CONVENTION
AUSTRALIA
Patents Act 1990
REQUEST FOR A STANDARD PATENT

The Applicant identified below requests the grant of a patent to the nominated person identified below for an invention described in the accompanying standard complete patent specification.

[70,71] Applicant and Nominated Person:
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[54] Invention Title:
URETHANE ACRYLATE SURFACED LAMINATE

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[31,33,32]
Details of basic application(s):
577,483 UNITED STATES OF AMERICA US 5 September 1990

The nominated person is not an opponent or eligible person described in section 33-36 of the Act

3 September 1991
PHILLIPS ORMONDE & FITZPATRICK
Attorneys for:
Ralph Wilson Plastics Co.

By: [Signature]
Our Ref: 227145
An Australian patent request must be supported by a statement of entitlement. The statement may be filed with the request or subsequently. This comprehensive form is suitable for convention or non-convention requests by individuals or corporate bodies. For non-convention applications omit paragraphs 2, 3 and 4. Where the applicant is a company the person making the statement should be an officer of the company. Otherwise the applicant should make the statement personally.

AUSTRALIA

Notice of Entitlement

Suitable for any request
No legalization or other witness required
Filable after application

DK-9150-M45

NOTICE OF ENTITLEMENT

INSTRUCTIONS
(a) Name of person making statement.
(b) Position of that person.
(c) Name of applicant.
(d) Address of applicant.
(e) Delete as necessary.
(f) Insert details if not covered by (i) or (ii).
(g) Delete as necessary.
(h) Delete for non-convention applications.
(i) Insert DATE of signing.
(j) Signature(s) of person making statement.

I(a) Thomas M. Roehlk
(b) Assistant Secretary
(c) RALPH WILSON PLASTICS CO.
(d) 600 General Bruce Drive
   Temple, Texas 76501 U.S.A.

State the following:-
1. The nominated person (applicant) is entitled to the grant of a patent
   (i) as assignee of the actual inventor(s)
   (ii) by contract of employment of the actual inventor(s)
   or (iii) by virtue of an assignment dated: September 4, 1990

2. The nominated person (applicant) is entitled to claim priority from the basic
   convention application(s).
   (i) as applicants of the said application(s)
   (ii) as the assignee of the applicants of the said application(s)
   (iii) with the consent of the applicants of the said application(s)

3. The basic convention application(s) was/were the first made in a Convention
   country in respect of the invention the subject of the application. (h)

Dated (i) August 28, 1991

Thomas M. Roehlk
Assistant Secretary

To: The Commissioner of Patents
1. A decorative laminate, comprising:
   a thermoset resin impregnated core layer;
   a thermoset resin impregnated decorative layer
   bonded to the core layer;
   a protective layer bonded to the decorative layer, wherein the protective layer comprises a mixture of:

   (1) a polyfunctional unsaturated aliphatic acrylate oligomer;
   (2) a photoinitiator;
   (3) a thermally activated catalyst; and
   (4) a coupling agent.
AUSTRALIA
Patents Act

COMPLETE SPECIFICATION
(ORIGINAL)

Application Number:
Lodged:

Complete Specification Lodged:
Accepted:
Published:

Priority

Related Art:

Name of Applicant:
Ralph Wilson Plastics Co.

Actual Inventor(s):
Andrew Dion

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Invention Title:
URETHANE ACRYLATE SURFACED LAMINATE

Our Ref : 227145
POF Code: 36782/149436

The following statement is a full description of this invention, including
the best method of performing it known to applicant(s):

- 1 -

6006
APPLICATION FOR PATENT

INVENTOR: ANDREW N. DION
TITLE: URETHANE ACRYLATE SURFACED LAMINATE

BACKGROUND OF THE INVENTION

1. Field of the Invention

This invention relates to a urethane acrylate surfaced laminate, and to a method for its manufacture. In particular, this invention is of a decorative laminate which has as its outer surface a layer of radiation-polymerized urethane acrylates featuring enhanced scratch, wear and chemical resistance, and which shows markedly decreased tendencies to delaminate, and of a method of manufacturing such a laminate.

2. Description of the Prior Art

Laminates are widely used in the building and furniture industries for coating walls, furniture, and the like, to enhance their scratch, wear and chemical resistance while at the same time enhancing the aesthetic appeal of the coated item.

Laminates are typically prepared by pressing together at elevated temperatures a core layer and a covering layer, or decorative layer. The core layer may itself consist of a thermopressed stack of resinated paper webs or it may be plastic-based.

