(54) Title: MULTILAYER PRESSURE SENSITIVE ADHESIVE

Fig. 1

(57) Abstract: Provided is a pressure sensitive adhesive article comprising (a) a substrate (Sa), (b) in contact with said substrate (Sa), a layer (Lb) of a tackifier-free composition (Cb) that comprises one or more acrylic polymer (POLb) having Tg of -10°C or lower, and (c) in contact with said layer (Lb), a layer (Lc) of a tackifier-containing composition (Cc) that comprises one or more tackifier and one or more hydrocarbon polymer (POLc) having Tg of 10°C. Also provided is a bonded article made by a process comprising contacting a substrate (Sd) with such a pressure sensitive adhesive article, wherein said substrate (Sd) is in contact with said layer (Lc).
MULTILAYER PRESSURE SENSITIVE ADHESIVE

[0001] Pressure sensitive adhesives (PSAs) that contain acrylic polymers have many desirable characteristics. For example, they generally have better resistance to chemical reagents and UV light than PSAs made from many other materials. It is often desired to provide a PSA that bonds well to polyolefin substrates. It is also desired to provide PSAs that use less tackifier than had been used in previously used PSAs. There is a need for PSAs that contain acrylic polymer, that show good adhesion to polyolefin substrates, and that have a relatively low amount of tackifier.

[0002] US 6,183,862 teaches a multilayer PSA that contains a facestock, a first adhesive layer, and a PSA layer. In the PSA taught by US 6,183,862, both the first adhesive layer and the PSA layer contain tackifier.

[0003] The following is a statement of the invention.

[0004] A first aspect of the present invention is a pressure sensitive adhesive article comprising

(a) a substrate (Sa),
(b) in contact with said substrate (Sa), a layer (Lb) of a tackifier-free composition (Cb) that comprises one or more acrylic polymer (POLb) having Tg of -10°C or lower, and
(c) in contact with said layer (Lb), a layer (Lc) of a tackifier-containing composition (Cc) that comprises one or more tackifier and one or more hydrocarbon polymer (POLc) having Tg of -10°C.

[0005] A second aspect of the present invention is a bonded article made by a process comprising contacting a substrate (Sd) with the article of the first aspect, wherein said substrate (Sd) is in contact with said layer (Lc).

[0006] Fig. 1 is a vertical cross section of a pressure sensitive adhesive article of the present invention showing substrate (Sa) (1); layer (Lb) (2) of a tackifier-free composition (Cb), which contains one or more acrylic polymer (POLb) having Tg of -10°C or lower; and layer (Lc) (3) of a tackifier-containing composition (Cc), which contains one or more tackifier and one or more hydrocarbon polymer (POLc) having Tg of -10°C or lower. Fig. 1 is not drawn to scale in any sense. For example, the size of the pressure sensitive adhesive article of the present invention in the horizontal direction shown in Fig. 1 may be larger by a factor of 1,000 or more than the size in the vertical direction shown in Fig. 1.
Fig. 2 (also not drawn to scale) depicts a preferred use to which the pressure sensitive adhesive article of the present invention may be put. Fig. 2 shows layer (Lc) (3) in contact with an additional substrate (Sd) (4).

The following is a detailed description of the invention.

As used herein, the following terms have the designated definitions, unless the context clearly indicates otherwise.

When a ratio is said herein to be X:1 or greater, it is meant that the ratio is Y:1, where Y is greater than or equal to X. For example, if a ratio is said to be 3:1 or greater, that ratio may be 3:1 or 5:1 or 100:1 but may not be 2:1. Similarly, when ratio is said herein to be W:1 or less, it is meant that the ratio is Z:1, where Z is less than or equal to W. For example, if a ratio is said to be 15:1 or less, that ratio may be 15:1 or 10:1 or 0:1:1 but may not be 20:1.

As used herein, Dynamic Mechanical Analysis (DMA) refers to measurements made in shear geometry in the linear viscoelastic range at frequency of 1 sec⁻¹. DMA measures the elastic modulus (G'), the loss modulus (G''), and tandelta (the quotient found by dividing G'' by G'), synonymously called "tan(δ)"). A curve showing tandelta as a function of temperature is known herein as a "tandelta curve." Elastic modulus is reported herein in units of kilopascals (kPa).

