

viscosity (IV) of the ethylene-propylene rubber (EPR) to the polypropylene matrix (IV_{EPR}/IV_{matrix}) is not higher than 0.85, more preferably not higher than 0.8. Preferably, the polypropylene matrix has an intrinsic viscosity (IV) of 1.8 to 4.0 dl/g, more preferably of 2.0 to 3.5 dl/g.

As stated above the intrinsic viscosity (IV) is of particular relevance in this application, as it influences the orientation effect needed. On the other hand the intrinsic viscosity (IV) is an indicator of the molecular weight. A high intrinsic viscosity (IV) results from a high molecular weight. Another way to express the molecular weight of a polymer is by way of its melt flow rate (MFR). The melt flow rate mainly depends on the average molecular weight. This is due to the fact that long molecules render the material a lower flow-tendency than short molecules. An increase in molecular weight means a decrease in the MFR-value. The melt flow rate (MFR) is measured in g/10 min of the polymer discharged through a defined dye under specified temperature and pressure conditions and the measure of viscosity of the polymer which, in turn, for each type of polymer is mainly influenced by its molecular weight but also by its degree of branching. The melt flow rate measured under a load of 2.16 kg (ISO 1133) is denoted as MFR₂. Hence as both measuring methods are interlinked by the molecular weight of the polymer, also the melt flow rate (MFR) has to be carefully considered.

Therefore it is required, that the polypropylene matrix shall have a specific melt flow rate (MFR). If the melt flow rate MFR₂ is too low (high molecular weight polymer matrix), i.e. lower than 1 g/10min, then, the easy tear features are far from target due to the high ductility of the matrix which promotes plastic deformation. Hence, the polypropylene matrix shall have an MFR₂ of not lower than 1 g/10min, more preferably not lower than 1.5 g/10min. On the other hand, if the MFR₂ is too high, i.e. higher than 30 g/10min, then the polypropylene matrix has inferior transparency and can not easily been processed. Moreover a too high melt flow rate (MFR₂) of the polypropylene matrix boosts the particle size of the ethylene-propylene rubber (EPR) which is

also detrimental for the orientation effect as well as for the transparency of the film. It is in particular preferred, that the MFR₂ of the polypropylene matrix is not higher than 20 g/10 min, more preferably not higher than 10 g/10min.

In addition it is preferred, that the composition has a MFR₂ of at least 1.5 g/10 min, more preferably of at least 2 g/10min.

The type of polypropylene matrix is not restricted to a specific one, as long as the matrix fulfills the above mentioned requirements. Hence, each homopolymer or copolymer can be used. In case a copolymer is used, the comonomer content, however, shall be less than 3 wt%, otherwise the film gets too ductile affecting negatively the easy tear features. Preferably, the comonomer is ethylene.

The choice of ethylene-propylene rubber (EPR) must conform with the orientation requirement and the transparency targeted. Hence an ethylene-propylene rubber (EPR) is needed having a high propylene amount, i.e. a propylene amount of at least 55 wt% in the rubber. More preferably the amount of propylene in the ethylene-propylene rubber (EPR) is at least 58 wt%, more preferably at least 60 wt%.

Moreover to achieve good tear features the weight ratio of the polypropylene matrix and the ethylene-propylene rubber (EPR) must be kept in a specific range. Accordingly, the composition does not have more than 20 wt% of the ethylene-propylene rubber (EPR), otherwise the film will be too soft and a linear crack cannot be guaranteed anymore. Therefore the amount of polypropylene matrix must range from 80 to 95 wt%, preferably in the range of 82 to 93 wt%. In turn, the ethylene-propylene rubber (EPR) must be in the range of 5 to 20 wt%, preferably in the range of 7 to 18 wt%.

To achieve the desired properties of the present invention, it has been further found out, that also the film must have a high cristallinity. A highly structured (crystal) polymer has improved optical and mechanical properties, i.e. a good tear performance expressed by a low tear energy.

Therefore the composition employed must comprise an alpha-nucleating agent in the amount of 0.0001 to 1 wt%, preferably in the amount of 0.001 to 0.5 wt%.

Alpha-nucleating agents of polypropylene are substances with the ability to act as substrates for crystal formation by epitaxy thus increasing the number of nuclei of the system. Thereby the nucleating agents improve the crystallization behavior in processing, i.e. decreasing the cycle time or increasing the line speed, increase the crystallinity in the final product and, additionally, improve the optical and mechanical properties in terms of improved stiffness and here of lower tear energy.