Paper-based laminates are susceptible to attack by mineral acids and, since the type of resin typically used in the covering layer is vulnerable to hydrolysis, are not sufficiently weather-resistant. Because of these shortcomings, laminates of this kind can only be used in certain limited benign applications. Whenever
it is desired to use laminates under more demanding circumstances, provisions must be made in order to improve the resistance of the laminates.

Plastic-based laminates, incorporating, for example, polyester or acrylate, are typically vulnerable to scratching and are not sufficiently resistant to organic solvents. With these laminates also, steps must be taken to improve the resistance of the laminates.

It is known in the art to enhance the scratch, wear, and chemical resistance of laminates by including as part of the laminate an outermost layer which has the desired properties. It is known to create such a protective layer from a synthetic resin of one or more radiation-polymerized components selected from unsaturated acrylates.

U.S. Patent No. 4,801,495, issued January 31, 1989, discloses a decorative laminate having a core layer and a decorative layer. In addition, the laminate features an outer layer made up of one or more radiation-polymerized components such as unsaturated acrylates or methacrylates. The outer layer has a scratch resistance of at least 1.5 Newtons. Also disclosed is a method of manufacturing the decorative laminate which includes applying the radiation-polymerizable components to the layer immediately beneath and then radiating the mixture to polymerize. The laminate is then consolidated by pressing together at elevated temperatures and pressures.

The technology disclosed in the '495 patent has several shortcomings, the most basic of which involves the tendency of the protective outer layer to delaminate, or disengage from the layers below. Because the bond between the outer protective layer and the layers immediately below is primarily mechanical,
the structure may be susceptible to a mode of failure wherein the outer protective layer, with time or upon bending or other stressing, peels off and separates from the layer beneath. Such a failure greatly reduces the commercial value of the laminate product.

It would thus be desirable to have a scratch, wear, and chemical resistant laminate wherein the protective outer layer is so tightly secured to the layer just beneath as to dramatically reduce the tendency of the product to delaminate, or separate into layers.

SUMMARY OF THE INVENTION

The present invention successfully addresses the shortcomings of the presently known decorative laminates by providing a laminate which features a protective outer layer providing superior scratch, wear, and chemical resistance, and which is so tightly bound to the layer immediately beneath as to dramatically reduce or eliminate the tendency of the protective layer to delaminate, or separate from the layer immediately below.

The laminate of this invention takes advantage of the highly desirable properties known to be associated with radiation-polymerizable urethane acrylate coated surfaces, yet provides a protective surface which is more securely bonded to the surface onto which the protective surface is coated than has heretofore been known to be possible.

This invention discloses a novel composition for the urethane acrylate protective layer and a novel method for creating and applying the protective layer to the surface to be protected.

The manufacturing process for the laminate of this invention involves two steps. The compounds are first
applied to a finish paper, or release sheet, and partly cured by UV radiation. The partly cured material is placed in contact with the surface to be protected, and the laminate is consolidated under heat and pressure. This lamination process completely cures the outer protective layer and bonds it to the surface.

The reaction mixture which will form the protective layer contains acrylate oligomers. The reaction mixture also contains one or more monomers. In addition, the mixture contains one or more photoinitiators which serve to initiate the polymerization during the UV curing. Also included are a thermal catalyst and a coupling agent. The thermal catalyst serves to promote the chemical cross-linking reaction during the thermopressing, while the coupling agent serves to chemically join the protective layer with the layer immediately beneath during the thermopressing step.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

The present invention is of a decorative laminate having enhanced scratch, wear, and chemical resistance properties as a result of the inclusion as the outermost layer of the laminate of a radiation-polymerized protective layer. The present invention is also a method for creating the protective layer and for securely bonding the protective layer onto the surface immediately beneath the protective layer.

It is known in the art that an outer layer made up of one or more synthetic resins which are polymerized by radiation can enhance the weather and scratch resistance of a laminate and can make it substantially less vulnerable to attack by acids and organic solvents.
In the preferred embodiment of this invention, a material made up of a urethane acrylate oligomer, one or more monomers, and one or more photoinitiator, thermally activated catalyst, and coupling agent, is applied to finish paper or release sheet. The material is then exposed to an appropriate amount of ultraviolet (UV) radiation. The photoinitiator serves to catalyze or initiate the polymerization reaction upon exposure to the radiation.

The extent of the exposure to radiation is controlled so as to limit the degree of polymerization of the resin to a desired extent. The partial curing serves to slightly harden the material, facilitating its handling during the subsequent manufacturing steps, while leaving a substantial amount of unpolymerized resin able to participate in the final cure during the thermopressing process step. During the final curing which takes place during the thermopressing, the protective layer is able to cross-link with the layer immediately beneath to create a strong bond of the two layers.

