Title: ONE-COMPONENT, WATERBORNE FILM-FORMING COMPOSITION

Abstract: A film-forming composition that includes a stable aqueous dispersion of a mixture of a polyester polyol and a carboxylic acid-containing acrylic polymer polyol and a crosslinking agent having functional groups reactive with hydroxyl groups. The weight ratio of polyester polyol to acrylic polymer polyol is in the range of 10:90 to 30:70. The crosslinking agent is one or more blocked isocyanates or aminoplasts. The equivalent ratio of hydroxyl groups to reactive functional groups on the crosslinking agent is from 0.8:1 to 1.7:1. The film forming composition can be used in multi-layer composite coating compositions that include a base coat layer and a substantially pigment-free topcoat deposited over at least a portion of the base coat layer, the topcoat composition including the present film-forming composition.
ONE-COMPONENT, WATERBORNE FILM-FORMING COMPOSITION

BACKGROUND OF THE INVENTION

1. Field of the Invention

The present invention relates to aqueous coating compositions as well as multi-component composite coating compositions including pigmented or colored base coats overcoated with transparent or clear topcoats, providing good smoothness and appearance in automotive coating applications while having a low volatile organic compound content.

2. Background of the Invention

Over the past decade, there has been a concerted effort to reduce atmospheric pollution caused by volatile solvents that are emitted during the painting process. However, it is often difficult to achieve high quality, smooth coating finishes, such as are required in the automotive industry, without using organic solvents which contribute greatly to flow and leveling of a coating.

One of the major goals of the coatings industry is to minimize the use of organic solvents by formulating waterborne coating compositions which provide a smooth, high gloss appearance, as well as good physical properties including resistance to acid rain. Unfortunately, many waterborne coating compositions do not provide acceptable appearance. Many automotive manufacturers are interested in coatings that contain low amounts of volatile organic compounds (VOCs).

Another challenge to formulators of waterborne coatings is to provide good acid resistance while maintaining acceptable physical properties. Lack of humidity resistance or blushing is another problem facing waterborne coating formulators.

U.S. Patent No. 4,350,809 to Fischer et al. discloses a process for the preparation of copolymers containing hydroxyl groups by free radical polymerization of at least two unsaturated copolymerizable monomers, at least one of which contains at least one carboxyl group, in the presence of at least one alkyl glycidyl ester of an aliphatic saturated monocarboxylic acid with a tertiary or quaternary α-carbon atom. The resulting polymers are advantageous because of their relatively low solution viscosities.
U.S. Patent No. 5,663,265 to Eppe et al. discloses low-viscosity copolymers useful in coating compositions. The copolymers contain hydroxyl and carboxyl groups and are obtained by free-radical polymerization of at least two olefinically unsaturated copolymerizable monomers of which at least one contains at least one carboxyl group and at least one is sterically hindered in the presence of one or more glycidyl esters of aliphatic saturated monocarboxylic acids having a tertiary or quaternary α-carbon atom. The copolymers are useful in clearcoat coating compositions, which demonstrate notable resistance to sulfuric acid and xylene, as well as a high degree of hardness.

U.S. Patent No. 5,596,057 to Eppe et al. discloses low viscosity copolymers prepared by bulk polymerization. The polymerization includes charging a material with functional groups that react with the carboxyl groups of the monomers and polymerizing at least two olefinically unsaturated copolymerizable monomers, at least one of which contains at least one carboxyl group and at least one of which is sterically-hindered. The copolymers are useful in coating compositions.

U.S. Patent No. 4,322,508 to Peng et al. discloses a thermosetting coating composition which includes a hydroxy functional component and a crosslinking agent capable of reacting with hydroxy functionality of the hydroxy functional component. The hydroxy functional component includes an oligoester formed by an esterification reaction between a carboxylic acid and an epoxide and a hydroxy functional copolymer. The hydroxy functional copolymer includes residues from one or more hydroxy functional monomers. The thermosetting compositions have a short shelf life, however, because the oligoester and hydroxy functional copolymer in the hydroxy functional component tend to separate over time.

It would be desirable to provide a coating composition which provides the benefits of including a combination of hydroxy functional polyesters and hydroxy functional acrylic copolymers, such that they do not tend to separate over time, providing a storage stable coating composition. The resultant coatings should have a balance of physical properties, including high gloss, hardness, impact resistance, flexibility, weatherability and chemical resistance.
SUMMARY OF THE INVENTION

The present invention is directed to a one-component, waterborne film-forming composition comprising:

(A) a stable aqueous dispersion of a mixture of a polyester polyl and a carboxylic acid-containing acrylic polymer polyl, wherein the weight ratio of polyester polyl to acrylic polymer polyl is in the range of 10:90 to 30:70; and

(B) a crosslinking agent having functional groups reactive with the hydroxyl groups in (A) selected from the group of blocked isocyanates and aminoplasts; wherein the equivalent ratio of hydroxyl groups in (A) to reactive functional groups in (B) is from 0.8:1 to 1.7:1.

The present invention is further directed to a multi-layer composite coating that includes:

(I) a base coat layer deposited from a pigmented film-forming base coat composition; and

(II) a substantially pigment free topcoat deposited over at least a portion of the base coat layer (I) from a topcoat composition that includes the one-component, waterborne film-forming composition described above.

The present invention is also directed to a coated substrate. The coated substrate includes a substrate and the one-component, waterborne film-forming composition described above over at least a portion of the substrate.

The present invention is additionally directed to a coated substrate that includes a substrate and the multi-layer composite coating composition described above over at least a portion of the substrate.

DETAILED DESCRIPTION OF THE INVENTION

Other than in the operating examples, or where otherwise indicated, all numbers expressing quantities of ingredients, reaction conditions and so forth used in the specification and claims are to be understood as being modified in all instances by the term "about." Accordingly, unless indicated to the contrary, the numerical
parameters set forth in the following specification and attached claims are approximations that may vary depending upon the desired properties sought to be obtained by the present invention. At the very least, and not as an attempt to limit the application of the doctrine of equivalents to the scope of the claims, each numerical parameter should at least be construed in light of the number of reported significant digits and by applying ordinary rounding techniques.

Notwithstanding that the numerical ranges and parameters setting forth the broad scope of the invention are approximations, the numerical values set forth in the specific examples are reported as precisely as possible. Any numerical values, however, inherently contain certain errors necessarily resulting from the standard deviation found in their respective testing measurements.

Also, it should be understood that any numerical range recited herein is intended to include all sub-ranges subsumed therein. For example, a range of “1 to 10” is intended to include all sub-ranges between and including the recited minimum value of 1 and the recited maximum value of 10, that is, having a minimum value equal to or greater than 1 and a maximum value of equal to or less than 10. Because the disclosed numerical ranges are continuous, they include every value between the minimum and maximum values. Unless expressly indicated otherwise, the various numerical ranges specified in this application are approximations.

As used herein, the term “substantially free” is meant to indicate that a material is present as an incidental impurity. In other words, the material is not intentionally added to an indicated composition, but may be present at minor or inconsequential levels because it was carried over as an impurity as part of an intended composition component.

As used herein, by “thermosetting composition” is meant one which “sets” irreversibly upon curing or crosslinking, wherein the polymer chains of the polymeric components are joined together by covalent bonds. This property is usually associated with a cross-linking reaction of the composition constituents often induced by heat or radiation. Hawley, Gessner G., The Condensed Chemical Dictionary, Ninth Edition, page 856; Surface Coatings, vol. 2, Oil and Colour Chemists' Association, Australia, TAFE Educational Books (1974). Once cured or crosslinked,
a thermosetting composition will not melt upon the application of heat and is insoluble in solvents. By contrast, a “thermoplastic composition” comprises polymeric components which are not joined by covalent bonds and thereby can undergo liquid flow upon heating and are soluble in solvents. Saunders, K.J., *Organic Polymer Chemistry*, pp. 41-42, Chapman and Hall, London (1973).

As used herein, “polymer” is meant to encompass oligomer, and includes without limitation both homopolymers and copolymers. Also, as used herein, “reactive” refers to a functional group that forms a covalent bond with another functional group under conditions sufficient to cure the composition. As used herein, “(meth)acrylate” and like terms are intended to include both acrylates and methacrylates.

As used herein, “substantially pigment-free coating composition” means a coating composition which forms a transparent coating, such as a clearcoat in a multi-component composite coating composition. Such compositions are sufficiently free of pigment or particles such that the optical properties of the resultant coatings are not seriously compromised. As used herein, “transparent” means that the cured coating has a BYK Haze index of less than 50 as measured using a BYK/Haze Gloss instrument.

As used herein, the phrase components “are different from each other” refers to components which do not have the same chemical structure as other components in the composition.

As used herein, “cure” as used in connection with a composition, e.g., “composition when cured,” means that any crosslinkable components of the composition are at least partially crosslinked. In certain embodiments of the present invention, the crosslink density of the crosslinkable components, i.e., the degree of crosslinking, ranges from 5% to 100% of complete crosslinking.

