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 (72) Inventeur/Inventor:
 SIEBERT, ELIZABETH J., US
 (73) Propriétaire/Owner:
 HENKEL CORPORATION, US
 (74) Agent: FETHERSTONHAUGH & CO.

(54) Titre : TRAITEMENT D'UN REVETEMENT AUTO-DEPOSE PAR ELECTROLYSE AU MOYEN D'UNE SOLUTION
 ALCALINE QUI CONTIENT DES ANIONS D'ACIDES ORGANIQUES MULTIFONCTIONNELS
 (54) Title: TREATING AN AUTODEPOSITED COATING WITH AN ALKALINE SOLUTION CONTAINING ANIONS OF
 MULTIFUNCTIONAL ORGANIC ACIDS

(57) **Abrégé/Abstract:**

The adhesion and/or corrosion resistance of an autodeposited coating can be improved by rinsing the uncured coating with an aqueous treatment solution that has a pH between 7 and 11 and contains between 0.05 and 5 w/o of anions derived from multifunctional organic acids, preferably anions of 1-hydroxyethylidene-1,1-diphosphonic acid, citric acid, tartaric acid, and/or oxalic acid. The method is particularly useful on leaf springs and other metallic objects with surfaces of high carbon and/or shot blasted steel, and is particularly useful in conjunction with an autodeposition bath containing internally stabilized poly (vinylidene chloride) latex, hydrofluoric acid, ferric fluoride, and hydrogen peroxide.





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The adhesion and/or corrosion resistance of an autodeposited coating can be improved by rinsing the uncured coating with an aqueous treatment solution that has a pH between 7 and 11 and contains between 0.05 and 5 w/o of anions derived from multifunctional organic acids, preferably anions of 1-hydroxyethylidene-1,1-diphosphonic acid, citric acid, tartaric acid, and/or oxalic acid. The method is particularly useful on leaf springs and other metallic objects with surfaces of high carbon and/or shot blasted steel, and is particularly useful in conjunction with an autodeposition bath containing internally stabilized poly (vinylidene chloride) latex, hydrofluoric acid, ferric fluoride, and hydrogen peroxide.

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**TREATING AN AUTODEPOSITED COATING WITH AN ALKALINE
SOLUTION CONTAINING ANIONS OF MULTIFUNCTIONAL ORGANIC
ACIDS**

BACKGROUND OF THE INVENTION

5 Field of the Invention

This invention relates to autodeposition. Autodeposition involves the use of an aqueous resinous coating composition of relatively low solids concentration (usually less than about 10 %) to form a coating of relatively high solids concentration (usually greater than about 10 %) on a metallic surface immersed therein, with the coating increasing in thickness and areal density (mass per unit area of coating) the longer the time the metallic surface is immersed in the composition. Autodeposition is somewhat similar to electrodeposition but does not require the aid of external electrical current to cause the resin particles to deposit on the metal surface.

15 In general, autodepositing compositions are aqueous acid solutions having solid resin particles dispersed

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therein in very finely divided form. The coating formed while the metal substrate used is immersed in the bath is generally wet and fairly weak, although sufficiently strong to maintain itself against gravity and moderate spraying forces. In this state the coating is described as "uncured". To make an autodeposition coated object suitable for normal practical use, the uncured coated is dried, usually with the aid of heat. The coating is then described as "cured".

The present invention relates more particularly to the chemical treatment of an uncured autodeposited coating for the purpose of improving various properties thereof, particularly the adhesion of the coating to the underlying metal substrate and the resistance to corrosion of the underlying metal provided by the cured autodeposited coating when the coated metal surfaced object is subjected to corrosive environments.