The glass transition temperature (Tg) of a material is determined by differential scanning calorimetry using the midpoint method and temperature scan rate of 10°C per minute according to test method ASTM D7426-08 (American Society of Testing and Materials, Conshohocken, PA, USA).

A "polymer," as used herein is a relatively large molecule made up of the reaction products of smaller chemical repeat units. Polymers may have structures that are linear, branched, star shaped, looped, hyperbranched, crosslinked, or a combination thereof; polymers may have a single type of repeat unit ("homopolymers") or they may have more than one type of repeat unit ("copolymers"). Copolymers may have the various types of repeat units arranged randomly, in sequence, in blocks, in other arrangements, or in any mixture or combination thereof.

Polymers molecular weights can be measured by standard methods such as, for example, size exclusion chromatography (SEC, also called gel permeation chromatography or GPC). Polymers have weight-average molecular weight (Mw) of 1000 or more. Polymers may have extremely high Mw; some polymers have Mw above 1,000,000; typical polymers have Mw of 1,000,000 or less. Some polymers are crosslinked, and crosslinked polymers are considered to have infinite Mn.
A s used herein "weight of polymer" means the dry weight of polymer.

Molecules that can react with each other to form the repeat units of a polymer are known herein as "monomers." The repeat units so formed are known herein as "polymerized units" of the monomer.

Vinyl monomers have the structure

\[
\begin{array}{c}
\text{R}^1 \quad \text{R}^2 \\
\text{R}^3 \quad \text{C} = \text{C} \\
\text{R}^4
\end{array}
\]

where each of \( \text{R}^1, \text{R}^2, \text{R}^3, \) and \( \text{R}^4 \) is, independently, a hydrogen, a halogen, an aliphatic group (such as, for example, an alkyl group), a substituted aliphatic group, an aryl group, a substituted aryl group, another substituted or unsubstituted organic group, or any combination thereof.

Some suitable vinyl monomers include, for example, styrene, substituted styrenes, dienes, ethylene, other alkenes, dienes, ethylene derivatives, and mixtures thereof. Ethylene derivatives include, for example, unsubstituted or substituted versions of the following: ethenyl esters of substituted or unsubstituted alkanoic acids (including, for example, vinyl acetate and vinyl neodecanoate), acrylonitrile, (meth)acrylic acids, (meth)acrylates, (meth)acrylamides, vinyl chloride, halogenated alkenes, and mixtures thereof. As used herein, "(meth)acrylic" means acrylic or methacrylic; "(meth)acrylate" means acrylate or methacrylate; and "(meth)acrylamide" means acrylamide or methacrylamide. "Substituted" means having at least one attached chemical group such as, for example, alkyl group, alkenyl group, vinyl group, hydroxyl group, carboxylic acid group, other functional groups, and combinations thereof. In some embodiments, substituted monomers include, for example, monomers with more than one carbon-carbon double bond, monomers with hydroxyl groups, monomers with other functional groups, and monomers with combinations of functional groups. (Meth)acrylates are substituted and unsubstituted esters or amides of (meth)acrylic acid.

As used herein, acrylic monomers are monomers selected from (meth)acrylic acid, alkyl esters of (meth)acrylic acid, alkyl esters of (meth)acrylic acid having one or more substituent on the alkyl group, (meth)acrylamide, N-substituted (meth)acrylamide, and mixtures thereof. As used herein, vinylaromatic monomers are monomers selected from styrene, alpha-alkyl styrenes, and mixtures thereof.

As used herein, an "acrylic " polymer is a polymer in which 30% or more of the polymerized units are selected from acrylic monomers and also in which 75% or more of the polymerized units are selected from the group consisting of acrylic monomers and vinylaromatic monomers. The percentages are by weight based on the weight of the polymer.
As used herein, an olefin polymer is a polymer in which 70% or more of the polymerized monomer units are selected from hydrocarbon alkenes, hydrocarbon dienes, and mixtures thereof, by weight based on the weight of the polymer.

As used herein, a hydrocarbon polymer is a polymer in which all of the atoms are either hydrogen or carbon.

A tackifier is an organic compound having molecular weight of 500 to 10,000 and having glass transition temperature of 0°C or higher.