As used herein, “stable dispersion” refers to a liquid having a liquid continuous phase and a dispersed phase, which may be a liquid, a solid or a combination thereof, where the dispersed phase does not agglomerate, coalesce, settle or separate from the continuous phase between the period of time the dispersion is
prepared and when it is used, typically a period of time not exceeding two years at ambient conditions.

The multi-component composite coating of the present invention is useful in a variety of coating applications, and is particularly useful in automotive coating applications. The multi-component composite coating composition comprises a base coat layer and a transparent or clear topcoat layer formed from an aqueous topcoat coating composition which is applied over the base coat.

The present one-component, waterborne film-forming composition includes:

(A) a stable aqueous dispersion of a combination of a polyester polyol and a carboxylic acid-containing acrylic polymer polyol; and

(B) a crosslinking agent having functional groups that are reactive with the hydroxyl groups in (A). The crosslinking agents are typically blocked polyisocyanates and/or aminoplasts. The film-forming compositions of the present invention typically are “thermosetting compositions” as defined above.

The polyester polyol component in (A) provides flow and leveling properties to the waterborne film-forming composition. These physical properties of the composition lead to a smooth topcoat film in the multi-component composite coating. The polyester polyol is present in an amount sufficient to provide a smooth film as indicated by the occurrence of no more than three craters in an 4 inch (10.2 cm) by 10 inch (25.4 cm) coated area.

The acrylic polymer polyol component in (A) provides film properties such as hardness, gloss, acid resistance and delamination resistance in the topcoat film in the multi-component composite coating composition. The acrylic polymer polyol is present in an amount sufficient to provide a 20° gloss as measured using a Novo Gloss Statistical Glossmeter (Paul N. Gardner Company, Inc., Pompano Beach, FL) of at least 90, typically at least 95 of the multi-component composite coating composition.

The weight ratio of the polyester polyol to the acrylic polymer polyol in (A) may be 10:90 or higher, in some cases 12.5:87.5 or higher, in other cases 15:85 or higher and at other times 17.5:82.5 or higher. Additionally, the weight ratio of the polyester polyol to the acrylic polymer polyol in (A) may be 30:70 or lower, in some cases 27.5:72.5 or lower, in other cases 25:75 or lower and at other times 22.5 to 77.5.
The weight ratio of the polyester polyol to the acrylic polymer polyol in (A) may be 20:80. The higher and lower designation refers to the level of the polyester polyol. The weight ratio of the polyester polyol to the acrylic polymer polyol is determined by the properties that are to be incorporated into the resulting coating. The weight ratio of the polyester polyol to the acrylic polymer polyol may be any value or any range of values inclusive of those stated above.

When the crosslinking agent of (B) comprises a blocked polyisocyanate, the equivalent ratio of hydroxyl groups in (A) to isocyanate groups in (B) may be 0.8:1 or higher, in some cases 1:1 or higher, in other cases 1:1.1 or higher, in some instances 1.2:1 or higher and at other times 1.35:1 or higher. Additionally, the equivalent ratio of hydroxyl groups in (A) to isocyanate groups in (B) may be 1.7:1 or lower, in some cases 1.65:1 or lower, in other cases 1.6:1 or lower, in some instances 1.55:1 or lower and at other times 1.5:1 or lower. The equivalent ratio of hydroxyl groups in (A) to isocyanate groups in (B) may be 1.4:1. The higher and lower designation refers to the level of the hydroxyl groups. The equivalent ratio of hydroxyl groups in (A) to isocyanate groups in (B) is determined by the properties that are to be incorporated into the resulting coating. When the amount of isocyanate functionality in (B) is too high, the composition may react and/or may be unstable. When the hydroxyl functionality is too high, an undesirable soft film coating may result. The equivalent ratio of hydroxyl groups in (A) to isocyanate groups in (B) may be any value or any range of values inclusive of those stated above.

As used herein, "equivalent ratio" means the ratio of chemical equivalents of hydroxyl functionality to chemical equivalents of functional groups that are reactive with hydroxyl groups. A chemical equivalent is the quantity of a material that supplies one mole of a functional group. T.L. Brown and H.E. LeMay, Chemistry: The Central Science, page 346, 1977.

When the crosslinking agent of (B) is an aminoplast, it can be present in the film-forming composition in an amount of at least 5 percent by weight, in some cases at least 10 percent by weight, in other cases at least 15 percent by weight, in some instances at least 20 percent by weight and at other times 20 percent by weight based on the total resin solids in the film-forming composition. Additionally, the amount of
the aminoplast crosslinking agent (B) may be up to 60 percent by weight, in some cases up to 55 percent by weight, in other cases up to 50 percent by weight, in some instances up to 40 percent by weight and at other times up to 30 percent by weight based on the total resin solids in the film-forming composition. The amount of aminoplast crosslinking agent (B) is determined by the properties that are to be incorporated into the resulting coating. When the amount of aminoplast is too high, the composition may react and/or may be unstable. When the amount of aminoplast is too low, an undesirable soft film coating may result. The amount of aminoplast crosslinking agent (B) present in the film-forming composition may be any value or any range of values inclusive of those stated above.

In an embodiment of the present invention, component (A) may be prepared by polymerizing a mixture of ethylenically unsaturated polymerizable monomers in the presence of at least one glycidyl ester of an aliphatic saturated monocarboxylic acid and at least one polyester polyl. In this embodiment, the monomer mixture includes at least one carboxylic acid group-containing monomer and at least one primary hydroxyl group-containing monomer, which form an acrylic polyl. The glycidyl ester of an aliphatic saturated monocarboxylic acid acts as a solvent for the reactants.

Typically, when the polymerization is complete, an amine is added to component (A) in an amount sufficient to provide a pH of from 7 to 10 when component (A) is dispersed in water. In an embodiment of the present invention, the pH is increased by using one or more suitable volatile amines. Examples of suitable volatile amines include, but are not limited to, dimethylethanolamine, ammonia, triethyl amine and diethyl propanol amine.

The polymerization described above is generally carried out by introducing the monomer mixture that includes at least one carboxylic acid group-containing monomer and at least one primary hydroxyl group-containing monomer to a suitable reactor to which at least one glycidyl ester of an aliphatic saturated monocarboxylic acid and at least one polyester polyl are added. A suitable free radical polymerization initiator is also added. In an embodiment of the present invention, the polymerization is run in the substantial absence of a solvent.
Any suitable free radical initiator may be used in the polymerization. Suitable free radical initiators include, but are not limited to, thermal initiators, photoinitiators and oxidation-reduction initiators. Examples of thermal initiators include, but are not limited to, azo compounds, peroxides and persulfates. Suitable persulfates include, but are not limited to, sodium persulfate and ammonium persulfate.

Oxidation-reduction initiators may include, as non-limiting examples perulfate-bisulfite systems as well as systems utilizing thermal initiators in combination with appropriate metal ions such as iron or copper.

Suitable azo compounds include, but are not limited to, non-water-soluble azo compounds such as 1-1'-azobis(cyclohexancarbonitrile), 2-2'-azobisisobutyronitrile, 2-2'-azobis(2-methylbutyronitrile), 2-2'-azobis(propionitrile), 2-2'-azobis(2,4-dimethylvaleronitrile), 2-2'-azobis(valeronitrile), 2-(carbamoylazo)-isobutyronitrile and mixtures thereof; and water-soluble azo compounds such as azobis tertiary alkyl compounds include, but are not limited to,

4-4'-azobis(4-cyanovlaric acid), 2-2'-azobis(2-methylpropionamidine) dihydrochloride, 2,2'-azobis[2-methyl-N-(2-hydroxyethyl)propionamide], 4,4'-azobis(4-cyanopentanoic acid), 2,2'-azobis(N,N'-dimethyleneisobutyramidine), 2,2'-azobis(2-amidinopropane) dihydrochloride, 2,2'-azobis(N,N'-dimethyleneisobutyramidine) dihydrochloride and mixtures thereof.

Suitable peroxides include, but are not limited to, hydrogen peroxide, methyl ethyl ketone peroxides, benzoyl peroxides, di-t-butyl peroxides, di-t-amyl peroxides, dicumyl peroxides, diacyl peroxides, decanoyl peroxide, lauroyl peroxide, peroxycarbonates, peroxysterers, dialkyl peroxides, hydroperoxides, peroxycetals and mixtures thereof.

Any suitable carboxylic acid group-containing monomer may be used to prepare the acrylic polyol used in the compositions of the present invention, so long as it can be polymerized under the conditions described above. Examples of suitable carboxylic acid group-containing monomers include, but are not limited to, (meth)acrylic acid, maleic acid and its corresponding anhydride, itaconic acid, aconitic acid, fumaric acid, alpha-halo acrylic acid, vinyl acetic acid and beta-carboxymethyl (meth)acrylate.
Any suitable monomer containing a primary hydroxyl group may be used to prepare the acrylic polyol used in the compositions of the present invention, so long as it can be polymerized under the conditions described above. Examples of suitable monomers containing a primary hydroxyl groups include, but are not limited to, hydroxyethylacrylate, hydroxyethylmethacrylate, hydroxypropyl acrylate, hydroxypropylmethacrylate, hydroxybutyl (meth)acrylate, glycerol allyl ether, ethylene oxide esters of (meth)acrylic acid and propylene oxide esters of (meth)acrylic acid.