Any suitable alpha-nucleating agent known in the art can be used. However, it is preferred to use at least one alpha-nucleating agent selected from the group consisting of sodium benzoate, 1,3:2,4-bis-(3,4-dimethyl-benzylidene)-sorbitol, sodium-2,2'-methylenebis-(4,6-di-tert-butyl-phenyl)-phosphate and hydroxybis-(2,4,8,10-tetra-tert-butyl-6-hydroxy-12h-dibenzo-(d,g)(1,3,2)-dioxaphosphocin-6-oxidato)-aluminium. A further α -nucleation method, herein referred to as "BNT", is a special reactor technique, where the catalyst is prepolymerised with monomers like vinylcyclohexane (VCH). This method is described in greater detail in e.g. EP 0 316 187 A2. For the purpose of this invention "BNT" is referred to as α -nucleating agent.

Preferably, the thickness of the film should not be too high since otherwise a significant loss in transparency will be the effect. Therefore, it is preferred that the film has a thickness of 5 to 200 μ m, more preferably of 10 to 150 μ m.

Preferably, the film shall have only few C_6 -extractables measured according to FDA-HHS 177.1520 where 2.5g of the sample are extracted at 50°C with 1 l hexane in a Soxhlet apparatus for 24 h. It is in particular preferred that the amount of C_6 -extractables shall be equal or lower than 2.6 wt%, more preferably equal or lower than 2.4 wt%.

It is in particular preferred that the film of the present invention has a tear resistance in machine direction (W_{MD}) determined by ISO 6383-2 of lower than 20 N/mm, more preferably of lower than 15 N/mm, most preferably of lower than 12.5 N/mm when measured on 100 μ m thick films.

More preferably, the inventive film has a W_{TD}/W_{MD} -ratio of more than 2, more preferably of more than 2.3, whereby W_{TD} is the tear-resistance of the film in a transverse direction determined by ISO 6383-2 when measured on 100 μ m thick films.

In addition, it is preferred that the film has a tear-angle (TA) in machine direction of equal or less than 2.5°, more preferably equal or less than 2.2°, most preferably equal or less than 2.0° when measured on 100 µm thick samples after having performed a test according to ISO 6383-2. Moreover, it is preferred that the film has tear angle (TA) in transverse direction of less than 4.5°, more preferably less than 4.0° when measured on 100 µm thick samples after having performed a test according to ISO 6383-2.

Further, it is preferred that the film has a haze of lower than 21%, more preferably lower than 18%, most preferably lower than 15%, determined according to ASTM D 1003-92 when measured on 100 μ m thick films.

Preferably, the film has a gloss of better than 42%, more preferably better than 48%, most preferably better than 52%, determined according to ISO 2813 when measured on $100 \mu m$ thick films.

Moreover, the present invention comprises the use of the composition as defined above for producing a cast film or blown film. In particular, it is preferred that the composition and the cast film or blown film as defined above is used for producing a pouch, label film, twist film, form-fill-and-seal (FFS) film and vertical-bag-form-fill-and-seal (VFFS) film.

Furthermore, the present invention is related to a process for producing a cast film as defined above comprising the steps of

(i) pouring or spreading a solution, hot melt or dispersion of a material onto a temporary carrier,

- (ii) hardening the material, and
- (iii) stripping the hardened film from the surface of the carrier, whereby the material is a composition as described above, in particular a composition comprising
- a. 80 to 95 wt% of a polypropylene matrix with an MFR₂ of 1 to 30 g/10min being a homopolymer or a copolymer, which has a comonomer content of less than 3 wt%;
- b. 5 to 20 wt% of an ethylene-propylene-rubber (EPR) with at least 55 wt% propylene having an intrinsic viscosity (IV) of 1 to 2.5 dl/g; and
- c. 0.0001 to 1 wt% of an α -nucleating agent.

Further, the present invention is related to a process for producing a blown film as described above thereby comprising the steps of

- (i) blowing up a tube of molten material with air perpendicularly to the upwards direction from a side-fed blown film die;
- (ii) cooling it down with air from the outside on a cooling ring;
- (iii) folding it and guiding it over deflector rolls onto the winder
 wherein the material is a composition as defined above, in particular a
 composition comprising
- a) 80 to 95 wt% of a polypropylene matrix with an MFR₂ of 1 to 30 g/10 min being a homopolymer or a copolymer, which has a comonomer content of less than 3 wt%;
- b) 5 to 20 wt% of ethylene-propylene rubber (EPR) with at least 55 wt% propylene having an intrinsic viscosity (IV) of 1 to 2.5 dl/g; and
- c) 0.0001 to 1 wt% of an alpha-nucleating agent.