The compatibility of a tackifier with a polymer is assessed according to the following test. First, the polymer is tested by DMA over the temperature range of -50°C to 110°C; the tandelta curve is recorded, and the number of peaks in that curve is noted. Also, a blend is made of the tackifier and the polymer having ratio of T% tackifier and P% polymer by weight (where T% plus P% is 100%), and the blend is subjected to DMA over the same temperature range, and the tandelta curve of the blend is observed. If the polymer and the tackifier are compatible at that ratio, the blend will appear visually to be homogeneous, and the number of peaks in the tandelta curve of the blend is the same as the number of peaks in the tandelta curve of the polymer alone, though one or more of the peaks from the polymer-alone tandelta curve may be shifted to a different temperature in the tandelta curve of the blend. If the polymer and the tackifier are incompatible at that ratio, either the blend will appear inhomogeneous, with apparent phase separation, or the tandelta curve of the blend will show at least one more peak than the number of peaks shown in the tandelta curve of the polymer alone, or both. The lowest value of T% for which the blend is incompatible is known herein as the "compatibility limit" of that tackifier in that polymer. When the compatibility limit is 20% or higher, the tackifier is said herein to be compatible with the polymer.

A Pressure Sensitive Adhesive (PSA) is an adhesive that forms a bond with a substrate when pressure is applied to bring the adhesive and the substrate into contact. The bond forms without addition of further materials or the application of heat. As used herein, a pressure sensitive adhesive article is an article in which a pressure sensitive adhesive is adhered to a first substrate and in which a surface of the PSA (the "available surface") is available to make contact with a second substrate. The available surface of the PSA may or may not be in contact with a release material. A release material is a material that forms a weak bond with the PSA and may be easily removed so that the available surface is exposed.

A composition is considered herein to be tackifier-free if the amount of all tackifiers in the composition is either zero or is less than 10% by weight based on the dry
weight of the composition. A composition herein is considered to contain tackifier (or synonymously to be "tackifier-containing") if the amount of tackifier in the composition is 10% or more by weight based on the dry weight of the composition.

[0028] A composition is herein considered to be "aqueous" if the composition contains water in the amount of 25% or more by weight based on the weight of the composition.

[0029] The present invention involves the use of a substrate, herein labeled substrate (Sa). The substrate (Sa) may be any material. Preferred are paper, polymer film, and metal foil. Among polymer films, preferred are polyester, polyvinyl chloride, polyethylene, and polypropylene films. Among polymer films, preferred are those in which at least one side has been treated by corona discharge.

[0030] In contact with substrate (Sa) is a layer of a tackifier-free composition, herein referred to as composition (Cb). Composition (Cb) contains one or more polymer, herein referred to as polymer (POLb). Polymer (POLb) has Tg of -10°C or lower; preferably -20°C or lower; more preferably -30°C or lower. Preferably polymer (POLb) has Tg of -100°C or higher.

[0031] Preferably, the amount of tackifier in composition (Cb) is, by weight based on the dry weight of composition (Cb), less than 10%; more preferably 3% or less; more preferably 1% or less; more preferably zero.

[0032] Preferably, polymer (POLb) is an acrylic polymer. Preferably, the amount of polymerized units of acrylic monomers in polymer (POLb) is, by weight based on the weight of polymer (b), 50% or more; more preferably 70% or more; more preferably 90% or more; more preferably 99% or more. Preferably, polymer (POLb) has Mw of 10,000 or higher; more preferably 50,000 or higher.

[0033] Preferably, polymer (POLb) contains polymerized units of one or more of n-butyl acrylate (n-BA), ethyl acrylate (EA), isooctyl acrylate (i-OA), or a mixture thereof. As used herein, "isooctyl" is an unsubstituted alkyl group that contains exactly 8 carbon atoms in a branched configuration. The term "isooctyl" includes all branched isomers of 8-carbon alkyl groups and all mixtures of such isomers, including, for example, the 2-ethylhexyl group, dimethyl-hexyl groups, methyl-heptyl groups, trimethyl-pentyl, and mixtures thereof. Preferably, the sum of the amounts of polymerized units of n-BA, polymerized units of EA, and polymerized units of i-OA in polymer (POLb), by weight based on the weight of polymer (POLb), is 50% or more; more preferably 75% or more; more preferably 90% or more.

[0034] Preferably, every polymer in composition (Cb) that has Mw of 10,000 or higher is an acrylic polymer.
[0035] Preferably, the amount of all tackifiers in composition (Cb), by weight based on the dry weight of composition (Cb), is 5% or less; more preferably 1% or less; more preferably zero.