Other suitable polymerizable ethylenically unsaturated monomers may be used to prepare the acrylic polyol of the present invention. Other suitable monomers include, but are not limited to, C₁₋₃₀ aliphatic alkyl esters of (meth)acrylic acid, non-limiting examples of which include methyl(meth)acrylate, ethyl(meth)acrylate, propyl(meth)acrylate, N-butyl(meth)acrylate, t-butyl(meth)acrylate, 2-ethylhexyl(meth)acrylate, isobornyl (meth)acrylate, glycidyl (meth)acrylate, dimethylaminoethyl (meth)acrylate, N-butoxy methyl (meth)acrylamide, lauryl (meth)acrylate, cyclohexyl (meth)acrylate and 3,3,5-trimethylcyclohexyl (meth)acrylate. Other non-limiting examples of suitable monomers include (meth)acrylamide, N,N dialkyl (meth)acrylamides dimethylaminoethyl (meth)acrylate, vinyl aromatic compounds such as styrene and vinyl toluene, nitriles such as (meth)acrylonitrile, vinyl and vinylidene halides such as vinyl chloride and vinylidene fluoride and vinyl esters such as vinyl acetate.

Carbamate functional groups can be included in the acrylic polyol polymer by copolymerizing the acrylic monomers with a carbamate functional vinyl monomer, such as a carbamate functional alkyl ester of methacrylic acid, or by reacting a hydroxyl functional acrylic polymer with a low molecular weight carbamate functional material, such as can be derived from an alcohol or glycol ether, via a transcarbamoylation reaction. Other useful carbamate functional monomers are disclosed in U.S. Patent No. 5,098,947 to Metzger et al., which is incorporated herein by reference.

While not being bound to a single theory, it is believed that forming the acrylic polyol in the presence of the polyester polyol results in the formation of a stable
polymer-polymer complex. The stability of the polymer-polymer complex may arise from the association of hydrophobic portions of the polyester polyl and acrylic polyl polymer backbones. On the other hand, a portion of the formed acrylic polyl may graft onto the polyester polyl backbone. Additionally, the acrylic polyl and polyester polyl may become inseparably commingled and entangled. Combinations of associations may exist to the point that an interpenetrating polymer network is formed between the polyester polyl and acrylic polyl. Regardless of the method or reason, the combination of the acrylic polyl and polyester polyl demonstrates superior storage stability when compared to prior art physical blends or mixtures of such materials.

In order to facilitate the formation of an interpenetrating polymer network structure, one or more crosslinking monomers may be utilized in the monomer mixture. The amount of crosslinking monomer used will depend on the degree of branching or crosslinking desired. If mild branching is desired, a relatively low level of crosslinking monomer will be used, whereas if a highly crosslinked acrylic polyl is desired a higher level of crosslinking monomer will be used. Suitable crosslinking monomers include compounds having two or more functional groups that will react with a free radical. As a non-limiting example, compounds having two or more reactive unsaturated groups may be used. Suitable crosslinking monomers that may be used in the monomer mix include, but are not limited to, ethylene glycol di(meth)acrylate, triethylene glycol di(meth)acrylate, tetraethylene glycol di(meth)acrylate, 1,3-butylene glycol di(meth)acrylate, trimethylolpropane tri(meth)acrylate, 1,4-butanediol di(meth)acrylate, neopentyl glycol di(meth)acrylate, 1,6-hexanediol di(meth)acrylate, pentaerythritol di(meth)acrylate, pentaerythritol tri(meth)acrylate, pentaerythritol tetra(meth)acrylate, glycerol di(meth)acrylate, glycerol allyloxy di(meth)acrylate, 1,1,1-tris(hydroxymethyl)ethane di(meth)acrylate, 1,1,1-tris(hydroxymethyl)ethane tri(meth)acrylate, 1,1,1-tris(hydroxymethyl)propane di(meth)acrylate, 1,1,1-tris(hydroxymethyl)propane tri(meth)acrylate, triallyl cyanurate, triallyl isocyanurate, triallyl trimellitate, diallyl phthalate, diallyl terephthalate, divinyl benzene, methylol (meth)acrylamide, triallylamine and methylenebis (meth)acrylamide.
In the polymerization method described above, the equivalent ratio of the carboxylic acid groups from the carboxylic acid group-containing monomer to the epoxy group of the glycidyl ester of an aliphatic saturated monocarboxylic acid may be greater than 1:1, in some cases at least 1.01:1, in other cases at least 1.1:1, in other instances 1.2:1 and in other cases at least 1.25:1. In a further embodiment, the carboxylic acid groups from the carboxylic acid group-containing monomer are present in excess of the epoxy group of the glycidyl ester of an aliphatic saturated monocarboxylic acid. Generally, the carboxylic acid functional groups of the carboxylic acid functional monomers react with the epoxy group of the glycidyl ester of an aliphatic saturated monocarboxylic acid, resulting in the formation of the corresponding ester group and a secondary hydroxyl group.

Any suitable glycidyl ester of an aliphatic saturated monocarboxylic acid may be used. In an embodiment of the present invention, the glycidyl ester of an aliphatic saturated monocarboxylic acid is a glycidyl ester having structure (I).

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\begin{array}{c}
\text{I} \\
R^1 \quad R^2 \quad O \quad C \quad C \quad O \quad CH_2 \quad CH_2 \quad CH_2 \\
R^3
\end{array}
\]

In structure (I), R\(^1\) is C\(_{1-18}\) linear or branched alkyl and R\(^2\) and R\(^3\) are independently selected from H and C\(_{1-18}\) linear or branched alkyl. Non-limiting examples of suitable glycidyl esters of carboxylic acids include VERSATIC ACID 911 and CARDURA E, each of which is commercially available from Resolution Performance Products.

The acrylic polyol used in the present invention may be of any suitable molecular weight. The molecular weight of the acrylic polyol may be at least 250, in some cases at least 500, in other cases at least 750, in some instances at least 1,000 and in other instances at least 1,500. Additionally, the molecular weight of the acrylic polyol may be up to 25,000, in some cases up to 15,000, in other cases up to 10,000, in some instances up to 7,500 and in other instances up to 5,000 as determined by
GPC using polystyrene standards. The molecular weight of the acrylic polyol is selected based on the flow properties desired in component A and the properties desired in the coating film resulting from the present one-component, waterborne film-forming composition. The molecular weight of the acrylic polyol may be any value or any range of values inclusive of those stated above.

As described above, the carboxylic acid functional monomer may be present in excess compared to the glycidyl ester of an aliphatic saturated monocarboxylic acid. Thus, the acrylic polyol may contain carboxylic acid functionality resulting from residual or unreacted carboxylic acid groups. The acrylic polyol may have an acid value of at least 1, in some cases at least 2, in other cases at least 5, in some instances at least 10 and in other instances at least 25 mg KOH/g resin. Additionally, the acrylic polyol may have an acid value of not more than 100, in some cases not more than 75, in other cases not more than 50, in some instances not more than 40 and in other instances not more than 35 mg KOH/g resin. The acid value (number of milligrams of KOH per gram of solid required to neutralize the acid functionality in the resin) is a measure of the amount of acid functionality in the resin. The acid value of the acrylic polyol may be any value or any range of values inclusive of those stated above.

Any suitable polyester polyol may be used in the present invention. As a non-limiting example, useful polyester polymers typically include the condensation products of polyhydric alcohols and polycarboxylic acids. Suitable polyhydric alcohols can include ethylene glycol, neopentyl glycol, trimethylol propane and pentaerythritol. Suitable polycarboxylic acids can include adipic acid, 1,4-cyclohexyl dicarboxylic acid and hexahydrophthalic acid. Besides the polycarboxylic acids mentioned above, functional equivalents of the acids such as anhydrides where they exist or lower alkyl esters of the acids such as the methyl esters can be used. Also, small amounts of monocarboxylic acids such as stearic acid can be used. The ratio of reactants and reaction conditions are selected to result in a polyester polymer with the desired pendent functionality, i.e., carboxyl or hydroxyl functionality.

As a non-limiting example, hydroxyl group-containing polyesters can be prepared by reacting an anhydride of a dicarboxylic acid such as hexahydrophthalic
anhydride with a diol such as neopentyl glycol in a 1:2 molar ratio. Where it is
desired to enhance air-drying, suitable drying oil fatty acids may be used and include
those derived from linseed oil, soya bean oil, tall oil, dehydrated castor oil or tung oil.

As a non-limiting example, the polyester polyol may be prepared by reacting
one or more polyepoxides with one or more polycarboxylic acids. Useful
polyepoxides contain at least two epoxy groups with the diepoxides being most
preferred. Among the preferred diepoxides useful in the preparation of such a
polyester polyol are diglycidyl ether of bisphenol A and butyl diglycidyl ether.
Useful polycarboxylic acids may be selected from aliphatic, cycloaliphatic, and
aromatic polycarboxylic acids and mixtures thereof, with those containing no
ethylenic unsaturation and bearing no hydroxy functionality being most preferred.
Exemplary of the many acids which may be employed are phthalic acid, isophthalic
acid, terephthalic acid, oxalic acid, malonic acid, succinic acid, glutaric acid, adipic
acid, pimelic acid, suberic acid, 1,4-napthalenedicarboxylic acid,
2,3-napthalenedicarboxylic acid, 2,6-napthalenedicarboxylic acid and the like.
Additionally, monocarboxylic acids may be included such as benzoic acid, t-butyl
benzoic acid and acetic acid.