Alternatively, the blown films may be produced by tubular quench technology with water cooling.

The propylene polymer blends of the present invention are preferably produced by combining the propylene polymer A in the form of powder or granules, the elastomeric copolymer B, and the α -nucleating agent C in a melt mixing device.

Melt mixing devices suited for this process are discontinuous and continuous kneaders, twin screw extruders and single screw extruders with special mixing sections and co-kneaders. The residence time must be chosen such that a sufficiently high degree of homogenisation is achieved.

Production of polypropylene matrix A

The propylene polymer may be produced by single- or multistage process polymerisation of propylene or propylene and α -olefin and/or ethylene such as bulk polymerisation, gas phase polymerisation, slurry polymerisation, solution polymerisation or combinations thereof using conventional catalysts. A homoor copolymer can be made either in loop reactors or in a combination of loop and gas phase reactor. Those processes are well known to one skilled in the art.

A suitable catalyst for the polymerisation of the propylene polymer is any stereospecific catalyst for propylene polymerisation which is capable of polymerising and copolymerising propylene and comonomers at a temperature of 40 to 110°C and at a pressure from 10 to 100 bar. Ziegler Natta catalysts as well as metallocene catalysts are suitable catalysts.

One skilled in the art is aware of the various possibilities to produce propylene homo- and copolymers and will simply find out a suitable procedure to produce suitable polymers which are used in the present invention.

Production of ethylene-propylene rubber

An ethylene-propylene-rubber (B) may be produced by known polymerisation processes such as solution, suspension and gas-phase polymerisation using conventional catalysts. Ziegler Natta catalysts as well as metallocene catalysts are suitable catalysts.

A widely used process is the solution polymerisation. Ethylene, propylene and catalyst systems are polymerised in an excess of hydrocarbon solvent. Stabilisers and oils, if used, are added directly after polymerisation. The solvent and unreacted monomers are then flashed off with hot water or steam, or with mechanical devolatilisation. The polymer, which is in crumb form, is dried with dewatering in screens, mechanical presses or drying ovens. The crumb is formed into wrapped bales or extruded into pellets.

The suspension polymerisation process is a modification of bulk polymerisation. The monomers and catalyst system are injected into the reactor filled with propylene. The polymerisation takes place immediately, forming crumbs of polymer that are not soluble in the propylene. Flashing off the propylene and comonomer completes the polymerisation process.

The gas-phase polymerisation technology consists of one or more vertical fluidised beds. Monomers and nitrogen in gas form along with catalyst are fed to the reactor and solid product is removed periodically. Heat of reaction is removed through the use of the circulating gas that also serves to fluidise the polymer bed. Solvents are not used, thereby eliminating the need for solvent stripping, washing and drying.

The production of ethylene propylene rubber (B) is also described in detail in e.g. US 3,300,459, US 5,919,877, EP 0 060 090 A1 and in a company publication by EniChem "DUTRAL, Ethylene-Propylene Elastomers", pages 1-4 (1991).

Alternatively, ethylene-propylene rubber (B), which are commercially available and which fulfil the indicated requirements, can be used.

Alternatively, polymers A and B may be produced in a series of reactors, i.e. starting with the production of polymer A in a loop reactor and transferring the product into a gas phase reactor, where copolymer B is polymerised.

It is additionally preferred that the films are oriented by monoaxial or biaxial orientation processes. In this way the tearability can be enhanced even further, especially beyond that of standard block-copolymers.

In another embodiment the polymer composition can be coextruded to afford a multi-layered film. The main layer thereby preferably comprises the described polymer composition, while one or more outside layers can be added. It is preferred that the outside layers have the same or a lower tearability. Exploiting this arrangement leads to highly tearable films. The outside layer can also be a thin layer of an especially transparent RACO (random copolymer) film.

Description of measurement methods

MFR

The melt flow rates were measured under a load of 2.16 kg at 230°C. The melt flow rate is that quantity of polymer in grams which the test apparatus standardized to ISO 1133 extrudes within 10 minutes at a temperature of 230°C under a load of 2.16 kg.

Intrinsic Viscosity

Intrinsic viscosity was measured according to DIN ISO 1628-1 (October 1999) in Decalin at 135°C.

Gloss

Gloss was determined according to ISO 2813 on 100 μm thick films at an angle of 20°.

Haze

Haze was determined according to ASTM D 1003-92 on 50 and 100 μm thick films. C_6 -solubles

C₆-solubles were determined according to FDA-HHS 177.1520 where 2.5 g of the sample are extracted at 50°C with 11 n-hexane in a Soxhlet apparatus for 24h.

