(54) Title: ORIENTED HDPE FILMS WITH SKIN LAYERS

(57) Abstract

Readily castable high density polyethylene (HDPE) films, preferably containing broad molecular weight distribution HDPE resin having treated upper skin layers and untreated lower skin layers, preferably coextruded, on each side. The treated upper skin layer is derived from a polymer, e.g. ethylene-propylene-butene-1 terpolymer, and the lower skin layer is derived from a polymer such as ethylene-propylene-butene-1 terpolymer and LDPE compounded with silicone oil. Such skin layers provide a film which is readily castable while maintaining the desirable properties of HDPE including dead fold, twist retention, TD tear and moisture barrier. The films are cast, and oriented in the solid state up to about two times in the machine direction and six times or more in the transverse direction to give films having good dead-fold characteristics making them highly suited for packaging, particularly for dry foods.
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ORIENTED HDPE FILMS WITH SKIN LAYERS

This invention relates to thermoplastic films and more particularly to films of oriented high density polyethylene (HDPE) which have good dead-fold characteristics, twistability, printability and receptivity of water-based coatings.

Blown films of HDPE having an ethylene-vinyl acetate heat seal coating are used for food packaging but such films must have a thickness of about two mils to meet the water vapor transmission (WVTR) requirements for packaging suitable for dry foods such as cereals. Moreover, blown HDPE films do not exhibit the dead-fold properties desirable in food packages particularly of the bag-in-box type.

U.S. Patents Nos. 4,870,122 and 4,916,025 describe HDPE films which have good WVTR properties at film thicknesses of about one mil. The films are produced from high density polyethylene (HDPE) having a density of 0.96 or higher and are biaxially oriented in an imbalanced manner to a degree of 1.25:1 to 2:1 in the machine direction (MD) and to a degree of 6:1 to 12:1 in the transverse direction (TD). These films have reduced water vapor transmission (WVTR), improved dead-fold characteristics and other physical characteristics which are markedly better than blown HDPE films which can be further improved by the addition of microcrystalline wax. When provided with a heat seal layer by co-extrusion or coating, the films are particularly well suited for use in packaging foods in bag-in-box operations conducted on vertical, form, fill and seal (VFFS) machinery. While the imbalanced orientation confers the desired overall combination of properties on the film, the high degree of orientation in the transverse direction tends to make the film weak in this direction so that it tears relatively easily.

The present invention relates to providing a film having good dead fold properties, optical clarity, good slip properties, and good receptivity to water based coatings, including water based inks, water based adhesives and other common water based coatings such as polyvinylidene chloride (PVDC), and acrylics.
The present invention relates to a film of high density polyethylene (HDPE) that has coextruded skin resins, laminated film or coatings on both sides of the HDPE base layer. Up to three layers of skin resins can be on each side of the base layer and can include heat sealable or non heat sealable type skins. For present purposes heat sealable skin resins are those which seal at a minimum seal temperature below the distortion temperature of HDPE base layer film.

More particularly, the present invention relates to an oriented polymer film which comprises:

a) a treated upper skin layer (a) capable of receiving a water base coating, layer (a) having a surface coextensively adherent to the upper surface of a base (or core) layer (b), and an exposed surface, the upper layer (a) comprising ethylene-propylene-butene-1 terpolymer, ethylene-propylene random copolymer, ethylene-propylene block copolymer, isotactic polypropylene, low density polyethylene (LDPE), very low density polyethylene (VLDPE), linear low density polyethylene (LLDPE), medium density polyethylene (MDPE), polyvinylidene chloride (PVDC), polyvinyl alcohol (PVOH) or an acrylic homo- or copolymer, the upper skin layer (a) being optionally compounded with an effective amount of anti-blocking agent, but being substantially devoid of silicone oil, the exposed surface of upper layer (a) containing a coefficient of friction-reducing amount of silicone oil transferred to the surface of (a) by contact with a silicone oil present upon the exposed surface of a lower skin layer (c);

b) a base layer (b) derived from HDPE, having an upper surface and lower surface,

c) lower skin layer (c) having a surface adherent to the lower surface of base layer (b), and an exposed surface, the layer (c) comprising ethylene-propylene-butene-1 terpolymer, ethylene-propylene random copolymer, ethylene-propylene block copolymer, isotactic polypropylene, low density polyethylene (LDPE), very low density polyethylene (VLDPE), linear low density polyethylene (LLDPE), medium density polyethylene (MDPE), the lower skin layer (c) being compounded with i) an
effective amount of anti-blocking agent, optionally comprising a particulate cross-linked hydrocarbyl-substituted polysiloxane, and ii) a quantity of silicone oil such that a coefficient of friction-reducing amount is present on the exposed surface of layer (c) as well as the exposed surface of layer (a) after mutual contact of said surfaces.