Once the material located on the finish paper has been partially cured, it can be cut up into appropriate sized sheets. Alternatively, the finish paper coated on one side with the partially cured material can be rolled up and stored for future use.

To apply the protective layer to a laminate, the finish paper, or release paper coated on one side with the partially cured material is placed on the surface to be protected, typically the decorative sheet, with the partially cured material in contact with the surface to be protected.

The protective layer and the underlying laminate are then pressed together at elevated temperatures. The heat activates a thermally sensitive catalyst in
the partially cured protective layer which, in turn, promotes the further polymerization of the resins in the protective layer.

A coupling agent present in the protective layer reacts with both the resins of the protective layer and that of the layer immediately beneath the protective layer, chemically binding the two layers.

The surface to be protected by the urethane acrylate outer layer typically forms a standard laminate composed of an internal core and an external decorative sheet. The core layer carries out the load-bearing function of the laminate. It can be made of wood or of other materials such as polyvinyl chloride, polyethylene, or other plastic materials. The core layer is more typically made up of sheets of thermopressed paper, such as soda kraft paper, impregnated with thermosetting synthetic resin, such as phenol-formaldehyde resin. Depending on the desired laminate thickness, from 1 to 100 or more sheets, stacked one on top of the other, can be thermopressed together to form the core.

The core layer may have on its outer surface an additional layer containing a thermosetting amino plastic or phenolic resin. Typically, a decorative sheet is applied directly onto the core layer or onto the additional layer. It is on the exterior surface of this decorative sheet that the radiation-polymerizable urethane acrylate protective layer is applied. The protective layer would typically be clear, transparent and pigment-free.

The decorative layer typically contains one or more thermosetting synthetic resins, such as amino plastic resin, and is located on core layers made up of the soda kraft paper, or of randomly deposited wood or cellulose fibers impregnated with phenol resin.
The radiation-polymerized synthetic resin protective layer is applied to the outer surface of the decorative layer. The protective layer contains urethane acrylates capable of being radically polymerized by UV radiation and which are present either individually or together in a polymerizable mixture.

The major component is a prepolymer or oligomer, which is polyfunctional, that is, which is multiply unsaturated. Preferred is an acrylate oligomer. The oligomer would ordinarily be a highly viscous, near-solid, with viscosity on the order of 2600 poise. In order to reduce the viscosity and render the oligomer easier to handle, a number of lower viscosity materials are typically added. Typically also present in the copolymerizable mixture, besides the predominant oligomeric component, may be additional components, usually diacrylates or similar monomers whose presence has a diluting effect on the mixture.

For example, the present invention can readily be practiced by using an acrylated urethane product known as Ebercryl 8800-20R, which is a commercial product of Radcure Specialties, Inc. of Louisville, Kentucky. According to the Material Safety Data Sheet, Ebercryl 8800-20R has the following composition:

<table>
<thead>
<tr>
<th>Component</th>
<th>Wt. %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acrylate oligomer</td>
<td>&gt;70.0</td>
</tr>
<tr>
<td>Ethoxyethoxyethyl acrylate</td>
<td>&lt; 8.0</td>
</tr>
<tr>
<td>Tripropylene glycol diacrylate (TRPGDA)</td>
<td>20.0</td>
</tr>
<tr>
<td>Polymeric isocyanate</td>
<td>&lt; 0.2</td>
</tr>
<tr>
<td>Hydroquinone</td>
<td>&lt; 0.1</td>
</tr>
</tbody>
</table>

The acrylate oligomer can be present in concentration of from about 40 to about 90% by weight, preferably from 40 to 70% by weight, and most preferably from 50 to 60% by weight. The Ebercryl 8800-20R has a viscosity at 65°C of from 1,800 to 3,000 cps. Depending on the amounts of additional components added, the
polyfunctional unsaturated aliphatic acrylate oligomer typically would constitute from about 30 to about 90 wt% of the total mixture, preferably from about 55 to about 85 wt% of the mixture.

The components used have a pronounced tendency to polymerize radically under the influence of UV radiation. The radically polymerizable prepolymer is a polyfunctional unsaturated aliphatic acrylate oligomer.