Tear Resistance

Tear resistance was determined at room temperature on 75*63*0.1 mm or 75*63*0.05 mm samples with a precut slit of 20 mm according to ISO 6383-2 (Elmendorf test). The force required to tear the specimen completely along its length was used to calculate the tear resistance.

Tear whitening

Tear whitening was determined on the fractured samples used to measure the tear resistance according to an internal Borealis method. It has been assessed with a ruler and rationalized with marks: 0 corresponds to no tear whitening at all, 1 to a tear whitening < 0.3 mm, 2 to a tear whitening in between 0.3-0.5 mm (0.5 mm excluded), 3 to a tear whitening in between 0.5-1.0 mm (1.0 mm excluded), 4 to a tear whitening in between 1.0-2.0 mm (2.0 mm excluded), 5 to a tear whitening above 2.0 mm (2 mm included).

Tear Plasticity

Tear plasticity was determined on the fractured samples used to measure the tear resistance according to an internal Borealis method. It has been assessed with a ruler and rationalized with marks: 0 correspond to no plasticity at all, 1 to a localized damage near the crack which size is < 1 mm, 2 to a damage localized in a zone of 1-2 mm around the main crack (2 mm excluded), 3 to a damage localized in a zone of 2-5 mm around the main crack (5 mm excluded), 4 to a damage localized in a zone of 5-10 mm around the main crack (10 mm excluded), 5 to a damage localized in a zone of ≥ 10 mm around the main crack

Tear Angle

Tear angle was determined on the fractured samples used to measure the tear resistance according to an internal Borealis method. It corresponds to the deviation of the crack towards a straight propagation. It has been measured with a protractor.

Dart Drop Impact

Dart Drop Impact (DDI) was determined according to ISO 7765-1 at room temperature. The energy (here expressed in form of weight in g) that causes films to fail under specified conditions of impact of a free-falling dart from a height of 660 mm that would result in 50% failure of the specimen is determined (F_{50}) . 20 specimens were tested for each weight.

Production of films

Cast films were produced:

A) on a single screw extruder with a barrel diameter of 30 mm and a slot die of 200 x 0.5 mm in combination with a chill- and a take up-roll. Melt temperature was 260 °C in the die; the chill-roll was kept at 15 °C and the take-up roll at 15 °C. The film thickness was adjusted through the ratio between extruder output (4,5 kg/h) and takeoff speed (10,5 m/min),

B) on a single screw extruder with a barrel diameter of 60 mm and a slot die of 800×0.9 mm in combination with a chill- and a take up-roll. Melt temperature was 250 °C in the die; the chill-roll was kept at 30 °C and the take-up roll at 22 °C. A film thickness of $50 \, \mu m$ was adjusted through the ratio between extruder output ($60 \, \text{kg/h}$) and takeoff speed ($30 \, \text{m/min}$).

Blown films were produced on a single screw extruder with a barrel diameter of 60 mm and a tubular die of 200 mm diameter x 1.2 mm die gap. Melt temperature was 190 °C in the die; the cooling air was kept at 15°C internal and external of the film bubble. A film thickness of 50 µm was adjusted through the ratio between extruder output (60 kg/h), blow up ratio (2:1) takeoff speed (20 m/min).

Examples

Preparation of polymers A

The propylene polymers A1-A4 and A6 used for the present invention were prepared according to the following procedure:

Raw Materials:

Hexane dried over molecular sieve (3/10A)

TEAL: 93 % from Sigma-Aldrich

Donor: Dicyclopentyldimethoxysilane: ex Wacker Chemie (99%).

N₂: supplier AGA, quality 5.0; purification with catalyst BASF R0311, catalyst

G132 (CuO/ZNO/C), molecular sieves (3/10A) and P₂O₅.

Propylene: polymerisation grade

Hydrogen: supplier AGA, quality 6.0

The catalyst ZN104 is commercially available from Basell.

Sandostab P-EPQ is commercially available from Clariant.

A 51 autoclave reactor has been purified by mechanical cleaning, washing with hexane and heating under vacuum/ N_2 cycles at 160 °C. After testing for leaks with 30 bar N_2 over night reactor has been vacuumed and filled with 1110 g propylene by weighing and 8 nl H_2 by pressure monitoring from a 50 l steel cylinder.