In another aspect, the present invention relates to a process for producing a film, which process comprises:

(1) providing the upper major surface of the above-described core layer (b) derived from a high density polyethylene with a coextensive water-based coating receiving upper skin layer (a) comprising ethylene-propylene-butene-1 terpolymer, ethylene-propylene random copolymer, ethylene-propylene block copolymer, isotactic polypropylene, low density polyethylene (LDPE), very low density polyethylene (VLDPE), linear low density polyethylene (LLDPE), medium density polyethylene (MDPE), polyvinylidene chloride (PVDC), polyvinyl alcohol (PVOH) or an acrylic homo- or copolymer, the upper skin layer (a) being optionally compounded with an effective amount of anti-blocking agent, but being substantially devoid of silicone oil;

(2) providing the lower major surface of core layer (b) with a coextensive surface lower skin layer (c) comprising ethylene-propylene-butene-1 terpolymer, ethylene-propylene random copolymer, ethylene-propylene block copolymer, isotactic polypropylene, low density polyethylene (LDPE), very low density polyethylene (VLDPE), linear low density polyethylene (LLDPE), medium density polyethylene (MDPE), said polymer being compounded with i) an effective amount of anti-blocking agent, optionally comprising a particulate cross-linked hydrocarbyl-substituted polysiloxane, and ii) a quantity of silicone oil such that a coefficient of friction-reducing amount will be present on the exposed surface of layer (c) as well as the exposed surface of layer (a) after mutual contact of said surfaces; and

(3) contacting the exposed surface of layer (c) with the exposed surface of layer (a) such that a coefficient of friction
reducing amount of silicone oil is transferred from the exposed surface of layer (c) to the exposed surface of layer (a).

The film composition of the present invention can comprise a biaxially oriented high density base layer of at least 50 weight percent of a high density polyethylene having a density of 0.96 or higher. The skin layers are used with HDPE films which have been biaxially oriented to provide product having good WVTR properties as well as other characteristics, e.g., barrier properties for gases such as oxygen, which are desirable in packaging films.

Those embodiments wherein a skin layer consists essentially of PVDC, PVOH or acrylic exhibit reduced water vapor transmission rate and oxygen transmission rate.

The skin layers help provide a film which is readily castable while maintaining the desirable properties of HDPE including dead fold, twist retention, TD tear and moisture barrier. The films can be cast, and oriented in the solid state up to about two times in the machine direction and six times or more in the transverse direction to give films having good dead-fold and water vapor transmission rate characteristics making them highly suited for packaging, particularly for dry foods.

The films with or without heat seal compositions in the skins can be used on a wide variety of packaging machines such as vertical form, fill and seal (VFFS) and high speed horizontal slug wrapping machines and can also be used for twist wrapping machines and laminations.

The films of the present invention are especially suited for applications requiring good dead fold characteristics. The present films can exhibit crease retention (as measured by ASTM D 920-49) in either or both directions of greater than 70%, greater than 85%, or even greater than 95% after 30 seconds, e.g., 70% to 85% TD and 75% to 95% MD.

The present films can possess both good lubricity and good receptivity for water-based surface coating materials and optical clarity allows the convertor/end user to utilize inks and/or adhesives which do not incorporate organic solvents. Organic solvents generally present health, explosion and fire
hazards and resort must be had to expensive ventilation and
safety equipment to provide an acceptable working environment
where they are used. Even with the use of such equipment, in
some jurisdictions, the atmospheric emission of organic solvents
is prohibited and expensive solvent recovery systems are
required to comply with legal limitations. The films herein
avoid organic solvent-containing coating materials thereby
eliminating the health, safety and environmental problems
attendant their use as well as the cost of the equipment
required to deal with the hazards which they pose.