[0036] Preferably, the amount of polymer (POLb) in composition (Cb), by weight based on the dry weight of composition (Cb), is 80% or more; more preferably 90% or more; more preferably 95% or more.

[0037] Preferably, composition (Cb) forms a continuous layer (Lb) on a face of substrate (Sa). Preferably, the thickness of the layer of composition (Cb) is 7.5 micrometer or more. Preferably, the thickness of the layer of composition (Cb) is 75 micrometer or less; more preferably 55 micrometer or less; more preferably 40 micrometer or less.

[0038] Preferably, composition (Cb) has the properties of a PSA. Preferably, composition (Cb) has elastic modulus of 20 kPa or higher over a temperature range that includes the range of 10°C to 40°C. Preferably, composition (Cb) has elastic modulus over a temperature range that includes the range of 10°C to 40°C of 1,000 kPa or lower; more preferably 500 kPa or lower.

[0039] A layer of composition (Cc) is in contact with the layer (Lb) of composition (Cb). Composition (Cc) contains one or more polymer (POLc). Polymer (POLc) is a hydrocarbon polymer.

[0040] Preferably, polymer (POLc) is selected from block polymers, statistical copolymers, and homopolymers, or mixtures thereof. Among homopolymers and statistical copolymers, preferred are styrene/butadiene copolymers, natural rubber, olefin polymers, and mixtures thereof.

[0041] Among embodiments in which polymer (POLc) is a homopolymer or statistical copolymer, preferably polymer (POLc) has Tg of -10°C or lower; more preferably -20°C or lower; more preferably -30°C or lower. Preferably polymer (POLc) has Tg of -100°C or higher.

[0042] Among embodiments in which polymer (POLc) is a block polymer, polymer (POLc) may have two blocks, three blocks, or more than three blocks. Preferred are triblock polymers, which are block polymers having exactly three blocks. Preferred triblock polymers have a block of polymerized units of styrene at each end and a middle block in between. Preferred middle blocks contain polymerized units of butadiene, isoprene, ethylene, propylene, butylene, or a mixture thereof; more preferred middle blocs contain polymerized units of isoprene.
When polymer (POLc) is a block polymer, it is expected that polymer (POLc) will exhibit more than one Tg. Preferably, at least one Tg is -10°C or lower; more preferably -20°C or lower; more preferably -30°C or lower.

Composition (Cc) contains one or more tackifier. Preferred tackifiers are rosin resins, derivatives of rosin resins, hydrocarbon resins, hydrogenated hydrocarbon resins, terpene resins, and derivatives of terpene resins. Preferably, composition (Cc) contains at least one tackifier that is compatible with at least one polymer (POLc). More preferably, every tackifier in composition (Cc) is compatible with every polymer (POLc). Preferred tackifiers are hydrocarbon resins and hydrogenated hydrocarbon resins; more preferred are hydrocarbon resins.

Preferably, the amount of tackifier in composition (Cc) is 5% or more by weight based on the dry weight of composition (Cc); more preferably 10% or more.

When polymer (POLc) is a block polymer, the preferred amount of tackifier is, by weight based on the weight of composition (Cc), 40% to 65%. When polymer (POLc) is a styrene/butadiene copolymer, the preferred amount of tackifier is, by weight based on the weight of composition (Cc), 5% to 50%, more preferably 10% to 50%. When polymer (POLc) is natural rubber, the preferred amount of tackifier is, by weight based on the weight of composition (Cc), 40% to 65%. When polymer (POLc) is a polyolefin, the preferred amount of tackifier is, by weight based on the weight of composition (Cc), 40% to 70%.

Preferably, composition (Cc) has the properties of a PSA. Preferably, composition (Cc) has elastic modulus of 20 kPa or higher over a temperature range that includes the range of 10°C to 40°C. Preferably, composition (Cc) has elastic modulus over a temperature range that includes the range of 10°C to 40°C of 1,000 kPa or lower; more preferably 500 kPa or lower.

Preferably, the thickness of the layer of composition (Cc) is 1.25 micrometer or more. Preferably, the thickness of the layer of composition (Cc) is 25 micrometer or less; more preferably 10 micrometer or less; more preferably 5 micrometer or less.