10 mg of ZN104-catalyst are activated for 10 minutes with a mixture of Triethylaluminium (TEAl; solution in hexane 1mol/l) and Dicyclopentyldimethoxysilane as donor (0.3 mol/l in hexane) - in a molar ratio of 5 after a contact time of 5 min - and 10 ml hexane in a catalyst feeder. The molar ratio of TEAl and Ti of catalyst is 250. After activation the catalyst is spilled with 300 g propylene into the stirred reactor with a temperature of 23 °C. Stirring speed is hold at 250 rpm. After 6 min prepolymerisation at 23 °C temperature is increased to 70 °C in about 14 min. After holding that

temperature for 1 hour polymerisation is stopped by flashing propylene and

cooling to room temperature.

After spilling the reactor with N₂ the homopolymer powder is transferred to a

steel container and stabilized with 0.1 wt% of Sandostab P-EPQ and 0.2 wt%

of Ionol in acetone and dried over night in a hood and additionally for 2 hours

at 50 °C under vacuum.

The amount of polymer powder (A3) was 113 g and the MFR (230 °C, 2.16 kg)

of the powder was 5 g/10 min.

Procedure for random copolymer

The propylene copolymer A5 used for the present invention was prepared

according to the following procedure:

Raw Materials:

Hexane dried over molecular sieve (3/10A)

TEAL: 93 % from Sigma-Aldrich

Donor: Dicyclopentyldimethoxysilane: ex Wacker Chemie (99%).

N₂: supplier AGA, quality 5.0; purification with catalyst BASF R0311, catalyst

G132 (CuO/ZNO/C), molecular sieves (3/10A) and P₂O₅.

Propylene: polymerisation grade

Ethylene: polymerisation grade

Hydrogen: supplier AGA, quality 6.0

Sandostab P-EPQ is commercially available from Clariant.

A 51 autoclave reactor has been purified by mechanical cleaning,

washing with hexane and heating under vacuum/N₂ cycles at 160 °C. After

testing for leaks with 30 bar N2 over night reactor has been vacuumed and

filled with 1110 g propylene by weighing and 5 nl H₂ by pressure monitoring

from a 50 l steel cylinder.

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8 mg of ZN104-catalyst are activated for 10 minutes with a mixture of Triethylaluminium (TEAl; solution in hexane 1 mol/lDicyclopentyldimethoxysilane as donor (0.3 mol/l in hexane) - in a molar ratio of 25 after a contact time of 5 min - and 10 ml hexane in a catalyst feeder. The molar ratio of TEAl and Ti of catalyst is 250. After activation the catalyst is spilled with 300 g propylene into the stirred reactor with a temperature of 23 °C. Stirring speed is hold at 350 rpm. After 6 min prepolymerisation at 23 °C temperature is increased to 70 °C in about 14 min. While heating the ethylene flow has been raised continuously up to the final flow of 340 ml/min during the prepolymerisation. After holding a polymerisation temperature of 70°C for 60 min at constant temperature, constant ethylene flow and constant stirrer rotation polymerisation is stopped by stopping the ethylene flow, by flashing propylene and cooling down the reactor to room temperature.

After spilling the reactor with N_2 the homopolymer powder is transferred to a steel container and stabilized with 0.1 wt% of Sandostab P-EPQ and 0.2 wt% of Ionol in acetone and dried over night in a hood and additionally for 2 hours at 50 °C under vacuum.

The amount of polymer powder (A5) was 98 g, the MFR (230 °C, 2.16 kg) of the powder was 3 g/10 min and its C2-content was 4.5 wt%.

The following homopolymers and random copolymers were prepared analogously according to the above procedure (Table 1):

Table 1. Characteristics of homopolymers and random copolymers.

	MFR	C2-content
	[g/10min]	[wt%]
A1	3	0
A2	1,5	0
A3	5	0
A4	10	0
A5	3	4.5
A6	2.5	0

Preparation of ethylene-propylene rubber (B)

The elastomeric copolymers of the present invention were prepared according to the following procedure:

A 5 l-reactor (autoclave) filled with about 0.2 barg propylene (polymerisation grade) is pressured up with the required amount of H₂ in order to achieve the targeted intrinsic viscosity of the elastomeric copolymer. Then 300 g of propylene are added.

5 mg of a ZN101 (supplied by Basell) catalyst is contacted with 0.3 ml white oil for about 16 hours and then activated for 5 minutes with a mixture of Triethylaluminium (TEAl; solution in hexane 1 mol/l) and an alkoxysilane (Dicyclopentyldimethoxysilan in the examples) as donor (0.3 mol/l in hexane) - in a molar ratio of 76 using a contact time of 5 min. The molar ratio of TEAl and Ti of catalyst was 380 and TEAl concentration in TEAl/donor mixture 12.6 mg/ml hexane. After activation the catalyst is transferred to the reactor by

spilling in with 500 g propylene. After 12 min pre-polymerisation at 30 °C a specified amount of ethylene is added to the reactor and the temperature is increased to the target polymerisation temperature (55 °C in the examples). During heating up additional ethylene dosing is started to achieve the target total pressure at the target polymerisation temperature. Total pressure is hold constantly via continuously dosing of ethylene during polymerisation. 30 min after end of prepolymerisation the reaction is stopped by flashing of monomers and cooling.