According to the present invention, the upper skin layer
is derived from at least one polymer selected from the group
consisting of ethylene-propylene-butene-1 terpolymer, ethylene-
propylene random copolymer, ethylene-propylene block copolymer,
isotactic polypropylene, low density polyethylene (LDPE), very
low density polyethylene (VLDPE), linear low density
polyethylene (LLDPE), medium density polyethylene (MDPE),
polyvinylidene chloride (PVDC), polyvinyl alcohol (PVOH) and
acrylic. Any polymer blends being of polymers which are
compatible with one another. It should be noted that PVDC, PVOH
and acrylic are incompatible with the other named polymers so
that the former are used singly.

In one embodiment the upper skin layer (a) is derived from
a random ethylene-propylene copolymer. This may be derived from
ethylene and one or more co-monomers. The propylene content of
these random copolymers is typically from 70 to 85 weight
percent, more usually from 75 to 85 percent, with the balance
of ethylene and any other comonomers such as butylene. Suitable
copolymers of this type are random copolymers of ethylene and
propylene or random terpolymers of ethylene, propylene and
butylene. Preferred copolymers of this type include the
following:

Ethylene-propylene copolymers containing 2 - 10 weight
percent random ethylene, e.g. 3 - 7 weight percent
ethylene.

Ethylene-propylene-butylene random terpolymers containing
1 - 5 weight percent random ethylene, 10 - 25 weight
percent random butylene. The amounts of the random ethylene and butylene components in these copolymers are typically in the range of 10 to 25 percent total (ethylene plus butylene). Typical terpolymers of this type include those with about 1 - 5 percent ethylene and 10 - 25 percent butylene.

These copolymers typically have a melt flow rate from 5 to 10 with a density of about 0.9 and a melting point from 115°C to 130°C.

In another embodiment the upper skin layer (a) is derived from a low density polyethylene. This may be a linear low density polyethylene (LLDPE) or a non-linear polyethylene. These polymers typically have a melt index of 1 to 10. The low density polyethylenes should have a density of 0.88 to 0.93 while the linear materials may have a density as high as 0.94, usually in the range 0.90 - 0.94, e.g. 0.918 or 0.921, with a melt index from 1 to 10. The linear low density polyethylenes may be derived from ethylene together with other higher comonomers such as hexene-1 or octene-1.

Prior to incorporation in the film, e.g., before extrusion, upper layer (a) can be compounded with an anti-blocking effective amount of an anti-blocking agent, e.g., silica, clays, talc and glass which are preferably provided in the form of approximately spheroidal particles. The major proportion of these particles, for example, from more than half to as high as 90 weight percent or more, will be of such a size that significant portion of their surface area, for example, from 10 to 70 percent thereof, will extend beyond the exposed surface of layer (a). In a preferred embodiment, the anti-blocking agent comprises particulate cross-linked hydrocarbyl-substituted polysiloxane anti-blocking agent. Particularly preferred particulate cross-linked hydrocarbyl-substituted polysiloxanes include the polydimethylsiloxanes. Most particularly preferred are non-meltably polysiloxanes characterized as having a mean particle size of 0.5 to 20.0 microns and a three dimensional structure of siloxane linkages. Such materials are
available from Toshiba Silicone Co., Ltd., worldwide, and in the United States from General Electric Co., and are marketed under the tradename Tospearl. Other commercial sources of similar suitable materials are also known to exist. Such materials are further described as non-meltable crosslinked organosiloxane resin powders in U.S. Patent 4,769,418. Effective amounts of the particulate cross-linked hydrocarbyl-substituted polysiloxane anti-blocking agent can range from 100 to 5000 ppm, preferably 1000 to 3000 ppm, say, from 2500 to 3000 ppm, based on loading of the resin from which the upper layer (a) is prepared.