Statement of Related Art

Basic constituents of an autodepositing composition are water, resin solids dispersed in the aqueous medium of the composition, and activator, that is, an ingredient or ingredients which convert the composition into one which will form on a metallic surface a resinous coating which increases in thickness or areal density as long as the surface is immersed in the composition. Various types of activators or activating systems are known, for example, as reported in the following U. S. Patent Nos.: 3,592,699; 3,709,743; 4,103,049; 4,347,172; and 4,373,050. The activating system generally comprises an acidic oxidizing system, for example: hydrogen peroxide and HF; HNO₃; a ferric-containing compound and HF; and other soluble metal-containing compounds, for example, silver fluoride, ferrous oxide, cupric sulfate, cobaltous nitrate, silver acetate, ferrous phosphate, chromium fluoride, cadmium fluoride,

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stannous fluoride, lead dioxide, and silver nitrate in an amount between about 0.025 and about 50 grams per liter ("g/l") and an acid, which can be used alone or in combination with hydrofluoric acid, and including, for example, sulfuric, hydrochloric, nitric, and phosphoric acid, and organic acids, including, for example, acetic, chloroacetic, and trichloroacetic acids.

Previously known autodepositing compositions can be used to form coatings which have good aesthetic properties and which protect the underlying metallic substrate from being degraded (for example, corroded by water). However, there are certain applications which require that the autodeposited coating have particularly good properties for satisfactory use. Various means have been developed to improve the properties of autodeposited coatings, including, for example: chemical pretreatment of the metallic surface prior to formation of the coating; selection of particular resins for use in forming the coating; addition to the autodepositing composition of chemical additives; and chemical treatment of the freshly formed or uncured coating, as described in detail in United States Patent No. 5,342,694.

There are several U.S. patents which disclose the treatment of freshly formed autodeposited coatings with acidic aqueous solutions of one or more chromium compounds to improve the corrosion-resistance and/or surface appearance of the cured coating. Among such patents are Nos: 3,795,546; 4,030,945; 4,411,950; and 4,637,839, all assigned to the same assignee as that of the present invention. The '546 and '945 patents disclose treating an uncured autodeposited coating with an acidic aqueous solution containing hexavalent chromium or hexavalent chromium and formaldehyde-reduced forms of hexavalent chromium to improve the corrosion-resistant properties of the cured form of the coating and to reduce the gloss of

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an otherwise glossy coating. According to these patents, the source of chromium can be chromium trioxide or water-soluble salts of chromium or dichromate, for example, sodium, potassium, and lithium salts thereof. Optional ingredients of such chromium-containing solutions include phosphoric acid (anti-gelling agent), sodium hydroxide (pH adjuster), and a water-soluble or water-dispersible polyacrylic acid (corrosion-resistant and paint-bonder improver). The '950 patent discloses the treatment of an uncured autodeposited coating with an aqueous chromium-containing solution which has dispersed therein particles of a resin which functions to impart to the cured form of the coating a reduced coefficient of friction. The patent discloses that the function of the chromium is to improve the corrosion-resistant properties of the cured coating, and the function of the resin, for example, polytetrafluoroethylene, is to increase the surface slip of the cured form of the coating. The '839 patent discloses the treatment of an uncured autodeposited coating with an acidic aqueous treating solution prepared by admixing a hexavalent chromium-containing compound (for example, ammonium and an alkali metal dichromate) with a hexavalent chromium/reduced chromium solution. In addition, the treating solution contains an acid or salt thereof, for example, hydrochloric acid, nitric acid, sulfuric acid, phosphoric acid, and ammonium, alkali metal, and alkaline earth metal salts of phosphoric acid. This patent discloses that the use of such a solution imparts a matte appearance to an autodeposited coating which otherwise would have a glossy appearance and improves the corrosion-resistant properties of the coating. In addition, U.S. Patent No. 3,647,567 discloses the use of an acidic aqueous solution of chromium trioxide or of water-soluble or acid-soluble chromates and dichromates to improve the corrosion resistance of the resinous coatings described therein. Exemplary chromates and dichromates are sodium, ammonium, lithium, magnesium, potassium

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and zinc.