The polymer is stabilized with 0.1 wt% of Sandostab P-EPQ and 0.2 wt% of Ionol in acetone and dried over night in a hood and additionally for 2 hours at 50 °C under vacuum.

Some example of polymerisation are provided in Table 2

Table 2. Polymerisation characteristics of ethylene-propylene rubbers.

Polymer	H2	C2	total	IV	C3
No.			pressure		
	barg	g	[barg]	[dl/g]	[wt %]
B1	3,7	60	30	1,8	71
B2	2,8	60	31	2,3	60
B4	1,6	50	30	3,2	56
B5	1,76	50	30	3,0	57

The following elastomeric ethylene-propylene copolymers were prepared according to the above procedure(s), except that H_2 and ethylene amounts were varied to achieve different intrinsic viscosities and comonomer concentrations. They are reported in Table 3.

Table 3. Characteristics of ethylene-propylene rubbers.

	IV	C3
	[dl/g]	[wt%]
B1	1,8	71
B2	2,3	60
B3	2,6	62
B4	3,2	56
B5	3	57
В6	1,3	65
B7	2,05	76
B8	1,95	62.5
В9	2,1	53
B10	2,0	43

The amounts of each component and the results of the measurements are shown in Tables 4 and 5. Tests were performed on 100 μ m thick cast films obtained according to method A.

Table 4. Polymer composition.

	Total	Component A		Component	В	Nucleation
ļ	MFR	Type	Content	Type	Content	
	[g/10min]		[wt%]		[wt%]	
E1	3,5	A1	86,4	B1	13,6	Y
CE1	1,2	A2	87,5	B2	12,5	N
CE2	3,5	A3	87	B3	13	Y
CE3	3	A4	82	B4	18	Y
CE4	0,8	A2	84,5	B5	15,5	Y
CE5	3,5	A5	89	В6	11	Y
E2	3,4	A1	89,1	B1	10,9	Y
E3	3,2	A1	91,8	B1	8,2	Y
E4	3,1	A1	94,6	B1	5,4	Y
CE6	3,5	A3	89,6	В3	10,4	Y
CE7	3,4	A3	92,2	В3	7,8	Y
CE8	3,2	A3	94,8	В3	5,2	Y
E5	2,5	A6	88,2	В7	11,8	Y
E6	2,5	A6	87,4	B8	12,6	Y
CE9	2,5	A6	86,6	В9	13,4	Y
CE10	2,5	A6	88,6	B10	11,4	Y
CE11	2,5	A6	88,2	B7	11,8	N
CE12	2,5	A6	87,4	B8	12,6	N
CE13	2,5	A6	86,6	В9	13,4	N
CE14	2,5	A6	88,6	B10	11,4_	N

Table 5. Performance of the different polymers.

Results have been obtained on 100 μm thick cast films produced according to method A.

	Elmendorf						Optical performa	nce	FDA .			
	Machine		on		Transver	se dire	ction					C6-
	W _{MD}	TA _{MD}	TW_{MD}	TP_{MD}	W _{TD}	TA_{TD}	TW_{TD}	TP _{TD}	W _{TD} /W _{MD}	Gloss20°	Haze	solubles
	[N/mm]	[°]	[-]	[-]	[N/mm]	[°]	[-]	[-]	[-]	[%]	[%]	[wt%]
E1	12,2	1,0	1	0	29,4	1,2	0	0	2,4	54,7	12,3	2,0
CE1	45,5	2,7	0	2	195,2	14,2	0	5	4,3	28	25,7	1,7
CE2	14,1	1,6	2	0	28,2	1,6	3	1	2,0	3,7	74,2	1,9
CE3	30,8	2,0	4	3	58,9	4,9	5	4	1,9	3,3	75,5	1,2
CE4	37,3	1,0	0	2	173,6	15,0	0	5	4,7	31,1	34	1,2
CE5	111,7	3,4	0	3,5	177,8	6,5	0	5	1,6	59,2	6,8	3,8
E2	10,7	0,8	1	0	26,3	0,4	0,5	0	2,5	54,5	12,2	na
E3	9,5	0	1	0	21,8	0,7	0,5	0	2,3	54,3	12,4	na
E4	8,1	0	0	0	20	0,6	0	0	2,5	55,5	12,5	na
CE6	13,9	1	2	1	21,5	1,2	3	1	1,5	8,3	68	na
CE7	12,2	1,3	2	1	18,6	1	2	1	1,5	16	58,2	na
CE8	8,6	1,5	0	1	16,7	1,1	1	1	1,9	30,3	44,7	na
E5	16,2	1,6	1	1	38,5	4,2	1	1	2,4	58,1	11,2	1,8
E6	19,9	1,6	1	1	43,1	3,8	1	2	2,2	57,9	12,1	1,6
CE9	57,5	2,5	3	4	34	3,7	3	2	0,6	57,2	14,1	1,5
CE10	80,7	3	3	5	41,9	4	3	2	0,5	54,5	9	1,4
CE11	40,3	3,2	0	2	107,9	2,7	0	3	2,7	51,5	16,6	1,8
CE12	38,3	2,8	0	3	100,5	3,5	0	3	2,6	50	16,9	1,6
CE13	57,5	2,6	0	5	74,2	5,8	0	4	1,3	48,6	18	1,5
CE14	1	2,8	1	5	51,9	4,6	_1	4	0,7	55,9	17,7	1,4_