It will be appreciated that various combinations of carboxylic acids and
epoxides within the scope of the invention other than those specifically discussed
above may be reacted in order to provide polyester polyols useful in the present
one-component, waterborne film-forming composition. For example, oligomers
bearing hydroxy functionality other than that produced by the esterification reaction
may be produced by reacting not just a hydroxy bearing carboxylic acid with an
epoxide as discussed above, but by reacting carboxylic acids and epoxides, either of
which bears hydroxy functionality.

The polyester polyol used in the present invention may be of any suitable
molecular weight. The molecular weight of the polyester polyol may be at least 250,
in some cases at least 500, in other cases at least 1,000, in some instances at least 750
and in other instances at least 1,000. Additionally, the molecular weight of the
polyester polyol may be up to 25,000, in some cases up to 15,000, in other cases up to
10,000, in some instances up to 5,000 and in other instances up to 3,000 as determined
by GPC using polystyrene standards. The molecular weight of the polyester polyol is selected based on the flow properties desired in component (A) and the properties desired in the coating film resulting from the present one-component, waterborne film-forming composition. The molecular weight of the polyester polyol may be any value or any range of values inclusive of those stated above.

The polymer-polymer complex of component (A) may have a hydroxyl equivalent weight of at least 200, in some cases at least 300, in other cases at least 400, in some instances at least 500 and in other instances at least 1,000 grams per equivalent. Additionally, the polymer-polymer complex of component (A) may have a hydroxyl equivalent weight of not more than 5,000, in some cases not more than 4,000, in other cases not more than 3,500, in some instances not more than 3,000 and in other instances not more than 2,000 grams per equivalent. The term “equivalent weight” is a calculated value based on the relative amounts of the various ingredients used in making the specified material and is based on the solids of the specified material. The relative amounts are those that result in the theoretical weight in grams of the material, such as a polymer produced from the ingredients, and give a theoretical number of the particular functional group that is present in the resulting polymer. The theoretical polymer weight is divided by the theoretical number to give the equivalent weight. For example, hydroxyl equivalent weight is based on the equivalents of reactive pendant and/or terminal hydroxyl groups in the hydroxyl-containing polymer.

As indicated above, in addition to (A), the present one-component, waterborne film-forming composition further includes a crosslinking agent (B) having functional groups that are reactive with the hydroxyl groups in (A). The crosslinking agents typically are blocked polyisocyanates and/or aminoplasts. By "blocked" or “capped”, is meant is that the isocyanate groups have been reversibly reacted with a compound so that the resultant blocked isocyanate group is stable to active hydrogens at ambient temperature but reactive with active hydrogens in the film-forming polymer at elevated temperatures, for example, at temperatures between 30°C and 200°C.

In an embodiment of the present invention, the blocked or capped isocyanate of (B) includes a compound comprising a blocking group and a polyisocyanate
compound. When a blocked or capped polyisocyanate is used, it may be blocked (or capped) using any suitable aliphatic, cycloaliphatic or aromatic alkyl monoalcohol phenol, or with another alcohol, or a beta diol known to those skilled in the art that can be used as a blocking agent for the polyisocyanate. Other suitable blocking agents include oximes and lactams. When used, the blocked polyisocyanate typically is present, when added to the other components which form the thermosetting film-forming composition of the present invention, in an amount ranging from 5 to 65 weight percent, and can be present in an amount ranging from 10 to 45 weight percent, and often is present in an amount ranging from 15 to 40 percent by weight based on the total weight of resin solids present in the film-forming composition.

Other blocked isocyanate compounds that may be used in the present invention include the tricarbamoyl triazine compounds described in detail in U.S. Patent No. 5,084,541, which is incorporated by reference herein. When used, such blocked isocyanates can be present, when added to the thermosetting composition, in an amount ranging up to 20 weight percent, and can be present in an amount ranging from 1 to 20 weight percent, based on the total weight of resin solids present in the film-forming composition.

The polyisocyanates can be fully blocked as described in U.S. Patent No. 3,984,299 column 1 lines 1 to 68, column 2 and column 3 lines 1 to 15, or partially blocked and reacted with the polymer backbone as described in U.S. Patent No. 3,947,338 column 2 lines 65 to 68, column 3 and column 4 lines 1 to 30, which are incorporated by reference herein.

Any suitable aliphatic, cycloaliphatic, or aromatic alkyl monoalcohol or phenolic compound may be used as a capping agent for the capped polyisocyanate curing agent in the coating composition of the present invention including, for example, lower aliphatic alcohols such as methanol, ethanol, and n-butanol; cycloaliphatic alcohols such as cyclohexanol; aromatic-alkyl alcohols such as phenyl carbinol and methylphenyl carbinol; and phenolic compounds such as phenol itself and substituted phenols wherein the substituents do not affect coating operations, such as cresol and nitrophenol.
Glycol ethers may also be used as capping agents. Suitable glycol ethers include ethylene glycol butyl ether, diethylene glycol butyl ether, ethylene glycol methyl ether and propylene glycol methyl ether.

Other suitable capping agents include oximes such as methyl ethyl ketoxime, acetone oxime and cyclohexanone oxime, lactams such as epsilon-caprolactam, and amines such as dibutyl amine.

In the present invention, the polyisocyanate of the blocked polyisocyanate may be one or more suitable polyisocyanates. Suitable polyisocyanates include, but are not limited to, 1,4-tetramethylene diisocyanate, 1,6-hexamethylene diisocyanate, isophorone diisocyanate, 4,4'-methylene-bis-(cyclohexyl isocyanate), p-phenylene diisocyanate, diphenylmethane-4,4'-diisocyanate, 2,4-toluene diisocyanate, 2,6-toluene diisocyanate, triphenylmethane-4,4',4''-triisocyanate, 1,2,4-benzene triisocyanate, polymethylene polyphenyl isocyanate, the isocyanurate of hexamethylene diisocyanate, the biuret of hexamethylene diisocyanate, the isocyanurate of isophorone diisocyanate meta- α,α,α',α'-tetramethylxlylenediisocyanate, and para- α,α,α',α'-tetramethylxlylenediisocyanate.

When the blocking or capping agent is a lower aliphatic alcohol, nonlimiting examples include methanol, ethanol, and n-butanol. When the blocking or capping agent is a cycloaliphatic alcohol, a nonlimiting example is cyclohexanol. When the blocking or capping agent is an aromatic-alkyl alcohol, nonlimiting examples include phenyl carbinol and methylphenyl carbinol. When the blocking or capping agent is a phenolic compound, non-limiting examples include phenol, substituted phenols, cresol and nitrophenol. When the blocking or capping agent is an oxime, non-limiting examples include methyl ethyl ketoxime, acetone oxime and cyclohexanone oxime.

When the blocking or capping agent is a lactam, a non-limiting example is epsilon-caprolactam. A non-limiting example of an amine blocking or capping agent is dibutyl amine.

Suitable aminoplasts for use as the crosslinking agent (B) may include the condensation products obtained from the reaction of alcohols and formaldehyde with one or more amines. In this embodiment, the condensation products obtained from the reaction of alcohols and formaldehyde with melamine, urea or benzoguanamine is
most commonly used. While the aldehyde employed is most often formaldehyde, other similar condensation products can be made from other aldehydes, such as acetaldehyde, crotonaldehyde, acrolein, benzaldehyde, furfural, glyoxal and the like.

Condensation products of other amines and amides can also be used, for example, aldehyde condensates of triazines, diazines, triazoles, guanadines, guanamines and alkyl- and aryl-substituted derivatives of such compounds, including alkyl- and aryl-substituted ureas and alkyl- and aryl-substituted melamines. Non-limiting examples of such compounds include N,N'-dimethyl urea, benzourea, dicyandiamide, formaguanamine, acetoguanamine, glycoluril, ammeline, 3,5-diaminotriazole, triaminopyrimidine, 2-mercapto-4,6-diaminopyrimidine and carbamoyl triazines of the formula C₃N₅(NHCOXR), where X is nitrogen, oxygen or carbon and R is a lower alkyl group having from one to twelve carbon atoms or mixtures of lower alkyl groups, such as methyl, ethyl, propyl, butyl, n-octyl and 2-ethylhexyl. Such compounds and their preparation are described in detail in U.S. Patent No. 5,084,541, which is hereby incorporated by reference.

The aminoplast resins can contain methylol or similar alkylol groups, and in most instances at least a portion of these alkylol groups are etherified by reaction with an alcohol. Any monohydric alcohol can be employed for this purpose, including methanol, ethanol, propanol, butanol, pentanol, hexanol or heptanol, as well as benzyl alcohol and other aromatic alcohols, cyclic alcohols such as cyclohexanol, monoethers of glycols, and halogen-substituted or other substituted alcohols such as 3-chloropropanol and butoxyethanol.