Examples of additional suitable monomers or oligomers that may be used in a radically copolymerizable mixture include, besides the prepolymer, a mono-, di-, tri-, tetra-, penta-, or hexacrylate. The acrylates in the mono- to hex-form are esters of mono or polyfunctional alcohols, i.e., polyols with 1 to 6 OH-groups with acrylic acid and are therefore also known as polyolacrylates. Suitable diacrylates are esters of acrylic acid with aliphatic divalent alcohols, especially ethylene glycol, 1,2-propylene glycol, 1,3-propylene glycol, butanediols, 1,6-hexanediol or neopentyl glycol; with aliphatic ether alcohols, in particular diethylene glycol, dipropylene glycol, dibutylene glycol, polyethylene glycals or polypropylene glycols; with oxyalkylated compounds of the above-named aliphatic alcohols and either alcohols; or also with aromatic dihydroxyl compounds, in particular bisphenol A, pyrocatechol, resorcinol, hydroquinone, p-xylylene glycol or p-hydroxybenzyl alcohol. Preferred diacrylates are 1,6-hexanediol diacrylate, tripropylene glycol diacrylate and 1,4-butanediol diacrylate. Preferred triacrylates are trimethylol propane triacrylate and pentacrythritol triacrylate.

Epoxy acrylate and silicone acrylate oligomers, which in addition to the urethane acrylate and unsaturated polyester acrylate oligomers mentioned above, are suitable polyfunctional prepolymers, can be used in
the radically copolymerizable mixture along with the
diacrylates or triacrylates already mentioned.

The prepolymers are produced, for example, from
hydroxylated copolymers in which the hydroxyl groups are
distributed along the copolymer chain. Unsaturated
acrylic copolymers are obtained from this copolymer by
esterification of the hydroxyl groups with acrylic acid.
To produce intermediate unsaturated acrylic copolymers,
the hydroxyl group is attached to the end of the chain
when the hydroxylated copolymers are produced. Urethane
acrylate oligomers are produced by the reaction of
acrylates or methacrylates containing hydroxy groups,
such as hydroxyethyl methacrylate, with multivalent
isocyanates, preferably diisocyanates. The di- or
polyisocyanates can preferably be reaction products of
diols, polyether diols, or polyester diols having a
stoichiometric excess of monomeric di- or polyisocyanate.

If the polyfunctional prepolymer predominates in the
polymerizable mixture, then, as the fundamental resin,
its chemical nature determines the characteristics of the
cured surface layer. As the diluting monomer, the added
acrylate or methacrylate in mono- to hex-form makes it
possible to adjust the viscosity of the mixture to be
cured, which is normally within the range from 20 to 100
poise (at 20°C), and participates fully in the radical
polymerization. In the process of irradiation, the
hardening of the coating is the result of the radical
polymerization at the double bonds of the prepolymer and,
if present, of the diluting monomers or oligomers.

As previously stated, it is useful to reduce the
viscosity of the uncured protective layer prior to
coating the finish paper with the material. The
reduction of the viscosity can be accomplished by any
convenient means, preferably by the addition of a lower
viscosity material, such as a di- or triacrylate. It is
preferable to add sufficient trimethylolpropane triacrylate (TMPTA) to lower the overall viscosity to 3000 cps, which is low enough to allow the mixture to be conveniently handled during processing yet high enough so that the mixture will not run off the finish paper prior to the partial curing promoted by the UV radiation. It is preferable to add sufficient TMPTA so that the concentration of the TMPTA in the overall mixture will be from 15 to 45% by weight, preferably from 25 to 35%, and most preferably from 32 to 34%. It should be noted that use of excessive amounts of TMPTA tends to render the protective layer brittle and more subject to cracking.

It may be desirable to use an acrylate of higher functionality as the diluting material. This could be used in addition to, or in place of, the di- or triacrylates. One possible high functionality acrylate is SR-399, sold by Sartomer Company of West Chester, Pennsylvania. SR-399 is the commercial name for dipentaerythritol hydroxypenta acrylate. The SR-399 has the advantage of providing enhanced chemical resistance over that which could be achieved with lower-functionality monomers.

The UV-polymerizable material must include at least one photoinitiator. A photoinitiator system comprising several components is usually used. Photoinitiators are catalysts which, upon absorption of ultraviolet light, form radicals, which, in turn, initiate the radical polymerization. Most photoinitiators include at least one carbonyl group, which carbonyl group is in conjunction with an aromatic ring.

The preferred photoinitiator is 2-hydroxy-2-methyl-1-phenyl-propan-1-one of the aromatic hydroxy ketone family. It is available commercially as Darocur 1173 from EM Industries, Inc. of Hawthorne, New York. Darocur
1173 has a chemical formula of C₁₀H₁₂O₂ and can be represented by the following chemical structure:

![Chemical Structure of 1173]

Darocur 1173 is a highly reactive photoinitiator. The compound is a hydrophobic liquid and can easily be incorporated in the UV-polymerizable material. Use of the photoinitiator does not result in yellowing of the transparent coating and the cured film shows excellent color stability. The Darocur 1173 has low viscosity and serves, in part, to slightly lower the overall viscosity of the uncured protective layer. The compound does not corrode the rollers used in the lamination process and its addition to UV-curable resins does not adversely affect the storage stability of the UV-curable system.