Although the polymer from which layer (a) is formed is not compounded with a silicone oil, this layer will ultimately acquire a coefficient of friction-reducing amount of silicone oil. Thus, when the finished film laminate containing silicone oil on the exposed surface of layer (c) is taken up on a winding coil, some of the silicone oil will be transferred from this surface to the exposed surface of layer (a), primarily to the exposed surfaces of the anti-blocking agent particles which protrude from layer (a). However, since the interior of layer (a) contains no amount of silicone oil which could interfere with the heat sealing or other properties of this layer (and ordinarily contains no silicone oil at all), the presence thereon of the transferred silicone oil serves to reduce the coefficient of friction of the layer without significantly impairing its receptivity to water-based coatings, its heat sealability or its optical clarity.

Optionally, layers (a) and/or (c) can contain a minor amount of a wax, e.g., a microcrystalline wax for added lubricity and/or reduced water vapor transmission rate. Amounts of wax from 2 to 15 weight percent of either or both layers (a) and (c) can be used if desired. Either or both of these layers can also contain pigments, fillers, stabilizers, light protective agents, antistatic agents, or other suitable modifying ingredients if desired.

While a variety of antistatic agents are available commercially which are suitable for use herein, the group of
ethoxylated amines and ethoxylated amides is preferred for use in the practice of the present invention. Ethoxylated amines are available from the Humco Chemical Division of Whitco Chemical Corp. under the trademark of Kemamine, from the Noury Chemical Company under the trademark of Armostat and from other sources. Ethoxylated amides are available from Akzo Chemie America under the trademark of Ethmid, from the Oxyxnol Chemical Company under the tradename of Oxynol and from other sources. Particularly preferred for their migratory properties is the group of ethoxylated amines.

The range of antistatic agent levels useful in the practice of the present invention is from 500 ppm to 2000 ppm of layers (a) and/or (c), with 1000 ppm to 1500 ppm being particularly preferred.

In order to enhance its receptivity for water-based coatings and inks, layer (a) can be treated by such known and conventional techniques as corona discharge and flame treating.

Lower layer (c) of the film laminate can be of substantially the same composition as that of upper surface layer (a) except that i) the polymer is not derived from components selected from the group consisting of PVDC, PVCH and acrylic, ii) the polymer constituting layer (c) is compounded with a silicone oil and coefficient of friction reducing amounts of an anti-blocking agent which preferably comprises particulate cross-linked hydrocarbyl-substituted polysiloxane as discussed above. The silicone oil advantageously possesses a viscosity of from 350 to 100,000 centistokes with 10,000 to 30,000 centistokes being especially preferred. Examples of suitable silicone oils are polydialkylsiloxanes, polyalkylphenylsiloxanes, olefin-modified siloxane oils, olefin/-polyether-modified silicone oils, epoxy modified silicone oils and alcohol-modified silicone oils, polydialkylsiloxanes which preferably have from about 1 to about 4 carbon atoms in the alkyl group, in particular polydimethylsiloxanes. Of the foregoing, the polydialkylsiloxanes, in particular a polydimethylsiloxane, are preferred for use herein.
The silicone oil can be added to lower skin layer (c) generally in the form of a dispersion or emulsion, the silicone being present within, as well as on the exposed surface of this layer as discrete microglobules, frequently of an average size from 1 to 2 microns. The silicone oil, which is generally substantially uniformly distributed on the exposed surface of layer (c), is responsible for imparting a reduced coefficient of friction to this surface as well as to the exposed surface of layer (a) when some of the oil is transferred thereto after these surfaces have been placed in mutual contact, e.g., as will occur when the laminate film has been wound on a winding coil.

Polydimethylsiloxane or other silicone oil can be present at from 0.15 to 1.5 weight percent of lower layer (c). Some of this silicone oil will, of course, be present on the exposed surface of layer (c). The amount selected should in any event be sufficient to provide a coefficient of friction of layers (a) and (c) (following transfer of silicone oil microglobules to the latter) of about 0.4 or less, preferably between 0.25 to 0.3 up to at least about 60°C. Because of the manner in which the silicone oil is applied to just the exposed surface of upper layer (a), such layer exhibits an improved coefficient of friction but not at the expense of its receptivity to water-based coatings, its heat sealability or its optical clarity.

The silicone oil should be incorporated as homogeneously as possible in the polymer constituting layer (c). This can be achieved by either incorporating the silicone oil as a dispersion or emulsion at room temperature and then heating the blend with the application of shearing forces or by incorporating the oil while the blend is being melted. The mixing temperature must be high enough to soften the blend and enable a very uniform mixture to be formed. The temperature required in a kneader or extruder is generally from 170 to 270°C.