Elsewhere, it is disclosed that the treatment of an uncured autodeposited coating with an aqueous solution or dispersion of a vulcanizing agent (for example, a sulfur-containing compound) or of a vulcanizing accelerator (for example, hexamethylenetetramine) to improve the solvent resistance of the cured coating.

Elsewhere, it is disclosed that adhesion of the freshly formed or wet coating to the underlying metallic substrate can be improved by contacting the coating with an acidic aqueous solution of an inorganic or organic acid or of an oxidizing agent (for example, sodium permanganate). This in turn leads to the provision of cured coatings which have a more uniform and appealing appearance. In addition to the use of chromium compounds, aforementioned U.S. Patent No. 3,647,567 teaches the use of an aqueous solution of phosphoric acid to improve the corrosion resistance of the resinous coating described therein.

In addition, it has been disclosed elsewhere that the treatment of an uncured autodeposited coating with an aqueous composition containing a water-miscible coalescing agent comprising a compound having two or more oxygen-containing functional groups such as ester groups, hydroxy groups, carbonyl groups and ether linkages. Examples of such classes of compounds include alcohols, ketones, alcohol esters, ketone esters, ketone ethers, and ester ethers. This Japanese patent discloses that the treatment of uncured autodeposited coatings with such coalescing agents inhibits or deters the tendency of the cured form of the coating to blister, crack and/or bridge.

It is an object of this invention to provide metallic surfaces, particularly surfaces that are made of one of the types of high carbon steel conventionally used for heavy duty springs and/or ferriferous surfaces that have been cold worked, especially by shot peening, grit blast-

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ing, or the like before being coated, with autodeposited coatings with better adhesion and/or better corrosion resistance than those obtained by following the teachings of the prior art.

5 DESCRIPTION OF THE INVENTION

In this description, except in the specific examples or where expressly indicated to the contrary, all numbers specifying amounts of materials or conditions of reaction or use are to be understood as modified by the term
10 "about" in determining the broadest scope of the invention. Practice of the invention within the exact numerical limits given is generally preferred.

Summary of the Invention

In a major embodiment of the present invention, im-
15 provements in properties of cured autodeposited coatings are achieved by contacting the uncured form of the coatings with an alkaline aqueous solution that also contains a component selected from the group consisting of anions of multifunctional organic acids, in an amount sufficient
20 to improve the corrosion resistance, adherence, or both corrosion resistance and adherence of the autodeposited coating after curing it. Organic acids are considered to be multifunctional for the purposes of this invention when their molecules each contain at least two electron
25 rich functional groups such as carboxyl and carbonyl, hydroxyl, ether, simple and substituted (but not quaternized) amino and amido, and phosphonyl. In general, molecules that contain different types of such functional groups are as useful as those that contain two or more of
30 the same functional group type. Non-limiting examples of suitable acids include citric, oxalic, tartaric, and diphosphonic acids.