The Darocur 1173 is used in an amount of from about 0.2 to about 1.5% by weight of the entire UV-polymerizable mixture, preferably from 0.2 to 0.8%, most preferably from 0.3 to 0.5%.

The UV-polymerizable system also contains a thermally activated catalyst. A wide variety of compounds can be used. These can include catalysts that are activated at temperatures between 150 and 350°F, such as benzoyl peroxide, 2,5-dimethyl-2,5-bis-(2-ethylhexyl peroxy)hexane, t-amyl peroxyoctoate, t-butyl peroxyoctoate, and lauroyl peroxide, 1,1-bis-t-butyl peroxycyclohexane, 1,1-bis-t-amyl peroxy cyclohexane, and dicumyl peroxide. Preferably, the thermally activated catalyst is tertiary-butyl peroxybenzoate.
The thermally activated catalyst is used to initiate polymerization reactions during the thermopressing step of the manufacture of the laminates. The thermopressing step cures the protective layer and bonds it to the decorative layer. The two layers cross-link so as to reduce or eliminate the tendency of the protective layer to delaminate, or peel off, from the layer immediately beneath.

The uncured protective layer also includes a coupling agent capable of chemically binding the polymerizable materials of the protective layer with those of the layer immediately beneath. Organic titanate esters and organosilane coupling agents enhance the bond between dissimilar resins due to their bifunctional nature. One end of the molecule reacts with the protective layer while the other reacts with the layer immediately beneath. Additives capable of functioning in this manner, include, but are not limited to, amino methoxy silanes, methacryloxypropyl tri-methoxy silane, and epoxy methoxy silanes. Another, and preferred, coupling agent is glycidoxypropyltrimethoxysilane, available commercially as Z-6040 Silane from Dow Corning. The Z-6040 Silane is used in concentrations of from 0.5 to 3% by weight of the total mixture, preferably from 0.75 to 1.5%, and most preferably from 1.0 to 1.3%. The chemical binding between the protective and the lower layers brought about by the Z-6040 Silane serves to dramatically reduce the tendency of the protective layer to peel off, or delaminate, with time and use. The use of the coupling agent serves to render the protective layer an integral part of the laminate and thus provides long-lasting protection of the laminate.

The synthetic resin polymerized by radiation may additionally contain the usual additives, such as plasticizers, fillers, pigments, and stabilizers, for
attaining the desired decorative, mechanical, and physical surface characteristics. Examples of these substances include barium sulfate, silica, aluminum oxide, and pigments that are colorfast when exposed to light. For most uses it would be desirable to have a transparent pigment-free protective layer.

To produce the decorative laminate panel, the transparent, liquid, radiation-polymerizable compounds and the various additives are applied in some convenient manner to the release sheet, or finish paper. The radiation-polymerizable mixture may, if desirable, contain certain pigments so as to give the protective layer a certain hue while still maintaining its transparency. Typically, a 4 mil thick layer of the radiation-polymerizable mixture is preferred, but any convenient thickness may be used.

The finish paper, or release sheet, containing the radiation-polymerizable mixture on one side is exposed to UV radiation to partially cure the resin in order to provide a tack free surface. This step is generally known as B-stage. To initiate the polymerization effected by radiation, a conventional source for forming free radicals, such as a photoinitiator, is preferred.

The partial curing of the resin effected by the photoinitiators facilitates the further handling of the finish paper and protective layer. The UV radiation can be generated by any convenient means, for example through medium-pressure mercury arc lamps having wavelengths of between about 200 and 400 nanometers. A typical lamp power output is in the range of about from 150 W/in to 300 W/in. Typical curing times are about in the range of 2-4 sec at line speeds of 400-500 in/min. The finish paper containing the partially cured protective layer can be rolled and stored for future use. Alternatively, it can be cut up into sheets of appropriate lengths. The
sheets can either be stored for future use or they can be immediately utilized in the manufacture of high resistance laminates.

In the next step of the process, the finish paper containing the partially cured protective layer is placed on top of a decorative layer which had been previously impregnated with a resin, such as melamine, with the partially cured protective layer in contact with the decorative layer. This assembly rests on one or more core sheets which provide the laminate with thickness.

The finish paper, the decorative print sheet, and the core layers are assembled and thermopressed together. In this step, the layers are pressed together under a pressure of between 800 and 1400 psi, preferably between 1000 and 1200 psi, at elevated temperatures of from 225 to 350°F, preferably from 260 to 280°F. The reaction time can range from about 30 to about 90 minutes. The exact times and temperatures for a specific product can readily be determined by one skilled in the art with routine experimentation.