Preferably, the sum of the amount of polymer (POLc) in composition (Cc) plus the amount of tackifier in composition (Cc) is, by weight based on the dry weight of composition (Cc), 80% or more; more preferably 90% or more; more preferably 95% or more.
Preferably, the amount of tackifier is, by weight based on the sum of the weights of composition (Cb) and composition (Cc), 0.5% or more; more preferably 1% or more; more preferably 2% or more. Preferably, the amount of tackifier is, by weight based on the sum of the weights of composition (Cb) and composition (Cc), 12% or less; more preferably 9% or less; more preferably 8% or less.

Either composition (Cb) or composition (Cc) may independently of each other contain one or more additional ingredients. Preferably such additional ingredients are chosen to improve the performance of the PSA article. Typical additional ingredients include plasticizers, fillers, thickeners, pigments, antioxidants, UV stabilizers, defoamers, surfactants, and mixtures thereof.

Preferably, the pressure sensitive adhesive article of the present invention has good resistance to migration the tackifier that is present in composition (Cc). It is useful to consider parameter "T-Compat-b", which is the compatibility limit of the tackifier in the specific polymer (POLb) in which the tackifier has the highest compatibility limit. It is also useful to consider the parameter "T-Compat-c," which is the compatibility limit of the tackifier in the specific polymer (POLc) in which the tackifier has the lowest compatibility limit. Preferably, T-Compat-b is lower than T-Compat-c. Preferably, the tackifier is incompatible in every polymer (POLb).

Preferably, the weight ratio of composition (Cb) to composition (Cc) is 1.01:1 or greater; more preferably 2:1 or greater; more preferably 4:1 or greater; more preferably 6:1 or greater. Preferably, the weight ratio of composition (Cb) to composition (Cc) is 20:1 or less; more preferably 15:1 or less; more preferably 10:1 or less.

The layer of composition (Cb) may be applied to substrate (Sa) by any method. Preferably, an aqueous composition (Cbl) is formed that contains particles of polymer (POLb) dispersed in an aqueous continuous medium. Preferably, composition (Cbl) is formed by aqueous emulsion polymerization to form a latex of particles of polymer (POLb). Preferably, the median particle size of the particles of polymer (POLb) is from 50 nm to 750 nm. Preferably, the amount of water in the continuous medium, by weight based on the weight of the continuous medium, is 75% or more; more preferably 90% or more.

One or more optional additional ingredients may be added to aqueous composition (Cbl). A layer of aqueous composition (bl) may be applied to substrate (a) by any method. Preferred methods are slide coating, curtain coating, and slot die coating.

After a layer (Lbl) of aqueous composition (Cbl) is applied to substrate (Sa), the layer of aqueous composition (Cbl) may be dried or allowed to dry. That is, the water may
be removed from the layer of aqueous composition (bl) to form a dry coating. Water is preferably removed by the application of heat or moving air or both.

[0058] The layer (Lc) of composition (Cc) may be applied to the layer of composition (Cb) by any method. Preferably, an aqueous composition (Ccl) is formed that contains tackifier and contains particles of polymer (POLc) dispersed in an aqueous medium. Preferably, the median particle size of the particles of polymer (POLc) is from 50 nm to 1200 nm. Preferably, the amount of water in the continuous medium, by weight based on the weight of the continuous medium, is 75% or more; more preferably 90% or more. One or more optional additional ingredients may be added to aqueous composition (Ccl).

[0059] A layer (Lcl) of aqueous composition (Ccl) may be applied to the layer of composition (Cb) by any method. Preferred methods are slide coating, curtain coating, and slot die coating. A layer of aqueous composition (Ccl) may either be applied to a wet layer of aqueous composition (Cbl) or applied to a dry layer of composition (Cb).

[0060] In some embodiments, after a layer (Lbl) of aqueous composition (Cbl) is applied to substrate (Sa) and dried, then a layer (Lcl) of aqueous composition (Ccl) is applied to the dry layer of composition (Cb), and then the layer (Lcl) of aqueous composition (Ccl) is dried.