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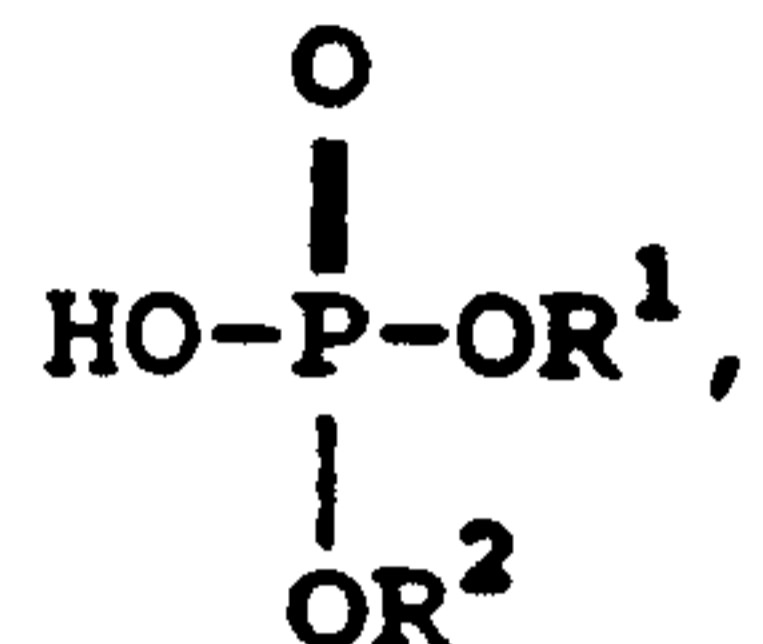
According to one aspect of the present invention, there is provided a process for forming an autodeposited organic coating on metallic parts of the surface of an object, said process comprising steps of contacting the
5 metallic surface to be coated with a liquid autodepositing composition to produce an uncured intermediate coating thereon and subsequently drying said uncured intermediate coating to produce the final autodeposited organic coating, characterized by contacting the uncured intermediate
10 coating, before drying it, with an aqueous adhesion and corrosion resistance promoting solution ("ACRPS") having a pH between 7 and 11 and comprising from 0.05 to 5 w/o of anions of multifunctional organic acids having two or more electron rich functional groups selected from the group
15 consisting of carboxyl; carbonyl; hydroxyl; ether; simple and substituted, but not quaternized, amino and amido; and phosphonyl groups.

An advantage of the present invention is that improvements in the properties of autodeposited coatings can
20 be realized by the use of a treating solution which does not require the presence of hexavalent chromium or a similarly toxic material which creates waste disposal prob-

lems.

Description of the Preferred Embodiments

One highly preferred type of acid from which anions needed in the treatment solutions according to this invention may be derived is the diphosphonic acids. The general formula of a phosphonic acid is:



where R^1 is a monovalent covalently bonded moiety containing at least one carbon atom and optionally also containing other functional groups, and R^2 is either a hydrogen atom or a monovalent covalently bonded moiety containing at least one carbon atom and optionally also containing other functional groups, and may be the same as R^1 or different. Anions for use in this invention are preferably derived from phosphonic acids in which R^2 in the formula above is hydrogen. More preferably, the anions used in this invention are derived from acids having at least two $(\text{H}_2\text{O}_3\text{P})$ groups attached to a single carbon atom, e.g., from 1,1-diphosphonic acids having the general formula $(\text{H}_2\text{O}_3\text{P})_2-\text{CR}^3\text{R}^4$, wherein each of R^3 and R^4 may be independently selected from hydrogen, hydroxyl, monovalent alkyl, monovalent substituted alkyl, and $(\text{H}_2\text{O}_3\text{P})$ groups. The most preferable anions are those of 1-hydroxyethylidene-1,1-diphosphonic acid, having the formula $\text{C}(\text{OH})(\text{CH}_3)(\text{PO}_3\text{H}_2)_2$.

Other preferred organic anions for use in the treating solutions according to this invention are anions derived from citric, tartaric, and oxalic acids.

The pH of the solution used for treating an uncured autodeposited coating according to this invention is between 7 and 11, preferably between 7.5 and 10, more preferably between 8.2 and 9.0. The concentration of the anions, expressed as their stoichiometric equivalent of the corresponding organic acid, is preferably between

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0.05 and 5 percent by weight ("w/o"), more preferably between 0.2 and 2 w/o, most preferably between 0.5 and 1.5 w/o.

5 In order to achieve the preferred pH values, the acid may be neutralized with a base, preferably a fugitive base, i.e., a base which volatilizes at or below the temperature used in curing of the autodeposited coating that is treated according to this invention, and additional base may be added to achieve an alkaline pH.
10 The most preferred base for use in preparing a treating solution according to this invention is ammonium hydroxide.