The fiber-containing layers of the stack, which preferably comprise soda kraft papers or a bonded fabric made of wood and/or cellulose fibers, contain the thermosetting resins typical in high pressure laminates, in particular phenol-formaldehyde resins, while the decorative papers which are also be present in the stack contain an amino plastic resin, in particular a malamine resin. The proportion of thermosetting resins amounts to 20 to 70% by weight, preferably about 50%, based on the weight of an individual layer. The thermosetting resins of the core and decorative layers are pre-cured and dried in the conventional manner.

The saturation or impregnation of the fiber-containing layers or paper layers is effected by, for example, immersion in a bath having a solution or
dispersion containing the thermosetting resin, or by applying or spraying by means of a dispensing system. The solvent or dispersing agent is of the aqueous alcohol, aqueous acetone, or aqueous type, depending on the synthetic resin used. It may also contain up to 20% by weight of a flame retardant.

As in the conventional manufacture of high pressure laminates, the packet, comprising the fiber-containing core layer, the decorative layer, and the radiation-polymerized surface layer is thermopressed to form a decorative panel causing the thermosetting resins to completely cure. However, if the core layer comprises a wood, plastic, or metal panel, then the temperature and pressure can usually be reduced to 80°C, and 5 bar, respectively.

The pressing is effected in either a stationary, once-through device, or in a continuous press apparatus. The number and thickness of the fiber-containing layers in the core layer, or the thickness of the core layer, is selected depending on the intended use of the panel.

During the thermopressing, the thermal catalyst, such as TBPB, serves to promote the reaction of the resins. In the presence of a coupling agent, such as the Z-6040 Silane, reaction also occurs between the resins of the protective and those of the decorative layers immediately beneath, which serves to chemically bind the two layers together. It is during the thermopressing process that the final consolidation and cure takes place for all components of the laminate. The acrylated urethane film is thus transferred from the finish paper to the decorative print sheet. At the conclusion of the thermopressing step, the finish paper, which imparts gloss and texture to the laminate surface, is removed and discarded.
The resulting laminate has improved scratch, chemical and wear resistance over conventional laminates not having a protective layer. Furthermore, the laminate of the present invention has superior delamination characteristics compared with laminates formed without the benefit of a coupling agent and/or a thermally activated catalyst.

EXAMPLE

A laminate was manufactured as described above. The formulation used was as follows:

<table>
<thead>
<tr>
<th>Amount (gms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ebercryl 8800-20R</td>
</tr>
<tr>
<td>TMPTA</td>
</tr>
<tr>
<td>Darocur 1173</td>
</tr>
<tr>
<td>TBPB</td>
</tr>
<tr>
<td>Z-6040 Silane</td>
</tr>
</tbody>
</table>

The urethane acrylate surface showed a scratch resistance of 7 Newtons, compared with 4 Newtons for a standard laminate. The scratch resistance test was carried out in accordance with method ISO 4586/2, Clause 14. In this test, the minimum load applied to a diamond scratching point, which produces a continuous surface mark visible to the naked eye, is the resistance to scratching of the surface. The minimum load is determined by applying successively decreasing loads to the diamond point and observing the effect.

The urethane acrylate surface of the present invention had a wear resistance of 1428, compared with 793 for a standard laminate. The wear resistance was measured in accordance with the method of NEMA LD3.1-1985. In this test, 180 grit sandpaper is applied to the wheels of a Taber abrading machine, Model 503. The number of revolutions is used to calculate the wear resistance.
The chemical resistance of the urethane acrylate surface of the present invention and of the standard laminate were tested by placing a drop of each chemical on the surface, covering the drop with a watchglass, and noting the effect, if any, on the laminate surface after 16 hours. The results were as follows:

<table>
<thead>
<tr>
<th>This Invention</th>
<th>Standard Laminate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slight Effect:</td>
<td>Glacial acetic acid</td>
</tr>
<tr>
<td>Glacial acetic acid</td>
<td>Acetic acid 98%</td>
</tr>
<tr>
<td>Sulfuric 25%</td>
<td>Uric acid</td>
</tr>
<tr>
<td>Sulfuric 60%</td>
<td>Sodium hydroxide 5%</td>
</tr>
<tr>
<td>Sulfuric 77%</td>
<td>Sodium hydroxide 10%</td>
</tr>
<tr>
<td>Sulfuric 96%</td>
<td>Ag eosin bluish 5%</td>
</tr>
<tr>
<td>Hydrochloric 37%</td>
<td>(in water)</td>
</tr>
<tr>
<td>Tincture of Iodine</td>
<td></td>
</tr>
</tbody>
</table>