In addition, cast and blown films of 50 μ m thickness of E1 and CE1 were tested to assess the difference of the different technologies. Results are listed in Table 6. A remarkable DDI performance for blown films can be seen with E1.

			E1 cast film A	cast film B	blown film	CE1 cast film A	cast film B	blown film
Elmendorf	W_{MD}	[N/mm]	19,2	5,9	3,7	45,4	23,5	4,0
	$\mathbf{W_{TD}}$	[N/mm]	190,6	30,6	10,4	194,4	165,4	36,3
Optics	Haze	[%]	6,9	4,0	14,2	17,9	15,5	36,3
DDI	Weight	[g]	>1200	303,0	174,0	>1200	>1200	176,0

Claims

- 1. A cast film or blown film comprising a composition comprising
 - a. 80 to 95 wt% of a polypropylene matrix (A) with an MFR₂ of 1 to 30 g/10min being a homopolymer or a copolymer, which has a comonomer content of less than 3 wt%;
 - b. 5 to 20 wt% of an ethylene-propylene-rubber (EPR) (B) with at least 55 wt% propylene having an intrinsic viscosity (IV) of 1 to 2.5 dl/g; and
 - c. 0.0001 to 1 wt% of an α -nucleating agent.
- 2. A film according to claim 1, wherein the comonomer is ethylene.
- 3. A film according to claim 1 or 2, wherein at least one α-nucleating agent is selected from the group consisting of sodium benzoate, 1,3:2,4-bis-(3,4-dimethyl-benzylidene)-sorbitol, sodium-2,2'-methylenebis-(4,6-di-tert-butyl-phenyl)-phosphate and hydroxybis-(2, 4,8, 10-tetra-tert-butyl-6-hydroxy-12h-dibenzo-(d, g) (1,3,2)-dioxaphosphocin-6-oxidato)-aluminium, or those based on prepolymerisation of the catalyst with monomers like vinylcyclohexane (VCH).
- 4. A film according to any one of the preceding claims 1 to 3, wherein the polypropylene matrix has a higher intrinsic viscosity (IV) than the ethylene-propylene-rubber (EPR).
- 5. A film according to any one of the preceding claims, wherein the polypropylene matrix has an intrinsic viscosity (IV) of 1.8 to 4.0 dl/g;
- 6. A film according to any one of the preceding claims, wherein the ratio (IV_{EPR}/IV_{matrix}) of the intrinsic viscosity (IV) of the ethylene-propylene rubber (EPR) to the polypropylene matrix is not higher than 0.85.
- 7. A film according to any one of the preceding claims, wherein the film has C₆-extractables equal or lower than 2.6 wt%.
- 8. A film according to any one of the preceding claims, wherein the composition has an MFR₂ of at least 2.0 g/10min.