Higher organic acid anion concentrations and higher pH values within the ranges given above are generally preferred for higher film thickness of the autodeposited coating to be treated according to the invention. Un-
15 cured film thickness treated are preferably from 12 to 50 micrometers (" μ "), more preferably from 18 to 31 μ .

Preferred coatings which are treated according to the process of the present invention are formed from an autodepositing composition in which particles of resin are dispersed in an aqueous acidic solution which is prepared by combining hydrofluoric acid and a soluble ferric iron-containing ingredient, most preferable ferric fluoride.
20
25

U.S. Patent Nos. 4,347,172 and 4,411,937, which disclose the activating system preferred for use to form the autodeposited coatings to be treated according to this invention, disclose the optional use in the composition of an oxidizing agent in an amount to provide from about
30 0.01 to about 0.2 oxidizing equivalent per liter of composition. Suitable oxidizing agents are those commonly known as depolarizers. Examples of oxidizing agents are hydrogen peroxide, dichromate, permanganate, nitrate, persulfate, perborate, p-benzoquinone and p-nitrophenol.
35 Hydrogen peroxide is preferred.

Preferred resins for use in forming autodeposited

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coatings which are treated according to the present invention comprise internally stabilized vinylidene chloride copolymers or externally stabilized vinylidene chloride copolymers containing in excess of 50 w/o, or more preferably at least 80 w/o, of vinylidene chloride. Most preferably, the vinylidene chloride copolymer is crystalline in nature. Exemplary crystalline resins are described in U.S. Patent No. 3,922,451 and aforementioned U.S. Patent No. 3,617,368. Generally speaking, crystalline vinylidene chloride-containing resins comprise a relatively high proportion of vinylidene chloride, for example, at least about 80 wt. % thereof. However, any resin suitable for use in an autodepositing composition can be used to form a coating to be treated according to this invention.

Internally stabilized polymers or resins include as part of their chemical structure a surfactant group which functions to maintain polymer particles or resin solids in a dispersed state in an aqueous medium, this being the function also performed by an "external surfactant", that is, by a material which has surface-active properties and which is absorbed on the surface of resin solids, such as those in colloidal dispersion. As is known, the presence of an external surfactant tends to increase the water sensitivity of coatings formed from aqueous resin dispersions containing the same and to adversely affect desired properties of the coatings. The presence of undue amounts of surfactant in autodepositing compositions can lead to problems, as described in U.S. Patent No. 4,191,676. As discussed in this patent, the presence of an undue amount of surfactant in autodepositing compositions can deter the build-up of resin particles on the metallic surface being coated. In

addition, the presence of undue amounts of surfactant can also adversely affect desired coating properties, for example, corrosion resistant properties. An advantage of internally stabilized vinylidene chloride-containing polymers is that stable aqueous dispersions, including acidic aqueous dispersions of the type comprising autodepositing compositions, can be prepared without utilizing external surfactants. (It is noted that there is a tendency in the literature to use interchangeably the following terms in connection with describing surface active materials which are used in polymerization processes for preparing polymers of the type to which the present invention relates: surfactant, wetting agent, emulsifier or emulsifying agent, and dispersing agent. As used herein, the term "surfactant" is intended to be synonymous with the aforementioned.) Various types of internally stabilized vinylidene chloride-containing polymers are known and species thereof are available commercially. Examples of such latexes are the Saran latexes such as, for example, SARANTM 143 and SARANTM 112 available from W. R. Grace Co. and the SERFENETM latexes available from Morton Chemical. In accordance with the present invention, these commercial latexes can be used to excellent advantage, and internally stabilized latexes in general are preferred.

Various surfactants which function to maintain polymeric particles in dispersed state in aqueous medium include organic compounds which contain ionizable groups in which the anionic group is bound to the principal organic moiety of the compound, with the cationic group being a constituent such as, for example, hydrogen, an alkali metal, and ammonium. Speaking generally, exemplary anionic groups of widely used surfactants contain sulfur or phosphorous, for example, in the form of sulfates, thio-sulfates, sulfonates, sulfinates, sulfaminates, phosphates, pyrophosphates and phosphonates. Such surfactants comprise inorganic ionizable groups linked to an

organic moiety.