Severe Effect: Safranin O

| Nitric 25% |
| Nitric 30% |
| Sulfuric 25% |
| Sulfuric 60% |
| Sulfuric 77% |
| Sulfuric 96% |
| Hydrochloric 20% |
| Hydrochloric 30% |
| Hydrochloric 37% |
| Phosphoric 75% |
| Phosphoric 90% |
| Formic 50% |
| Formic 90% |
| Hydrofluoric acid 48% |
| Aqua Regia |
| Perchloric acid |
| Sodium hypochlorite 5% |
| Cupra ammonia |
| Zinc chloride |
| Salt mixture |
| Cleaning Solution |
| Phosphorus pentoxide |
| Silver nitrate 10% |

The delamination characteristics of the urethane acrylate surface of the present invention and of the standard laminate were tested by immersion into boiling water for five hours. The coating containing the coupling agent did not blister or delaminate from the laminate surface.
While the invention is described in accordance with the specific examples below, it is understood that these are only for illustrative purposes. Many alternatives, modifications and variations will be apparent to those skilled in the art in the light of the below examples and as such will fall within the general scope of the claims.

The present invention and the embodiments disclosed herein are well adapted to carry out the objectives and obtain the ends set forth at the outset. Certain changes can be made in the method without departing from the spirit and the scope of this invention. It is realized that changes are possible and it is further intended that each element recited in any of the following claims is to be understood as referring to all equivalent elements for accomplishing substantially the same results in substantially the same or equivalent manner. It is intended to cover the invention broadly in whatever form its principles may be utilized.
The claims defining the invention are as follows:

1. A decorative laminate, comprising:
   a thermoset resin impregnated core layer;
   a thermoset resin impregnated decorative layer bonded to the core layer;
   a protective layer bonded to the decorative layer, wherein the protective layer comprises a mixture of:
   (1) a polyfunctional unsaturated aliphatic acrylate oligomer;
   (2) a photoinitiator;
   (3) a thermally activated catalyst; and
   (4) a coupling agent.

2. A decorative laminate of claim 1 wherein the mixture further includes a monomer.

3. A decorative laminate of claim 2 wherein the monomer is a mono-, di-, tri-, tetra-, penta-, or hexacrylate.

4. A decorative laminate of claim 2 wherein the monomer is 1,6-hexanediol diacrylate, tripropylene glycol diacrylate, 1,4-butanediol diacrylate, trimethylol propane triacrylate, or pentacrythritol triacrylate.

5. A decorative laminate of claim 4 wherein the monomer is trimethylolpropane triacrylate.

6. A decorative laminate of claim 2 wherein the monomer is dipentaerythritol hydroxypenta acrylate.

7. A decorative laminate of claim 1 wherein the monomer is from about 15 to about 45 wt% of the mixture.
8. A decorative laminate of claim 1 wherein the monomer is from about 25 to about 35 wt% of the mixture.

9. A decorative laminate of claim 1 wherein the mixture further includes a plasticizer, filler, pigment, or stabilizer.

10. A decorative laminate of claim 1 wherein the polyfunctional unsaturated aliphatic acrylate oligomer is an acrylated urethane.

11. A decorative laminate of claim 1 wherein the polyfunctional unsaturated aliphatic acrylate oligomer is from about 55 to about 85 wt% of the mixture.

12. A decorative laminate of claim 1 wherein the photoinitiator is an aromatic hydroxy compound.

13. A decorative laminate of claim 1 wherein the photoinitiator is 2-hydroxy-2-methyl-1-phenyl-propan-1-one.

14. A decorative laminate of claim 1 wherein the photoinitiator is from about 0.2 to about 1.5 wt% of the mixture.

15. A decorative laminate of claim 1 wherein the photoinitiator is from about 0.2 to about 0.8 wt% of the mixture.
16. A decorative laminate of claim 1 wherein the thermally activated catalyst is benzoyl peroxide, 2,5-dimethyl-2,5-bis-(2-ethylhexyl peroxy)hexane, t-amyl peroxyoctoate, t-butyl peroxyoctoate, lauroyl peroxide, 1,1-bis-t-butyl peroxy cyclohexane, 1,1-bis-t-amyl peroxy cyclohexane, or dicumyl peroxide.

17. A decorative laminate of claim 1 wherein the thermally activated catalyst is tertiary-butyl peroxybenzoate.

18. A decorative laminate of claim 1 wherein the thermally activated catalyst is from about 0.2 to about 1.5 wt% of the mixture.

19. A decorative laminate of claim 1 wherein the thermal activated catalyst is from about 0.2 to about 0.8 wt% of the mixture.