In the present film-forming composition, (A) and (B) can react to effect cure at suitable temperatures when the one-component, waterborne film-forming composition of the present invention is applied to a substrate. The film-forming composition may cure at 60°C; on the other hand, it may cure at 70°C or above; if required it may cure at 80°C or above; in some cases it will be required to cure at 100°C or above and in other situations at 100°C or above. When the crosslinking agent of (B) is a blocked polyisocyanate, the cure temperature will typically be at least 100°C. Additionally, the film-forming composition may cure at up to 200°C; on the other hand, it may cure at up to 190°C; if required it may cure at up to 180°C; in some cases it will be
required to cure at up to 170°C and in other situations at up to 160°C. The cure
temperature of the film-forming composition may be any value or any range of values
inclusive of those stated above.

Generally, component (A) is present in the film-forming composition of the
present invention in an amount of at least 25 weight percent, in some cases at least 30
weight percent, in other cases at least 35 weight percent, in some instances at least 40
weight percent and in other instances at least 50 weight percent based on total resin
solids of the coating composition. Additionally, component (A) is present in the film-
forming composition in an amount of up to 99 weight percent, in some cases up to 95
weight percent, in other cases up to 90 weight percent, in some instances up to 80
weight percent and in other instances up to 70 weight percent based on total resin
solids of the coating composition. The amount of component (A) is determined by
variables such as the properties desired in the final coated film and the speed of cure.
The amount of component (A) present in the film-forming composition may be any
value or any range of values inclusive of those stated above.

Generally, component (B) is present in the film-forming composition of the
present invention in an amount of at least 1 weight percent, in some cases at least 5
weight percent, in other cases at least 10 weight percent, in some instances at least 20
weight percent and in other instances at least 30 weight percent based on total resin
solids of the coating composition. Additionally, component (B) may be present in the
film-forming composition in an amount of up to 75 weight percent, in some cases up
to 70 weight percent, in other cases up to 65 weight percent, in some instances up to
60 weight percent and in other instances up to 50 weight percent based on total resin
solids of the coating composition. The amount of component (B) is determined by
variables such as the properties desired in the final film and the speed of cure. The
amount of component (B) present in the film-forming composition may be any value
or any range of values inclusive of those stated above.

If desired, the present film-forming composition can comprise other optional
materials well known in the art of formulated surface coatings, such as surfactants,
flow control agents, thixotropic agents such as bentonite clay, fillers, organic
cosolvents, catalysts, including phosphonic acids and other customary auxiliaries.
These materials can constitute up to 40 percent by weight of the total weight of the coating composition.

The solids content of the present film-forming composition may be at least 20 weight percent, in some cases at least 25 weight percent, in other cases at least 30 weight percent, in some instances at least 35 weight percent and in other instances at least 40 weight percent based on the total weight of the film-forming composition. Additionally, the solids content of the present film-forming composition may be up to 75 weight percent, in some cases up to 70 weight percent, in other cases up to 65 weight percent, in some instances up to 60 weight percent and in other instances up to 50 weight percent based on the total weight of the film-forming composition. The solids content of the film-forming composition may be any value or any range of values inclusive of those stated above.

Any of the film-forming compositions previously described can be used advantageously to form a clear topcoat in a multi-component composite coating composition, such as a color-plus-clear composite coating. A color-plus-clear composite coating typically comprises a base coat deposited from a pigmented or colored film-forming composition, and a transparent or clear topcoat applied over the base coat. A further embodiment of the present invention relates to such a multi-layer composite coating. The multi-layer composite coating includes a base coat layer deposited from a pigmented film-forming base coat composition; and a substantially pigment free topcoat deposited over at least a portion of the base coat layer from a topcoat composition, where the topcoat composition comprises the present one-component, waterborne film-forming composition.

The multi-component composite coating of the present invention can be applied to various substrates to which they adhere, including wood, metals, glass, cloth, polymeric substrates and the like. They are particularly useful for coating metals and elastomeric substrates that are found on motor vehicles. The compositions can be applied by conventional means including brushing, dipping, flow coating, spraying and the like, but they are most often applied by spraying. The usual spray techniques and equipment for air spraying and electrostatic spraying and either manual or automatic methods can be used. During application of the coating
composition to the substrate, ambient relative humidity can range from about 30 to about 60 percent. The coating composition of the present invention is particularly advantageous when applied at an ambient relative humidity ranging from about 40 to about 60 percent, yielding very smooth finishes.

Another embodiment of the present invention is directed to a coated substrate that includes a substrate and the multi-layer composite coating composition described above over at least a portion of the substrate. Any suitable substrate may be used. As non-limiting examples, suitable substrates can include a metallic substrate or an elastomeric substrate.

When preparing the present multi-layer composite coating, a base coat coating composition is first applied to the surface of the substrate to be coated to form a base coat layer thereon. The base coat coating composition can be waterborne, solventborne or powdered, and typically includes a film-forming resin, crosslinking material (such as are discussed above) and pigment. Non-limiting examples of suitable base coat coating compositions include waterborne base coats for color-plus-nuclear composites such as are disclosed in U.S. Patent Numbers 4,403,003; 4,147,679; and 5,071,904, each of which is incorporated by reference herein.

After application of the base coating to the substrate, a film is formed on the surface of the substrate by driving water out of the film by heating or by an air-drying period. Typically, the coating thickness ranges from about 0.1 to about 5 mils (about 2.54 to about 127 microns), and preferably about 0.4 to about 1.5 mils (about 10.16 to about 38.1 microns) in thickness.

The heating will preferably be only for a short period of time and will be sufficient to ensure that the topcoat can be applied to the base coat if desired without the former dissolving the base coat composition. Suitable drying conditions will depend on the particular base coat composition and on the ambient humidity, but in general a drying time of from about 1 to 5 minutes at a temperature of about 80-250°F (20-121°C) will be adequate to ensure that mixing of the two coats is minimized. At the same time, the base coat film is adequately wetted by the topcoat composition so that satisfactory intercoat adhesion is obtained. Also, more than one base coat and multiple topcoats may be applied to develop the optimum appearance. Usually
between coats, the previously applied coat is flashed; that is, exposed to ambient conditions for about 1 to 20 minutes.

After application of the base coat in a composition, a topcoat can be formed by depositing any of the film-forming compositions described in detail above thereover.

Preferably, the topcoat coating composition is chemically different or contains different relative amounts of ingredients from the base coat coating composition.

The film-forming (topcoat) coating composition can be applied to the surface of the base coat by any of the coating processes discussed above for applying the base coat coating composition to the substrate. The coated substrate may be heated to cure the coating layers. In the curing operation, solvents are driven off and the film-forming materials of the clearcoat and the base coat are each crosslinked. The heating or curing operation may be carried out at 60°C or above; on the other hand, it may be carried out at 70°C or above; if required it may be carried out at 80°C or above; in some cases it will be required to be carried out at 100°C or above and in other situations at 120°C or above. When the crosslinking agent is a blocked polyisocyanate, the curing operation is typically carried out at a temperature of at least 100°C. Additionally, the heating or curing operation of the film-forming materials of the clearcoat and the base coat may be carried out at up to 200°C; on the other hand, it may be carried out at up to 190°C; if required it may be carried out at up to 180°C; in some cases it will be required to be carried out at up to 170°C and in other situations at up to 160°C. The heating or curing operation may be carried out at any temperature or any range of temperatures inclusive of those stated above.

The thickness of the topcoat may be at least 0.25 mils (6.3 microns), in some cases at least 0.5 mils (12.7 microns), in other cases at least 0.75 mils (19.1 microns), in some instances at least 1.0 mils (25.4 microns) and in other instances at least 1.25 mils (31.7 microns). Additionally, the thickness of the clearcoat may be up to 7 mils (177.8 microns), in some cases up to 5 mils (127 microns), in other cases up to 4 mils (101.6 microns), in some instances up to 3 mils (76.2 microns) and in other instances up to 2.5 mils (63.5 microns). The thickness of the topcoat may be any thickness or any range of thicknesses inclusive of those stated above.
In certain embodiments of the present invention, the crosslink density of the cured film-forming composition, i.e., the degree of crosslinking, ranges from 5% to 100% of complete crosslinking. In other embodiments, the crosslink density ranges from 35% to 85% of full crosslinking. In other embodiments, the crosslink density ranges from 50% to 85% of full crosslinking. One skilled in the art will understand that the presence and degree of crosslinking, i.e., the crosslink density, can be determined by a variety of methods, such as dynamic mechanical thermal analysis (DMTA) using a Polymer Laboratories MK III DMTA analyzer conducted under nitrogen. This method determines the glass transition temperature and crosslink density of free films of coatings or polymers. These physical properties of a cured material are related to the structure of the crosslinked network.

According to this method, the length, width and thickness of a sample to be analyzed are first measured, the sample is tightly mounted to the Polymer Laboratories MK III apparatus, and the dimensional measurements are entered into the apparatus. A thermal scan is run at a heating rate of 3°C/min, a frequency of 1 Hz, a strain of 120%, and a static force of 0.01N, and sample measurements occur every two seconds. The mode of deformation, glass transition temperature and crosslink density of the sample can be determined according to this method. Higher crosslink density values indicate a higher degree of crosslinking in the coating.