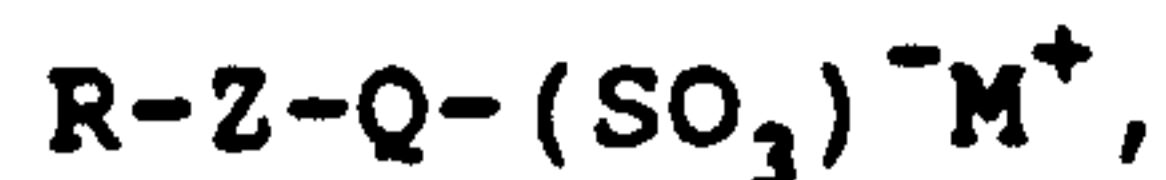
Although various ways may be used to introduce into the molecular structure of the vinylidene chloride resin such ionizable groups, it is believed that the most widely used method for preparing such resins will involve reacting vinylidene chloride with a monomeric surfactant and optionally one or more other monomers. In such a reaction, the monomeric surfactant comprises a material which is polymerizable with monomeric vinylidene chloride or with a monomeric material which is polymerizable with monomeric vinylidene chloride and which is ionizable in the reaction mixture and in the acidic aqueous medium comprising an autodepositing composition.

With respect to particular resins that can be used in the coating composition of the present invention, a preferred class can be prepared by copolymerizing (A) vinylidene chloride monomer with (B) monomers such as methacrylic acid, methyl methacrylate, acrylonitrile, and vinyl chloride and (C) a water soluble ionic material such as sodium sulfoethyl methacrylate. Although the constituents comprising the above-desired resin can vary over a relatively wide range, in general the resin will comprise the polymerized constituents in the following amounts:

- 1) between 45 and about 99 weight percent based on the total weight of monomers used of vinylidene chloride monomer;
- 2) from about 0.5 to 30 weight percent based on the total weight of (1) and (2) of a second relatively more hydrophilic ethylenically unsaturated monomeric material wherein such monomeric material has a solubility in both the water phase and the oil phase of the polymer latex of at least 1 weight percent at the temperature of polymerization; and
- 3) from about 0.1 to about 5 weight percent based on the total weight of other monomers of an ionic, significantly water-soluble material which is copolym-

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erizable with (2) and is selected from the group of sulfonic acids and their salts having the formula:



5 wherein the radical "R" is selected from the group consisting of vinyl and substituted vinyl, for example, alkyl-substituted vinyl; the symbol "Z" represents a difunctional linking group which will activate the double bond in the vinyl group; -Q- is a divalent hydrocarbon moiety having its valence bonds
10 on different carbon atoms; and the symbol "M⁺" represents a cation.

Examples of resins prepared from such monomers are disclosed in U.S. Patent No. 3,617,368.

The relatively hydrophilic monomers of (2) above include those materials which are readily copolymerizable
15 with (1) in aqueous dispersion, that is, which copolymerize within a period of about 40 hours at a temperature ranging from the freezing point of the monomeric serum up to about 100 ° C, and which have a solubility in both the
20 water and the oil phase of the polymer latex of at least 1 weight percent at the temperature of polymerization. Exemplary of preferred materials, particularly when used in conjunction with monomeric vinylidene chloride are methacrylic acid and methyl methacrylate. Other monomers
25 which may be advantageously employed include the hydroxyethyl and propyl acrylates, hydroxyethylmethacrylate, ethyl hexylacrylate, acrylic acid, acrylonitrile, methacrylonitrile, acrylamide, and the lower alkyl and dialkylacrylamides, acrolein, methyl vinyl ketone, and vinyl
30 acetate.