9. A film according to any one of the preceding claims, wherein the film has a thickness of 5 to 200 μm.

- 10. A film according to any one of the preceding claims, wherein the film has a tear resistance in the machine direction (W_{MD}) determined by ISO 6383-2 of lower than 20 N/mm when measured on 100 μm thick samples.
- 11. A film according to any one of the preceding claims, wherein the film has a W_{TD}/W_{MD}-ratio of more than 2, whereby W_{TD} is the tear resistance of the film in the transverse direction determined by ISO 6383-2 when measured on 100 μm thick samples.
- 12. A film according to any one of the preceding claims, wherein the film has a tear angle (TA) in machine direction equal or less than 2.5° and a tear angle (TA) in transverse direction of less than 4.5° when measured on 100 μm thick samples after having performed a test according to ISO 6383-2.
- 13. A film according to any one of the preceding claims, wherein the film has a haze of lower than 21 % determined according to ASTM D 1003-92 when measured on 100 µm thick samples.
- 14. A film according to any one of the preceding claims, wherein the film has a gloss of better than 42 % determined according to ISO 2813 when measured on 100 µm thick samples.
- 15. Process for producing a cast film according to any one of the preceding claims 1 to 14, comprising the steps of
 - i) pouring or spreading a solution, hot-melt or dispersion of a material onto a temporary carrier
 - ii) hardening the material, and
 - iii) stripping the hardened film from the surface of the carrier whereby the material is a composition comprising:
 - a. 80 to 95 wt% of a polypropylene matrix with an MFR₂ of 1 to 30 g/10min being a homopolymer or a copolymer, which has a comonomer content of less than 3 wt%;
 - b. 5 to 20 wt% of an ethylene-propylene-rubber (EPR) with at least 55 wt% propylene having an intrinsic viscosity (IV) of 1 to 2.5 dl/g; and
 - c. 0.0001 to 1 wt% of an α -nucleating agent.

16. Process for producing a blown film according to any one of the preceding claims 1 to 14, comprising the steps of

- (i) blowing up a tube of molten material with air perpendicularly to the upwards direction from a side-fed blown film die;
- (ii) cooling it down with air from the outside on a cooling ring;
- (iii) folding it and guiding it over deflector rolls onto the winder wherein the material is a composition comprising:
- a. 80 to 95 wt% of a polypropylene matrix with an MFR₂ of 1 to 30 g/10min being a homopolymer or a copolymer, which has a comonomer content of less than 3 wt%;
- b. 5 to 20 wt% of an ethylene-propylene-rubber (EPR) with at least 55 wt% propylene having an intrinsic viscosity (IV) of 1 to 2.5 dl/g; and
- c. 0.0001 to 1 wt% of an α -nucleating agent.
- 17. Process according to claim 15 or 16, wherein the composition is further defined by any one of the claims 2 to 8.
- 18. Use of a composition for producing a cast film or blown film, wherein the composition comprises
 - a. 80 to 95 wt% of a polypropylene matrix with an MFR_2 of 1 to 30 g/10min being a homopolymer or a copolymer, which has a comonomer content of less than 3 wt%;
 - b. 5 to 20 wt% of an ethylene-propylene-rubber (EPR) with at least 55 wt% propylene having an intrinsic viscosity (IV) of 1 to 2.5 dl/g; and
 - c. 0.0001 to 1 wt% of an α -nucleating agent.
- 19. Use according to claim 18, wherein the composition is further defined by any one of the claims 2 to 8.
- 20. Use of the film according to any one of the claims 1 to 14 as a pouch, label film, twist film, form-fill-and-seal (FFS) film and vertical-bag-form-fill-and-seal (VFFS) film.

INTERNATIONAL SEARCH REPORT

International application No PCT/EP2006/004129

A. CLASSIFICATION OF SUBJECT MATTER INV. C08J5/18 C08L23/10 B29D7/01 B29C47/00 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) C08J C08L B29D B29C Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Category' Citation of document, with indication, where appropriate, of the relevant passages X 1 - 20WO 2004/055101 A (BOREALIS TECHNOLOGY OY; TORGERSEN, ULF; DE MINK, PAUL; BERNREITNER, KL) 1 July 2004 (2004-07-01) claims 1-9 examples 1-3,5; table 1 page 5, line 14 - page 8, line 20 page 8, line 26 - line 28 EP 0 863 183 A (CHISSO CORPORATION) 1 - 20X 9 September 1998 (1998-09-09) claims 1,14,25-30 page 25; examples 7-12; table 3
page 23, line 14 - line 27 page 15 - page 18; table 1 US 6 747 103 B1 (VESTBERG TORVALD ET AL) 1 - 20Α 8 June 2004 (2004-06-08) the whole document Further documents are listed in the continuation of Box C. See patent family annex. Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention earlier document but published on or after the international filing date "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled document referring to an oral disclosure, use, exhibition or other means in the art. document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 30/06/2006 23 June 2006 Authorized officer Name and mailing address of the ISA/ European Patent Office, P.B. 5818 Patentlaan 2 NL – 2280 HV Rijswijk Tel. (+31–70) 340–2040, Tx. 31 651 epo nl, Fax: (+31–70) 340–3016 Lippert, S

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