[0061] In some embodiments, a layer (Lbl) of aqueous composition (Cbl) is applied to substrate (Sa) and, while the layer (Lbl) of aqueous composition (Cbl) is still wet, a layer (Lcl) of aqueous composition (Ccl) is applied on top of the layer (Lbl) of aqueous composition (Cbl), and then the entire ensemble is then dried. Among such embodiments, it is preferred that a multilayer coating device is used that simultaneously applies a layer (Lbl) of aqueous composition (Cbl) to substrate (Sa) and also applies a layer (Lcl) of aqueous composition (Ccl) onto the layer (Lbl) of aqueous composition (Cbl), and then the entire ensemble is dried. One suitable coating device is a slide coater. A slide coater forms a liquid composite in which there is a layer (Lbl) of aqueous composition (Cbl) underneath a layer (Lcl) of aqueous composition (Ccl); while keeping those layers intact, the slide coater applies a layer of the entire composite onto a substrate (Sa) in a way that brings layer (Lbl) into contact with substrate (Sa) and leaves layer (Lcl) in contact with air; then the entire article is dried to remove water from the aqueous compositions. Also contemplated are transfer coating methods, for example methods that involve making a coating layer of composition (Cb) on a release liner, then making a coating layer of composition (Cc) on top of the layer of composition (Cb), then contacting the layer of composition (Cc) with substrate (Sa) (preferably under pressure), and then removing the release liner.
In addition to adhesion to polyolefin surfaces, the pressure sensitive adhesive article of the present invention desirably shows other useful characteristics, including, for example, a useful degree of adhesion to other types of surfaces and resistance to shear. Also, it is desirable that, when the pressure sensitive adhesive article of the present invention is stored for a time, little or no tackifier migrates from composition (Cc) into composition (Cb).

While the present invention is not bound by any theory, it is contemplated that one reason why acrylic PSAs often have adhesion to polyolefin surfaces that is considered to be poor is that acrylic polymers have many chemical groups on the chain that are relatively polar. It is contemplated that the surface of an acrylic PSA has a relatively high surface energy as a result. In contrast, polyolefin surfaces are comparatively non-polar and are considered to have relatively low surface energy. It is contemplated that this mismatch of surface energy values prevents the acrylic PSA from effectively wetting the surface of the polyolefin (i.e., from making intimate contact on a molecular scale). It is contemplated that the presence of tackifier provides a lower surface energy on the surface of the PSA, thereby improving wetting of polyolefin surface, which in turn improves adhesion to polyolefin surfaces.

It is contemplated that the pressure sensitive adhesive article of the present invention will be put to use by bringing it into contact with an additional substrate (Sd). It is contemplated that pressure will be applied to bring composition (Cc) and substrate (Sd) into intimate contact and then released. It is contemplated that the result will be a bonded article in which the pressure sensitive adhesive article is still intact and in which composition (Cc) is bonded with substrate (Sd). Substrate (Sd) may be any substance. Preferably, Substrate (Sd) is a polyolefin. Preferably, substrate (Sd) has Tg of 50°C or higher.

The following are examples of the present invention.

The materials used in the following examples are as follows.
<table>
<thead>
<tr>
<th>Label</th>
<th>Material</th>
<th>Description</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>PSA-L</td>
<td>PSA, low-shear type acrylic aqueous latex polymer</td>
<td>Polymerized Units more than 95%: mixture of BA and EA remainder: other acrylic monomers</td>
<td></td>
</tr>
<tr>
<td>PSA-H</td>
<td>PSA, low-shear type acrylic aqueous latex polymer</td>
<td>Polymerized Units more than 95%: mixture of BA and EA remainder: other acrylic monomers</td>
<td></td>
</tr>
<tr>
<td>SIS</td>
<td>Kraton™ D1161 P Copolymer</td>
<td>SIS triblock polymer aqueous dispersion</td>
<td>Kraton Performance Polymers</td>
</tr>
<tr>
<td>T1</td>
<td>Tacolyn™ 5003 resin</td>
<td>hydrocarbon resin tackifier aqueous dispersion</td>
<td>Eastman Chemical Co.</td>
</tr>
<tr>
<td>PET</td>
<td>PET film</td>
<td>polyethylene terephthalate film, thickness 50.8 μm</td>
<td>Chemsultants</td>
</tr>
<tr>
<td>HDPE</td>
<td>HDPE</td>
<td>high density polyethylene</td>
<td>Chemsultants</td>
</tr>
<tr>
<td>Blend 1</td>
<td>mixture of aqueous dispersions</td>
<td>Solids weight ratio of 45 SIS / 55 T1</td>
<td></td>
</tr>
</tbody>
</table>

[0067] The HDPE Peel test was PSTC Test Method 101 (Pressure Sensitive Tape council, Naperville, IL, USA), with test substrate of HDPE, 20 minutes dwell time, at 180°, reported in units of Newtons per 25 mm of width (N/25mm).