20. A decorative laminate of claim 1 wherein the coupling agent is an organic titanate ester or an organosilane.

21. A decorative laminate of claim 1 wherein the coupling agent is an amino methoxy silane, methacryloxypropyl tri-methoxy silane, or an epoxy methoxy silanes.

22. A decorative laminate of claim 1 wherein the coupling agent is glycidoxypropyltrimethoxy silane.

23. A decorative laminate of claim 1 wherein the coupling agent is from about 0.5 to about 3.0 wt% of the mixture.
24. A process for manufacturing a decorative laminate, comprising:
   coating on a release sheet a mixture which comprises:
   (1) a polyfunctional unsaturated aliphatic acrylate oligomer;
   (2) a photoinitiator;
   (3) a thermally activated catalyst; and
   (4) a coupling agent;
   partially curing the mixture on the release sheet with ultraviolet radiation, to form a partially cured protective layer;
   placing the release sheet containing the partially cured protective layer on top of a partially cured thermosetting resin impregnated decorative layer with the protective coating in contact with the top surface of the decorative layer;
   placing the decorative layer over a partially cured thermosetting resin impregnated core layer; and
   thermopressing the protective layer, decorative layer, and core layer, to form the decorative laminate.

25. A process for manufacturing a decorative laminate of claim 24 wherein the mixture further includes a monomer.

26. A process for manufacturing a decorative laminate of claim 25 wherein the monomer is a mono-, di-, tri-, tetra-, penta-, or hexacrylate.

27. A process for manufacturing a decorative laminate of claim 25 wherein the monomer is 1,6-hexanediol diacrylate, tripropylene glycol diacrylate, 1,4-butanediol diacrylate, trimethylol propane triacrylate, or pentacrythritol triacrylate.
28. A process for manufacturing a decorative laminate of claim 27 wherein the monomer is trimethylolpropane triacrylate.

29. A process for manufacturing a decorative laminate of claim 25 wherein the monomer is dipentaerythritol hydroxypenta acrylate.

30. A process for manufacturing a decorative laminate of claim 24 wherein the monomer is from about 15 to about 45 wt% of the mixture.

31. A process for manufacturing a decorative laminate of claim 24 wherein the monomer is from about 25 to about 35 wt% of the mixture.

32. A process for manufacturing a decorative laminate of claim 24 wherein the mixture further includes a plasticizer, filler, pigment, or stabilizer.

33. A process for manufacturing a decorative laminate of claim 24 wherein the polyfunctional unsaturated aliphatic acrylate oligomer is an acrylated urethane.

34. A process for manufacturing a decorative laminate of claim 24 wherein the polyfunctional unsaturated aliphatic acrylate oligomer is from about 55 to about 85 wt% of the mixture.

35. A process for manufacturing a decorative laminate of claim 24 wherein the photoinitiator is an aromatic hydroxy compound.
36. A process for manufacturing a decorative laminate of claim 24 wherein the photoinitiator is 2-hydroxy-2-methyl-1-phenyl-propan-1-one.

37. A process for manufacturing a decorative laminate of claim 24 wherein the photoinitiator is from about 0.2 to about 1.5 wt% of the mixture.

38. A process for manufacturing a decorative laminate of claim 24 wherein the photoinitiator is from about 0.2 to about 0.8 wt% of the mixture.

39. A process for manufacturing a decorative laminate of claim 24 wherein the thermally activated catalyst is benzoyl peroxide, 2,5-dimethyl-2,5-bis-(2-ethylhexyl peroxy)hexane, t-amyl peroxyoctoate, t-butyl peroxyoctoate, lauroyl peroxide, 1,1-bis-t-butyl peroxy cyclohexane, 1,1-bis-t-amyl peroxy cyclohexane, or dicumyl peroxide.

40. A process for manufacturing a decorative laminate of claim 24 wherein the thermally activated catalyst is tertiary-butyl peroxybenzoate.

41. A process for manufacturing a decorative laminate of claim 24 wherein the thermally activated catalyst is from about 0.2 to about 1.5 wt% of the mixture.

42. A process for manufacturing a decorative laminate of claim 24 wherein the coupling agent is an organic titanate ester or an organosilane.
43. A process for manufacturing a decorative laminate of claim 24 wherein the coupling agent is an amino methoxy silane, methacryloxypropyl tri-methoxy silane, or an epoxy methoxy silanes.

44. A process for manufacturing a decorative laminate of claim 24 wherein the coupling agent is glycidoxypropyltrimethoxysilane.

45. A process for manufacturing a decorative laminate of claim 24 wherein the coupling agent is from about 0.5 to about 3.0 wt% of the mixture.

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