In addition to silicone oil, layer (c) is compounded with an effective anti-blocking and coefficient of friction reducing amount of an anti-blocking agent which comprises particulate cross-linked hydrocarbyl-substituted polysiloxane, as described
above. Effective amounts can range from 500 to 5000 ppm, preferably 1000 to 3000 ppm, say, from 2500 to 3000 ppm, based on loading of the resin from which the layer is prepared.

Core layer (b) is derived from imbalanced biaxially oriented film base layer made from a major proportion of a high density polyethylene (HDPE) having a density of at least 0.96. The film can be composed exclusively of a single HDPE resin, a mixture of HDPE resins, or of HDPE containing a minor proportion of other resource polymers. These high density polyethylenes typically have a melt index from 0.5 to 10, usually from 0.7 to 2. The mixture of HDPE resins gives better processing characteristics in the extruder by reducing extruder torque. Films made with a blend of HDPE resins reduce splittiness of the film which manifests itself as the tendency of the film to break in the TD direction during operation on vertical, form, fill and seal (VFFS) machinery.

The blends of HDPE polymers can comprise two or more polymers all of which preferably have densities of 0.96 or greater. Blends of HDPE polymers advantageously comprise a major proportion of HDPE having a melt index of 0.5 to 6 and one or more polymers having a different melt index.

Terblends have been found particularly desirable. Suitable terblends generally comprise 50 to 98 weight percent, preferably 84 to 96 weight percent of HDPE having a density of 0.96 or higher and a melt index of greater than 0.5 to about 1.0; 1 to 25 weight percent, preferably 3 to 8 weight percent of HDPE having a density of 0.96 or greater and a melt index of 0.1 to 0.5; and 1 to 25 weight percent, preferably 3 to 8 weight percent, of HDPE having a density of 0.96 or higher and a melt index of greater than 2 to about 8. Preferably, the second and third HDPE polymers which are minor components are present in about equal amounts.

In a preferred embodiment, the film of the present invention has both its upper skin layer and lower skin layer comprising a mixture of ethylene-propylene-butene-1 terpolymer and low density polyethylene (LDPE). For example, the upper skin layer and lower skin layer can comprise 85 to 95 wt%, say
90 wt%, of ethylene-propylene-butene-1 terpolymer and 5 to 15 wt%, say 10 wt% low density polyethylene (LDPE).

The proper degree of orientation in the film contributes to the desired physical properties, as well as good WVTR and dead-fold characteristics. For example, it has been determined that films with a thickness of 1.4 to 4 mils will have acceptable WVTR (g-mil/100 in2/24 hr - 1 atm) of less than about 0.2/mil whereas a somewhat heavier gauge (1.5 times thicker or more) is needed in a blown HDPE film to achieve the same WVTR.

The benefits of reduced WVTR are due to the improvements obtained by biaxial orientation below the HDPE melting point. Although higher density HDPE resin having a density of 0.957 or greater can be made directly into thin films by cast extrusion, problems of curling, uniformity, flatness and high WVTR remain as obstacles. Accordingly, thin HDPE films of 0.8 to 1.5 mils, such as 0.5 to 1.2 mils, having the best balance of properties, particularly for VFFS applications, are obtained with imbalanced biaxially oriented films prepared from films having a cast gauge of 12 to 20 mils reduced to the desired gauge by orientation.

The final film gauge will typically be not more than about 2.5 mils.

The HDPE base film is oriented either before or after the skins are applied to a degree of 1:1:1 to 2:1, usually from 1.25:1 to 2:1, in the machine direction (MD), and to a degree of 6:1 to 12:1 in the transverse direction (TD). The films exhibit improved dead-fold, and other physical properties which are markedly better than cast and blown HDPE films, even when the total film thickness is reduced to less than 1 or 2 mils. When provided with a skin layer as described below, the films are particularly suited for use in packaging, especially of dry foodstuffs. The films may be used in a wide variety of packaging equipment including vertical form, fill and seal (VFFS), high speed horizontal slug wrapping, as well as twist folding packaging machines.