The one-component, waterborne film-forming composition of the present invention, when used as a coating composition, demonstrates excellent gloss, film hardness, solvent and acid resistance while containing minimal volatile organic solvents.

The present invention will further be described by reference to the following examples. The following examples are merely illustrative of the invention and are not intended to be limiting. Unless otherwise indicated, all percentages are by weight.

Example 1:

The ingredients listed below were used to prepare a polyester polyol.

<table>
<thead>
<tr>
<th>Component</th>
<th>Weight (grams)</th>
</tr>
</thead>
</table>

1,4-Cyclohexane & 344.0
Dicarboxylic Acid & 
Isostearic Acid & 568.0
Trimethylol propane & 540.0
Triphenyl phosphite & 3.6
Dibutyltin Oxide & 2.6

The components were charged into a three-liter, four-necked round bottom flask equipped with a motor driven stainless steel paddle agitator, a thermocouple to record batch temperature, a Dean-Stark water trap connected with a condenser to collect distillate evolved and a nitrogen sparge tube.

The synthesis was performed using azeotropic conditions with xylene (3% on solids). Heat was applied to a Glas-Col heating mantle and the temperature was gradually increased to about 220°C and held until an acid value of less than four, as measured by potentiometric titration with KOH, was obtained.

Example 2:

This example demonstrates the synthesis of the stable aqueous dispersion of a combination of a polyester polyol and a carboxylic acid-containing acrylic polymer polyol of the present invention. The synthesis used the ingredients shown below:

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Amount (grams)</th>
<th>Resin Solids (weight %)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Charge 1</strong></td>
<td>Glycidyl ester of branched decanoic acid</td>
<td>961.9</td>
</tr>
<tr>
<td><strong>Charge 2</strong></td>
<td>di-t-amyl peroxide</td>
<td>62.7</td>
</tr>
<tr>
<td></td>
<td>m-styrene dimer</td>
<td>91.2</td>
</tr>
<tr>
<td><strong>Charge 3</strong></td>
<td>Butyl acrylate</td>
<td>326.5</td>
</tr>
<tr>
<td></td>
<td>Styrene</td>
<td>714.1</td>
</tr>
<tr>
<td></td>
<td>Hydroxyethyl acrylate</td>
<td>668.2</td>
</tr>
<tr>
<td></td>
<td>2-ethylhexyl acrylate</td>
<td>231.0</td>
</tr>
<tr>
<td></td>
<td>Acrylic acid</td>
<td>363.6</td>
</tr>
<tr>
<td><strong>Charge 3A</strong></td>
<td>Polyester Polyol of Example 1</td>
<td>422.1</td>
</tr>
</tbody>
</table>
Charge 4
Dimethyl ethanol amine 120.0

Charge 5
Deionized water 5538.7

1Cardura™ Resin E-10, Resolution Performance Products, Houston, TX.

Charge 1 was added to a reaction vessel equipped with a reflux condenser and nitrogen blanket and heated to 160°C. Charge 3 and Charge 3A were mixed together and the mixture was added to the reaction vessel over a four-hour period. Beginning at the same time as the mixture, charge 2 was added to the reaction vessel over a 4.5-hour period, after which time the resulting product continued to be mixed in the vessel at 160°C. The product was cooled to 100°C and charge 4 was added over a 30-minute period of time. Charge 5 was pre-heated to 70°C and added to the vessel over a 30-minute period. The product was mixed for one hour at 100°C and decanted. The resulting product had total solids of 38.9 weight percent (1 hour at 110°C), pH of 8.4 and viscosity of 170cps (Brookfield, RVT spindle #1, 30 rpm at 22.3°C). The average particle size of the dispersed particles was 0.08 – 0.09 μm determined by Horiba LA-900 Laser Scattering Particle Size Distribution Analyzer, available from Horiba Instruments, Irvine, California.

The resulting latex did not demonstrate any visible sign of separation after standing for 4 months at ambient conditions.
Example 3:

This is a comparison example demonstrating the synthesis of an aqueous dispersion where a polyester polyol and a carboxylic acid-containing acrylic polymer polyol are physically mixed after synthesis of the acrylic polyol. An acrylic polyol was prepared using the ingredients shown below:

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Amount (grams)</th>
<th>Resin Solids (weight %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Charge 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glycidyl ester of branched decanoic acid&lt;sup&gt;1&lt;/sup&gt;</td>
<td>961.9</td>
<td>29.5</td>
</tr>
<tr>
<td>Charge 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>di-t-amyl peroxide</td>
<td>62.7</td>
<td></td>
</tr>
<tr>
<td>m-styrene dimer</td>
<td>91.2</td>
<td></td>
</tr>
<tr>
<td>Charge 3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Butyl acrylate</td>
<td>326.5</td>
<td>10.0</td>
</tr>
<tr>
<td>Styrene</td>
<td>714.1</td>
<td>21.9</td>
</tr>
<tr>
<td>Hydroxyethyl acrylate</td>
<td>668.2</td>
<td>20.5</td>
</tr>
<tr>
<td>2-ethylhexyl acrylate</td>
<td>231.0</td>
<td>7.07</td>
</tr>
<tr>
<td>Acrylic acid</td>
<td>363.6</td>
<td>11.1</td>
</tr>
<tr>
<td>Charge 4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dimethyl ethanol amine</td>
<td>120.0</td>
<td></td>
</tr>
<tr>
<td>Charge 5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deionized water</td>
<td>5538.7</td>
<td></td>
</tr>
</tbody>
</table>

<sup>1</sup> Cardura<sup>TM</sup> Resin E-10, Resolution Performance Products, Houston, TX.

Charge 1 was added to a reaction vessel equipped with a reflux condenser and nitrogen blanket and heated to 160°C. Charge 3 was added to the reaction vessel over a four-hour period. Beginning at the same time as the mixture, charge 2 was added to the reaction vessel over a 4.5-hour period, after which time the resulting product continued to be mixed in the vessel at 160°C. The product was cooled to 100°C and charge 4 was added over a 30-minute period of time. Charge 5 was pre-heated to 70°C and added to the vessel over a 30-minute period. The product was mixed for one hour at 100°C and decanted. The average particle size of the dispersed particles was 0.08 – 0.09 μm, determined by Horiba LA-900 Laser Scattering Particle Size.
Distribution Analyzer. The resulting latex did not demonstrate any visible sign of separation after standing 4 months at ambient conditions.

The acrylic polyl described above was physically mixed with the polyester polyl of example 1, at a solids ratio of 9:1 acrylic polyl to polyester polyl, using an overhead mixer for 20 minutes. The average particle size of the particles in the resulting dispersion was about 10 μm, determined by Horiba LA-900 Laser Scattering Particle Size Distribution Analyzer. The dispersion separated within a week into two layers, demonstrating its instability compared to the dispersion of the present invention shown in Example 2.

Examples 4 and 5:

A waterbased clearcoat (for Example 4) was prepared from the following ingredients:

<table>
<thead>
<tr>
<th>INGREDIENTS</th>
<th>AMOUNTS (grams)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Component I:</strong></td>
<td></td>
</tr>
<tr>
<td>Polymer of Example 2</td>
<td>71.5</td>
</tr>
<tr>
<td><strong>Component II:</strong></td>
<td></td>
</tr>
<tr>
<td>Cymel 327&lt;sup&gt;2&lt;/sup&gt;</td>
<td>8.72</td>
</tr>
<tr>
<td>DMP-Blocked HDI trimer&lt;sup&gt;3&lt;/sup&gt;</td>
<td>5.60</td>
</tr>
<tr>
<td>Tinuvin 1130&lt;sup&gt;4&lt;/sup&gt;</td>
<td>0.80</td>
</tr>
<tr>
<td>Tinuvin 292&lt;sup&gt;4&lt;/sup&gt;</td>
<td>0.50</td>
</tr>
<tr>
<td>Silicone surfactant&lt;sup&gt;5&lt;/sup&gt;</td>
<td>0.40</td>
</tr>
<tr>
<td>Silicone oil&lt;sup&gt;6&lt;/sup&gt;</td>
<td>0.02</td>
</tr>
<tr>
<td>High boiling alcohol&lt;sup&gt;7&lt;/sup&gt;</td>
<td>2.00</td>
</tr>
<tr>
<td><strong>Component III:</strong></td>
<td></td>
</tr>
<tr>
<td>Deionized water</td>
<td>10.46</td>
</tr>
</tbody>
</table>

<sup>2</sup>Melamine crosslinker, available from Cytec Industries, West Paterson, New Jersey.
<sup>4</sup>Available from Ciba Specialty Chemicals Corporation, Tarrytown, NY.
<sup>5</sup>Available as Byk 345 from the Byk Chemie division of Altana AG, Bad Homburg, Germany.
<sup>6</sup>Available as Byk 333 from Byk Chemie.
<sup>7</sup>Isolol 18T, available from Sasol Industries, Hamburg, Germany.