These monomers, which can be employed in amounts of from 0.5 to 30 weight percent, based on the total weight of the nonionic monomers used, provide for the necessary reactivity with the copolymerizable ionic material of (3)
35 and also provide for the required water solubility of the interpolymer in water. Thus, such materials may be referred to as "go-between" monomers. It is to be under-

stood that the optimum amount of such relatively hydrophilic monomers may vary somewhat within the prescribed range depending upon the amount of hydrophobic monomer used in preparing the resin, as well as upon the amount and type of the copolymerizable ionic monomer used.

The copolymerizable ionic monomers used in preparing the aforementioned type resins are those monomeric materials which contain in their structure both an ionizable group and a reactive double bond, are significantly soluble in water, are copolymerizable with the hydrophilic monomer constituent (2) and in which the substituent on the double bond is chemically stable under the conditions normally encountered in emulsion polymerization.

Examples of the aforementioned divalent hydrocarbon moiety having its valence bonds on different carbon atoms include alkylene and arylene divalent hydrocarbon radicals. Although the alkylene (CH_2) group can contain up to about 20 carbon atoms, it preferably has 2 to about 8 carbon atoms.

The solubility of the defined copolymerizable ionic material as described herein is strongly influenced by the cation M^+ . Exemplary cations are the free acids, alkali metal salts, ammonium and amine salts and sulfonium and quaternary ammonium salts. Preferred are the free acids, alkali metal salts, particularly sodium and potassium, and ammonium salts.

It is further noted that, with one of the ions above, and the usual choices for R and Z, the solubility of the monomer depends on Q. As indicated, this group can be either aliphatic or aromatic and its size will determine the hydrophilic/ hydrophobic balance in the molecule, that is, if Q is relatively small, the monomer is water soluble, but as Q becomes progressively larger, the surface activity of such monomer increases until it becomes a soap and ultimately a water insoluble wax. It is to be understood, however, that the limiting size of Q depends on R, Z, and M^+ . As exemplary of the above, it

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has been found that sodium sulfoethyl methacrylate is a highly acceptable copolymerizable ionic material for use in the present invention.

5 Further, the selection of R and Z is governed by the reactivity needed, and the selection of Q is usually determined by the reaction used to attach the sulfonic acid to the base monomer (or vice versa).

Processes for preparing latexes containing resins of the aforementioned type are known, such latexes being commercially available and being referred to herein as "self-stabilizing latexes", that is, latexes, the polymeric particles of which contain in the polymer molecule functional groups that are effective in maintaining the polymeric particles dispersed in the aqueous phase of the latex. As mentioned above, such latexes do not require the presence of an external surfactant to maintain the particles in their dispersed state. Latexes of this type generally have a surface tension very close to that of water (about 72 dynes/cm). It has been observed that autodepositing compositions containing such latexes form coatings which build up at a relatively fast rate.

An exemplary method for preparing such latexes involves preparation of an aqueous dispersion by an essentially continuous, carefully controlled addition of the requisite polymerization constituents (including polymerization initiator systems, if desired) to the aqueous medium having the desired pH value, followed by the subsequent addition of the necessary polymerization initiator, to form a polymeric seed latex in order to aid in the control of particle size. When forming such polymeric seed latexes, very small amounts of conventional surfactants, such as alkali soaps or the like, may be incorporated in the aqueous medium to further aid in the attainment of particles of desired size. The addition of such surfactants, however, is not critical for the production of the highly stable, internally stabilized, aqueous colloidal dispersions of polymeric particles of the type