The skin layers can be applied to the HDPE base film in any conventional manner, for example, by coating or coextrusion before orientation or by coating the HDPE after one or both of
the biaxial orientation operations. Generally, the skin layers of the film of the invention comprise less than 50 wt%, preferably less than 15 wt%, say about 10 wt% of the total film.

The films may be produced and oriented in the conventional manner. When the skin layer is present on one or both sides of the HDPE film, cast extrusion is generally accomplished using a standard cast roll and water bath system.

In the usual manner the film is heated to its orientation temperature and first subjected to MD orientation between two sets of nip rolls, the second rotating at a greater speed than the first in an amount equal to the desired draw ratio. Then the film is TD oriented by heating and subjecting it to transverse stretching in a tenter frame. Typically MD orientation is conducted at 60° to 120°C and TD orientation at 110° to 160°C.

The skin layers and/or base layer may contain stabilizers to minimize oxidative and thermal degradation, as well as other additives to achieve other functionality including, but not limited to, static reduction, ease of processing, ink receptivity, etc.

The following Examples, in which all parts are by weight unless otherwise specified, illustrate the invention.

**Example 1**

A three-layer biaxially oriented film having a 1.15 mil final thickness was prepared by coextruding:

a base layer HDPE (M-6211 obtained from Oxychem of Houston, Texas) comprising 90% of the total film thickness;

a first skin layer on one side of the base layer comprising 5% of the total film thickness which is Chisso 7510, obtained from Chisso Corp. of Japan, consisting of ethylene-propylene-butene-1 terpolymer and 2300 ppm Syloloc 44™, an antiblock agent available from W. R. Grace (USA), the first skin layer being subjected to treatment by corona discharge; and

a second skin layer on the other side of the base layer (Chisso 7820, obtained from Chisso Corporation of Japan), comprising 5% of the total film thickness, whose resin component
is a blend of about 90% ethylene-propylene-butene-1 terpolymer and 10% LDPE with 2300 ppm Sylobloc 42\textsuperscript{TM}, an antiblock agent available from W. R. Grace (USA), which second skin layer further contains about 1 wt% of silicone oil.

The film was prepared in a standard cast roll system and then oriented 1.3 times in the MD at about 115°C, and 9.0 times in the TD direction at 110°-160°C in a tenter frame.

The film was tested for dead fold properties by ASTM D-920-49 (\% crease retention after 30 seconds). In the transverse direction, the film of the present invention (HDPE with terpolymer skins) exhibited about 75\% crease retention, and about 85\% crease retention in the machine direction. This compares favorably with 0.80 mil polypropylene film with terpolymer skins (80 BS\textsuperscript{TM}G-ONE available from Mobil Chemical Co. of Pittsford, NY (8\% TD, 6\% MD)); 1.25 mil 100\% polypropylene film (240 B available from Mobil Chemical Co. of Pittsford, NY (15\% TD, 13\% MD)); 1.85 mil HDPE film with ionomer skin (Blown HD, available from American National Can Co. of Neenah, WI (47\% TD, 36\% MD)); and 2.0 mil wax coated glassine (55\%).

Example 2

A three-layer biaxially oriented film having a 1.15 mil final thickness was prepared in accordance with Example 1 except that 3000 ppm of Tospearl SF-344 available from GE, with a particle size of 4.5 microns, was compounded with the resin used for both skin layers prior to coextruding.

Example 3

A three-layer biaxially oriented film having a 1.15 mil final thickness was prepared in accordance with Example 1 except that 1000 ppm of Tospearl SF-344 available from GE, with a particle size of 4.5 microns, was compounded with the resin used for the second skin layer prior to coextruding.

Example 4

A three-layer biaxially oriented film having a 1.15 mil final thickness was prepared in accordance with Example 1 except that
2000 ppm of Tospearl SF-344 available from GE, with a particle size of 4.5 microns, was compounded with the resin used for the second skin layer prior to coextruding.

**Example 5**

A three-layer biaxially oriented film having a 1.15 mil final thickness was prepared in accordance with Example 1 except that 3000 ppm of Tospearl SF-344 available from GE, with a particle size of 4.5 microns, was compounded with the resin used for the second skin layer prior to coextruding.