[0068] The SS Shear test was PSTC Test Method 107 (Pressure Sensitive Tape council, Naperville, IL, USA), with test substrate of stainless steel, test area was 25mm X 25 mm, mass was 1 kg. The results reported are time to failure (the symbol “>” means that the test was stopped prior to failure at the time shown).

[0069] Single-layer PSAs were made as follows. A layer of PSA-L was applied to the corona-treated side of PET film. The coating was then dried by placing in a oven for 5 minutes at 80°C. Dry thickness was 22.5 micrometers.

[0070] Two-layer PSAs were made as follows. A layer of PSA-L was applied and dried as in the single-layer PSA method. Thickness of the dry first coating was 20.0 micrometers.
Then a layer of Blendl was applied and dried as in the single-layer PSA method. Thickness of the dry second coating was 2.5 micrometers.

[0071] Example 1: Modification of a low-shear adhesive.

The results were as follows.

<table>
<thead>
<tr>
<th>Type</th>
<th>HDPE Peel (N/25mm)</th>
<th>SS Shear (hours)</th>
</tr>
</thead>
<tbody>
<tr>
<td>single layer</td>
<td>3.9</td>
<td>4.6</td>
</tr>
<tr>
<td>(comparative)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Two-Layer</td>
<td>5.4</td>
<td>5.4</td>
</tr>
</tbody>
</table>

[0073] The two-layer example of the present invention shows improvements in both peel strength on HDPE and in shear bonding to stainless steel.


Coatings were made as in Example 1 except that in all cases PSA-H was used in place of PSA-L.

[0075] The results were as follows.

<table>
<thead>
<tr>
<th>Type</th>
<th>HDPE Peel (N/25mm)</th>
<th>SS Shear (hours)</th>
</tr>
</thead>
<tbody>
<tr>
<td>single layer</td>
<td>1.2</td>
<td>58.5</td>
</tr>
<tr>
<td>(comparative)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Two-Layer</td>
<td>2.3</td>
<td>&gt;77</td>
</tr>
</tbody>
</table>

[0076] The two-layer example of the present invention shows improvements in both peel strength on HDPE and in shear bonding to stainless steel.
CLAIMS

1. A pressure sensitive adhesive article comprising
   (a) a substrate (Sa),
   (b) in contact with said substrate (Sa), a layer (Lb) of a tackifier-free composition (Cb) that comprises one or more acrylic polymer (POLb) having Tg of -10°C or lower, and
   (c) in contact with said layer (Lb), a layer (Lc) of a tackifier-containing composition (Cc) that comprises one or more tackifier and one or more hydrocarbon polymer (POLc) having Tg of -10°C.

2. The article of claim 1, wherein the amount of said tackifier in said article is 12% or less by weight, based on the sum of the weights of said composition (Cb) and said composition (Cc).

3. The article of claim 1, wherein said acrylic polymer (POLb) comprises polymerized units of n-butyl acrylate, iso-octyl acrylate, or a mixture thereof.

4. The article of claim 1, wherein said tackifier is incompatible with said acrylic polymer (POLb).

5. A bonded article made by a process comprising contacting a substrate (Sd) with the article of claim 1, wherein said substrate (Sd) is in contact with said layer (Lc).
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

INV. C09J7/02

ADD.

According to International Patent Classification (IPC) onto both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
C09J C09K

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)

EPO-Internal, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No.

X J.P S63 215787 A (YASUHARA YUSHI KOGYO KK) 8 September 1988 (1988-09-08) abstract

- - - -


- - - -

paragraph [0031] - paragraph [0034]

paragraph [0044] - paragraph [0046]

claims 1, 16, 17, 43, 16, 17, 51, 57, 67, 71

- - - -

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

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Date of the actual completion of the international search

2 June 2014

Date of mailing of the international search report

10/06/2014

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Si emsen, Beatrice
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| A        | A WO 00/17285 AI (AVERY DENNISON CORP [US]; K0 CHAN U [US]; CHUANG HSIAO K [US]; UGOLICK) 30 March 2000 (2000-03-30) page 1, line 5 - line 9  
            page 3, line 8 - line 11  
            page 3, line 35 - page 4, line 4  
            page 4, line 32 - line 36  
            claims 1, 13 | 1-5 |
            paragraph [0005]  
            paragraph [0012]  
            paragraph [0014]  
            paragraph [0017] - paragraph [0018]  
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