Waterbased clearcoat compositions were prepared from the components indicated above. Charge II was mixed separately under agitation for 1 hour at room
temperature. Charge II was added into charge I under agitation for 15 minutes and kept stirring for an additional 1 hour. The dispersion was then allowed to sit overnight. Charge III was added slowly under agitation to bring the dispersion to the spraying viscosity of 30 seconds efflux cup DIN 4 23°C.

Application:

The waterborne clearcoat described above (Example 4) and a commercial waterborne clearcoat, HYDROKLARLACK PPG 97020 available from PPG Industries, Inc., Pittsburgh, PA (Example 5), were applied in a humidity and temperature controlled spray booth at 60% relative humidity (“RH”) and 70°F (21°C) onto cold rolled steel substrates which had been previously electrocoated with ED5000 and primed with 1177225A gray primer (both products available commercially from PPG Industries, Inc.), the primed panels having been prepared by ACT Laboratories Inc., of Hillsdale MI. Each of the waterborne clearcoat formulations was spray-applied using the SATA LP90 gun with a MSB nozzle and 135 air cap. The coatings were applied over a commercial silver metallic waterborne basecoat available from PPG Industries, Inc. The basecoat was applied in two coats, with 60 second flash between coats and then prebaked for 10 minutes at 176°F (80°C). The clearcoat was then applied in two coats without any flash. The clearcoated panels were allowed to flash for five to ten minutes at ambient condition and baked for 10 minutes at 140°F (60°C) and finally, for 30 minutes at 293-302°F (145-150°C). Panels were baked in a horizontal position. The film build was approximately 2.0 mils.

Delamination test:

Deminerlized water was dripped onto test panels at 63°C for 72 hours. The test panels were then taken off of the water drip and dried with a clean cloth. Subsequently the surface was scratched using a Cross-Cut Tester (Byk-Chemie, Wesel, Germany) by cutting through the film to the substrate in one steady motion. A second cut was made, perpendicular to and centered on the first cut. Next, a lap of tape (Scotch Brand 800, 3M, St. Paul, Minnesota) was placed and rubbed firmly over
the cut. The tape was removed in a rapid upward motion. The tape and removal step was then repeated. Separation of the coating from the substrate should be no more than 1mm along the scratch with no separation of the coating on the remaining area.

5 Acid test 1:

A Gradient oven from Byk Gardner was used at a temperature range of 41-81°C. Several drops (each with a volume of 250μl) of a solution of 10% by weight sulfuric acid were applied to a preheated test panel at distance corresponding to the heating elements. Application of the acid solution was completed within 60 seconds, after which temperature stressing of the test panel for a period of 30 minutes was begun. The test panel was then removed from the gradient oven, then rinsed in deionized water and dried under a cold air stream. The evaluation was performed visually. The temperature at which the test panel showed the first signs of damage was recorded. The temperature at which the first visible signs of damage occur should be greater than 60°C after 30 minutes of storage time.

10 Acid test 2:

Several drops of 38% by weight sulfuric acid are applied to a test panel. No visible signs of damage should be seen before 72 hours at room temperature.

20 20° Gloss:

Specular gloss was measured at 20° with a Novo Gloss Statistical Glossmeter (Paul N. Gardner Company, Inc., Pompano Beach, FL) where higher numbers indicate better performance.

25 Hardness:

Hardness was measured using the Tukon Microhardness Instrument Model 300 (Wilson Instruments Division of Instron Corporation, Canton, MA). Higher numbers indicate better performance.
Results from the evaluations of the coated panels baked at 140°C (285°F) for gloss, indentation hardness, delamination, humidity resistance, adhesion and acid resistance are summarized in the table below.

<table>
<thead>
<tr>
<th></th>
<th>Example 4</th>
<th>Example 5 (Comparative)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gloss 20°</td>
<td>96.3</td>
<td>91.5</td>
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Example 4 included 13% solvent, while the conventional coating of Comparative Example 5 included 40% by weight solvent. These data show that the film-forming composition of the present invention (Example 4) provided better gloss than that of Comparative Example 5 made using a commercially available polymer, and in other respects provided a comparable coating, while containing significantly less volatile solvent.

Those skilled in the art will recognize that changes may be made to the embodiments described above without departing from the broad inventive concept thereof. It is understood, therefore, that this invention is not limited to the particular embodiments disclosed, but it is intended to cover modifications that are within the spirit and scope of the invention, as defined by the appended claims.
WE CLAIM:

1. A one-component, waterborne film-forming composition comprising:
   (A) a stable aqueous dispersion of a mixture of a polyester polyol
   and a carboxylic acid-containing acrylic polymer polyol, wherein the weight ratio of
   polyester polyol to acrylic polymer polyol is in the range of 10:90 to 30:70; and
   (B) a crosslinking agent having functional groups reactive with the
   hydroxyl groups in (A) selected from the group of blocked isocyanates and
   aminoplasts.

2. The film-forming composition of claim 1, wherein the crosslinking
   agent in (B) is a blocked polyisocyanate and the equivalent ratio of hydroxyl groups in
   (A) to isocyanate functional groups in (B) is from 0.8:1 to 1.7:1.

3. The film-forming composition of claim 1, wherein the crosslinking
   agent in (B) is an aminoplast and the aminoplast is present at from 10 to 60 percent by
   weight of the total resin solids of the film-forming composition.

4. The film-forming composition of claim 1, wherein component (A) is
   prepared by polymerizing a mixture of ethylenically unsaturated polymerizable
   monomers comprising:
   (1) at least one carboxylic acid group-containing monomer;
   (2) at least one primary alcohol group-containing monomer;
   (3) at least one glycidyl ester of an aliphatic saturated
   monocarboxylic acid; and
   (4) at least one polyester polyol.

5. The film-forming composition of claim 1, wherein the polyester polyol
   and the carboxylic acid-containing acrylic polymer polyol in (A) are present as a
   stable polymer-polymer complex.
6. The film-forming composition of claim 2, wherein after polymerization is complete, an amine is added to component (A) in an amount sufficient to provide a pH of from 7 to 10 when component (A) is dispersed in water.

7. The film-forming composition of claim 6, wherein the pH is increased by using one or more volatile amines.

8. The film-forming composition of claim 7, wherein the amines are selected from the group consisting of dimethylethanolamine, ammonia, triethyl amine and diethyl propanol amine.

9. The film forming composition of claim 4, wherein the equivalent ratio of the carboxylic acid groups from the carboxylic acid group-containing monomer to the epoxy group of the glycidyl ester of an aliphatic saturated monocarboxylic acid is greater than 1:1.

10. The film-forming composition of claim 4, wherein the polymerization comprises the steps of:

(a) introducing components (1) and (2) to a suitable reactor; and

(b) adding components (3) and (4) to the reactor, along with a suitable free radical polymerization initiator, in the substantial absence of a solvent.

11. The film-forming resin of claim 4, wherein the glycidyl ester (3) is a glycidyl ester having the following structure:

![Diagram](attachment:image.png)
wherein $R^1$ is $C_{1-18}$ linear or branched alkyl and $R^2$ and $R^3$ are independently selected from H and $C_{1-18}$ linear or branched alkyls.

12. The film-forming resin composition of claim 4, wherein the carboxylic acid group-containing monomer (1) is one or more selected from the group consisting of acrylic acid and (meth)acrylic acid and the monomer containing a primary alcohol group is one or more selected from the group consisting of hydroxyethylacrylate, hydroxyethylmethacrylate, hydroxypropylacrylate, hydroxypropylmethacrylate and hydroxybutyl (meth)acrylate.

13. The film-forming composition of claim 1, wherein (A) and (B) are curable at a temperature of from 60°C to 140°C.

14. The film-forming composition of claim 1, wherein the blocked isocyanate of (B) comprises a compound comprising a blocking group and a polyisocyanate compound.

15. The film-forming composition of claim 14, wherein the polyisocyanate is at least one selected from the group consisting of 1,4-tetramethylene diisocyanate, 1,6-hexamethylenediisocyanate, isophorone diisocyanate, 4,4'-methylene-bis-(cyclohexyl isocyanate), p-phenylene diisocyanate, diphenylmethane-4,4'-diisocyanate, 2,4-toluene diisocyanate, 2,6-toluene diisocyanate, triphenylmethane-4,4',4''-triisocyanate, 1,2,4-benzene triisocyanate, polymethylene polyphenyl isocyanate, the isocyanurate of hexamethylene diisocyanate, the biuret of hexamethylene diisocyanate, the isocyanurate of isophorone diisocyanate meta-$\alpha,\alpha',\alpha'$-tetramethylylenediisocyanate, and para-$\alpha,\alpha',\alpha'$-tetramethylylenediisocyanate.

16. The film-forming composition of claim 14, wherein the compound comprising a blocking group is at least one selected from the group consisting of aromatic alkyl monoalcohols, phenolic compounds, lower aliphatic alcohols,
cycloaliphatic alcohols, aromatic-alkyl alcohols; glycol ethers, amines, lactams and oximes.