**Example 6**

The films of Examples 1 to 5 were tested for kinetic coefficient of friction values for both their treated and untreated skins according to the procedure of ASTM D 1894-78. The results are set out below in the Table.

<table>
<thead>
<tr>
<th>TABLE</th>
<th>Untreated Skin COF</th>
<th>Treated Skin COF</th>
</tr>
</thead>
<tbody>
<tr>
<td>20 EX.1 (0 ppm Tospearl)</td>
<td>0.47</td>
<td>0.33</td>
</tr>
<tr>
<td>EX.2 (3000ppm Tospearl)</td>
<td>0.23</td>
<td>0.21</td>
</tr>
<tr>
<td>(both layers)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>EX.3 (1000 ppm Tospearl)</td>
<td>0.34</td>
<td>0.35</td>
</tr>
<tr>
<td>EX.4 (2000ppm Tospearl)</td>
<td>0.34</td>
<td>0.35</td>
</tr>
<tr>
<td>25 EX.5 (3000ppm Tospearl)</td>
<td>0.23</td>
<td>0.35</td>
</tr>
</tbody>
</table>

The addition of particulate cross-linked hydrocarbyl-substituted polysiloxane as an antiblocking/slip agent in the resin of both layers (Ex. 2) results in a significant decrease in coefficient of friction values both for untreated surface to untreated surface and treated surface to treated surface. The addition of particulate cross-linked hydrocarbyl-substituted polysiloxane as an antiblocking/slip agent in the resin of the lower untreated skin layer (Ex. 3 to 5) results in a significant decrease in coefficient of friction values for untreated surface to untreated surface. This provides enhanced machinability of the film for both horizontal and vertical packaging machines.
CLAIMS

1. An oriented polymer film which comprises:
   a) a treated upper skin layer (a) capable of receiving a water base coating, layer (a) having a surface coextensively adherent to the upper surface of a base layer (b), and an exposed surface, the upper layer (a) comprising ethylene-propylene-butene-1 terpolymer, ethylene-propylene random copolymer, ethylene-propylene block copolymer, isotactic polypropylene, low density polyethylene (LDPE), very low density polyethylene (VLDPE), linear low density polyethylene (LLDPE), medium density polyethylene (MDPE), polyvinylidene chloride (PVDC), polyvinyl alcohol (PVOH) or an acrylic, homo- or copolymer, the upper skin layer (a) being optionally compounded with an effective amount of anti-blocking agent, but being substantially devoid of silicone oil, the exposed surface of upper layer (a) optionally containing a coefficient of friction-reducing amount of silicone oil transferred to the surface of (a) by contact with a silicone oil present upon the exposed surface of a lower skin layer (c);
   b) a base layer (b) derived from HDPE, having an upper surface and lower surface,
   c) a lower skin layer (c) having a surface adherent to the lower surface of base layer (b), and an exposed surface, the layer (c) comprising ethylene-propylene-butene-1 terpolymer, ethylene-propylene random copolymer, ethylene-propylene block copolymer, isotactic polypropylene, low density polyethylene (LDPE), very low density polyethylene (VLDPE), linear low density polyethylene (LLDPE), medium density polyethylene (MDPE), the lower skin layer (c) being compounded with i) an effective amount of anti-blocking agent, optionally comprising a particulate cross-linked hydrocarbyl substituted polysiloxane, and ii) a quantity of silicone oil such that a coefficient of friction-reducing amount is present on the exposed surface of layer (c) as well as the exposed surface of layer (a) after mutual contact of said surfaces.

2. A film according to claim 1 wherein the base layer (b)
comprises a biaxially oriented base layer of at least 50 weight percent of a high density polyethylene having a density of 0.96 or higher.

5 3. A film according to claim 1 or 2 which is oriented to a degree of 1:1 to 2:1 in the machine direction.

4. A film according to claim 2 or 3 which is oriented to a degree of 6:1 to 12:1 in the transverse direction.

5. A film according to any preceding claim wherein at least one skin layer comprises ethylene-propylene-butene-1 terpolymer, or a mixture of ethylene-propylene-butene-1 terpolymer and low density polyethylene (LDPE).

6. A film according to any preceding claim wherein the upper skin layer (a) comprises PVDC, PVOH or acrylic homo- or copolymer.

7. A film according to any preceding claim wherein upper skin layer (a) is compounded with an anti-blocking agent.

8. A film according to claim 7 wherein the anti-blocking agent comprises a particulate cross-linked hydrocarbyl substituted polysiloxane.

9. A film according to any preceding claim wherein lower skin layer (c) is compounded with, as anti-blocking agent, a particulate cross-linked hydrocarbyl-substituted polysiloxane.

10. A film according to any preceding claim wherein the anti-blocking agent is present in upper skin layer (a) and/or lower skin layer (c) in an amount from 1000 to 3000 ppm of the layer.

11. A film according to any preceding claim wherein the exposed surface of the upper skin layer (a) is treated by flame
12. A film according to any preceding claim which has
treated retention of at least 70% after 30 seconds.

13. A film according to any preceding claim wherein the
lower skin layer (c) is compounded with an anti-static agent.

14. A process for producing a film, which process
comprises:

(1) providing the upper major surface of a core layer (b)
derived from a high density polyethylene with a coextensive
water-based coating receiving upper skin layer (a) comprising
ethylene-propylene-butene-1 terpolymer, ethylene-propylene
random copolymer, ethylene-propylene block copolymer, isotactic
polypropylene, low density polyethylene (LDPE), very low density
polyethylene (VLDPE), linear low density polyethylene (LLDPE),
medium density polyethylene (MDPE), poly(vinylidene chloride
(PVDC), polyvinyl alcohol (PVOH) and acrylic, said upper skin
layer (a) being optionally compounded with an effective amount
of anti-blocking agent, but being substantially devoid of
silicone oil;

(2) providing the lower major surface of core layer (b)
with a coextensive surface layer (c) comprising ethylene-
propylene-butene-1 terpolymer, ethylene-propylene random
copolymer, ethylene-propylene block copolymer, isotactic
polypropylene, low density polyethylene (LDPE), very low density
polyethylene (VLDPE), linear low density polyethylene (LLDPE),
medium density polyethylene (MDPE), said polymer being
compounded with i) an effective amount of anti-blocking agent,
optionally comprising a particulate cross-linked hydrocarbyl-
substituted polysiloxane, and ii) a quantity of silicone oil
such that a coefficient of friction-reducing amount will be
present on the exposed surface of layer (c) as well as the
exposed surface of layer (a) after mutual contact of said
surfaces; and

(3) contacting the exposed surface of layer (c) with the
exposed surface of layer (a) such that a coefficient of friction reducing amount of silicone oil is transferred from the exposed surface of layer (c) to the exposed surface of layer (a).
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

IPC(6) : B32B 1/00, 27/00, 70/00, 5/00, 31/00; B29C 55/00

US CL : Please See Extra Sheet.

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)

U.S. : 428/35.7, 349, 500, 516, 523, 323, 220, 447, 515, 518, 910; 156/229

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 practicable, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category*</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y</td>
<td>US, A, 4,692,379 (KEUNG ET AL) 08 September 1987, abstract; column 2, line 43 to column 3, line 39; column 4, lines 9-34.</td>
<td>1-3, 14</td>
</tr>
<tr>
<td>Y</td>
<td>US, A, 4,870,122 (LU) 26 September 1989, abstract; column 2, lines 17-31.</td>
<td>1-3</td>
</tr>
<tr>
<td>A</td>
<td>US, A, 4,966,933 (KAWAKAMI ET AL) 30 October 1990, see entire document.</td>
<td>1-3</td>
</tr>
<tr>
<td>A</td>
<td>US, A, 4,652,618 (SUMIDA ET AL) 24 March 1987, see entire document.</td>
<td>1-3</td>
</tr>
</tbody>
</table>

☐ Further documents are listed in the continuation of Box C. ☐ See patent family annex.

* Special categories of cited documents:

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"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 invention

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"*" document member of the same patent family

Date of the actual completion of the international search: 07 FEBRUARY 1995

Date of mailing of the international search report: 07 MAR 1995

Name and mailing address of the ISA/US

Commissioner of Patents and Trademarks

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A. CLASSIFICATION OF SUBJECT MATTER:
US CL:
428/35.7, 349, 500, 516, 523, 323, 220, 447, 515, 518, 910; 156/229