17. The film-forming composition of claim 16, wherein the lower aliphatic alcohols are selected from the group consisting of methanol, ethanol, and n-butanol; the cycloaliphatic alcohols is cyclohexanol; the aromatic-alkyl alcohols are selected from the group consisting of phenyl carbinol and methylphenyl carbinol; the phenolic compounds are selected from the group consisting of phenol, substituted phenols, cresol and nitrophenol; the glycol ethers are selected from the group consisting of ethylene glycol butyl ether, diethylene glycol butyl ether, ethylene glycol methyl ether and propylene glycol methyl ether; the oximes are selected from the group consisting of methyl ethyl ketoxime, acetone oxime and cyclohexanone oxime, the lactam is epsilon-caprolactam, and the amine is dibutyl amine.

18. The film-forming composition of claim 1, wherein the aminoplast of (B) comprises the condensation products obtained from the reaction of alcohols and formaldehyde with one or more amines.

19. The film-forming composition of claim 18, wherein the amines are one or more selected from the group consisting of melamine, urea, benzoguanamine, triazines, diazines, triazoles, guanadines and guanamines.

20. The film-forming composition of claim 18, wherein the alcohols are at least one selected from the group consisting of aromatic alkyl monoalcohols, phenolic compounds, lower aliphatic alcohols, cycloaliphatic alcohols, aromatic-alkyl alcohols and glycol ethers.

21. The film-forming resin of claim 4, wherein the mixture of monomers further comprises (5) a crosslinking monomer.

22. A multi-layer composite coating comprising:
(I) a base coat layer deposited from a pigmented film-forming base coat composition; and

(II) a substantially pigment-free topcoat deposited over at least a portion of said base coat layer from a topcoat composition is a one-component, waterborne film-forming composition comprising:

(A) a stable aqueous dispersion of a mixture of a polyester polyol and a carboxylic acid-containing acrylic polymer polyol, wherein the weight ratio of polyester polyol to acrylic polymer polyol is in the range of 10:90 to 30:70; and

(B) a crosslinking agent having functional groups reactive with the hydroxyl groups in (A) selected from the group of blocked isocyanates and aminoplasts.

23. The multi-layer composite coating of claim 22, wherein the crosslinking agent in (B) is a blocked polyisocyanate and the equivalent ratio of hydroxyl groups in (A) to isocyanate functional groups in (B) is from 0.8:1 to 1.7:1.

24. The multi-layer composite coating of claim 22, wherein the crosslinking agent in (B) is an aminoplast and the aminoplast is present at from 10 to 60 percent by weight of the total resin solids of the film-forming composition.

25. The multi-layer composite coating of claim 22, wherein the first component (A) is prepared by polymerizing a mixture of ethylenically unsaturated polymerizable monomers comprising:

1. at least one carboxylic acid group-containing monomer;

2. at least one primary alcohol group-containing monomer;

3. at least one glycidyl ester of an aliphatic saturated monocarboxylic acid; and

4. at least one polyester polyol.
26. The multi-layer composite coating of claim 25, wherein the polyester polyl and the carboxylic acid-containing acrylic polymer polyl in (A) are present as a stable polymer-polymer complex.

27. The multi-layer composite coating of claim 25, wherein after polymerization is complete, an amine is added to component (A) in an amount sufficient to provide a pH of from 7 to 10 when component (A) is dispersed in water.

28. The multi-layer composite coating of claim 27, wherein the pH is increased by using one or more volatile amines.

29. The multi-layer composite coating of claim 28, wherein the amines are selected from the group consisting of dimethylethanolamine, ammonia, triethyl amine and diethyl propanol amine.

30. The multi-layer composite coating of claim 25, wherein the equivalent ratio of the carboxylic acid groups from the carboxylic acid group-containing monomer to the epoxy group of the glycidyl ester of an aliphatic saturated monocarboxylic acid is greater than 1:1.

31. The multi-layer composite coating of claim 25, wherein the polymerization comprises the steps of:
   (a) introducing components (1) and (2) to a suitable reactor; and
   (b) adding components (3) and (4) to the reactor, along with a suitable free radical polymerization initiator, in the substantial absence of a solvent.

32. The multi-layer composite coating of claim 25, wherein the glycidyl ester (3) is a glycidyl ester having the following structure:
wherein $R^1$ is $C_1$-$C_{18}$ linear or branched alkyl and $R^2$ and $R^3$ are independently selected from H and $C_1$-$C_{18}$ linear or branched alkyls.

33. The multi-layer composite coating of claim 25, wherein the carboxylic acid group-containing monomer (1) is one or more selected from the group consisting of acrylic acid and (meth)acrylic acid and the monomer containing a primary alcohol group is one or more selected from the group consisting of hydroxyethylacrylate, hydroxyethylmethacrylate, hydroxypropylacrylate, hydroxypropylmethacrylate and hydroxybutyl (meth)acrylate.

34. The multi-layer composite coating of claim 25, wherein (A) and (B) are curable at a temperature of from 60°C to 200°C.

35. The multi-layer composite coating of claim 25, wherein the blocked isocyanate of (B) comprises a compound comprising a blocking group and a polyisocyanate compound.

36. The multi-layer composite coating of claim 35, wherein the polyisocyanate is at least one selected from the group consisting of 1,4-tetramethylene diisocyanate, 1,6-hexamethylene diisocyanate, isophorone diisocyanate, 4,4'-methylene-bis-(cyclohexyl isocyanate), p-phenylene diisocyanate, diphenylmethane-4,4'-diisocyanate, 2,4-toluene diisocyanate, 2,6-toluene diisocyanate, triphenylmethane-4,4',4''-triisocyanate, 1,2,4-benzene triisocyanate, polymethylene polyphenyl isocyanate, the isocyanurate of hexamethylene diisocyanate, the biuret of hexamethylene diisocyanate, the isocyanurate of
isophorone diisocyanate meta-α,α,α',α'-tetramethylxylylenediisocyanate and
para-α,α,α',α'-tetramethylxylylenediisocyanate.

37. The multi-layer composite coating of claim 35, wherein the compound
comprising a blocking group is at least one selected from the group consisting of
aromatic alkyl monoalcohols, phenolic compounds, lower aliphatic alcohols,
cycloaliphatic alcohols, aromatic-alkyl alcohols, glycol ethers, amines, lactams and
oximes.

38. The multi-layer composite coating of claim 37, wherein the lower
aliphatic alcohols are selected from the group consisting of methanol, ethanol, and
n-butanol; the cycloaliphatic alcohols is cyclohexanol; the aromatic-alkyl alcohols are
selected from the group consisting of phenyl carbinol and methylphenyl carbinol; the
phenolic compounds are selected from the group consisting of phenol, substituted
phenols, cresol and nitrophenol; the glycol ethers are selected from the group
consisting of ethylene glycol butyl ether, diethylene glycol butyl ether, ethylene
glycol methyl ether and propylene glycol methyl ether; the oximes are selected from
the group consisting of methyl ethyl ketoxime, acetone oxime and cyclohexanone
oxime; the lactam is epsilon-caprolactam and the amine is dibutyl amine.

39. The multi-layer composite of claim 35, wherein the aminoplast of (B)
comprises the condensation products obtained from the reaction of alcohols and
formaldehyde with one or more amines.

40. The multi-layer composite of claim 39, wherein the amines are one or
more selected from the group consisting of melamine, urea, benzoguanamine,
triazines, diazines, triazoles, guanadines and guanamines.

41. The multi-layer composite of claim 39, wherein the alcohols are at
least one selected from the group consisting of aromatic alkyl monoalcohols, phenolic
compounds, lower aliphatic alcohols, cycloaliphatic alcohols, aromatic-alkyl alcohols and glycol ethers.

42. The multi-layer composite coating of claim 22, wherein the mixture of 5 monomers further comprises (5) a crosslinking monomer.

43. A coated substrate comprising:
   (A) a substrate, and
   (B) the multi-layer composite coating of claim 22 over at least a 10 portion of the substrate.

44. The coated substrate of claim 43, wherein the substrate is selected from a metallic substrate and an elastomeric substrate.

45. A coated substrate comprising:
   (A) a substrate, and
   (B) the one-component, waterborne film-forming composition of 15 claim 1 over at least a portion of the substrate.

46. The coated substrate of claim 45, wherein the substrate is selected from a metallic substrate and an elastomeric substrate.
**INTERNATIONAL SEARCH REPORT**

**PCT/US 03/15724**

**A. CLASSIFICATION OF SUBJECT MATTER**

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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)

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Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched.

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, PAJ, WPI Data

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

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**Further documents are listed in the continuation of box C.**

**Patent family members are listed in annex.**

* Special categories of cited documents:
  *A* document defining the general state of the art which is not considered to be of particular relevance.
  *E* earlier document published on or after the international filing date.
  *L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified).
  *O* document referring to an oral disclosure, use, exhibition or other means.
  *P* document published prior to the international filing date but later than the priority date claimed.

*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.

**Date of the actual completion of the international search**

25 July 2003

**Date of mailing of the international search report**

03/09/2003

**Name and mailing address of the ISA**

European Patent Office, P.B. 5818 Patentilaan 2 NL - 2280 HV Rijswijk

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**Authorized officer**

Lanz, S
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