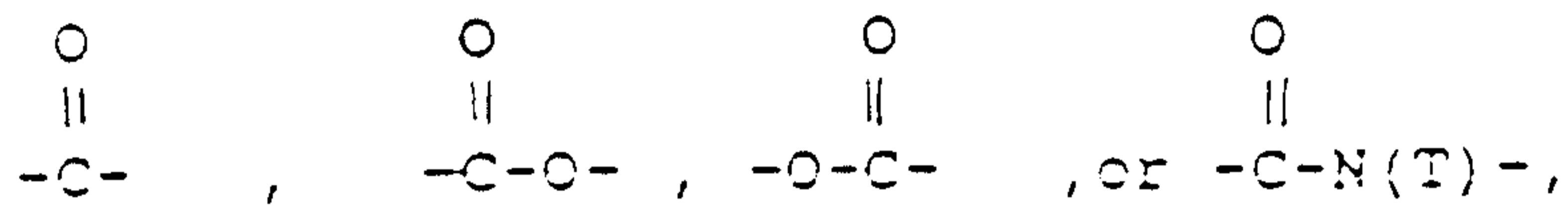
described above. In any event, additions of surfactants are limited so that the total amount present in the aqueous phase of the final coating solution is less than the critical micelle concentration, as taught in U.S. Patent No. 4,191,676. Following the formation of the polymeric seed latex, the remaining polymerization constituents are simultaneously and continuously added under carefully controlled conditions to the aqueous medium.

Highly stable polymer latexes for use in the present invention are characterized by the virtual absence of undesirable coagulum which often results when polymeric latexes are stabilized by conventional water soluble surfactants. Thus, such latexes combine the highly beneficial properties of optimum colloidal stability, reduced viscosities at relatively high polymer solids content, low foaming tendencies, and excellent product uniformity and reproducibility. Such highly stable latexes which are internally stabilized are disclosed, for example, in U.S. Patent No. 3,617,368.

A preferred embodiment of this invention comprises the use of vinylidene chloride-containing latexes in which a water soluble ionic material such as, for example, sodium sulfoethyl methacrylate is copolymerized with the comonomers comprising the copolymer. Sodium sulfoethyl methacrylate is particularly effective for use with monomeric vinylidene chloride and the relatively hydrophilic monomers methyl methacrylate or methacrylic acid when used in the amounts and in the manner described herein.

Particularly preferred latexes for use in this invention are latexes with about 35 to about 60 weight % solids comprising a polymeric composition prepared by emulsion polymerization of vinylidene chloride with one or more comonomers selected from the group consisting of vinyl chloride, acrylic acid, a lower alkyl acrylate (such as methyl acrylate, ethyl acrylate, butyl acrylate), methacrylic acid, methyl methacrylate, acryloni-

trile, methacrylonitrile, acrylamide, and methacrylamide and stabilized with sulfonic acid or sulfonic acid salt of the formula $R-Z-(CH_2)_n-(SO_3)^-M^+$, wherein R represents vinyl or lower alkyl-substituted vinyl; Z represents one
5 of the difunctional groups:



where T represents hydrogen or an alkyl group; n is an
10 integer from 1 to 20 (preferably 1 to 6), and M^+ is hydrogen or an alkali metal cation, preferably sodium or potassium.

A subgroup of preferred polymers are those having at least about 50% by weight of vinylidene chloride, but
15 less than about 70%, and about 5 to about 35% vinyl chloride, and about 5 to about 20% of a vinyl compound selected from the group consisting of acrylic acid, methyl acrylate, ethyl acrylate, butyl acrylate, methacrylic acid, methyl methacrylate, acrylonitrile, methacrylonitrile, acrylamide and methacrylamide, and combinations thereof, and about 1 to about 3% by weight of sulfoethyl methacrylate.

A particularly preferred group of latexes, however, are latexes containing about 30 to about 70 weight % of
25 solids formed by emulsion polymerization of about 50 to about 99 % vinylidene chloride based on total weight of polymer and about 0.1 to about 5% by weight of sulfoethyl methacrylate, with optionally other comonomers selected from the group consisting of vinyl chloride, acrylic and
30 methacrylic monomers such as acrylonitriles, acrylamides, methacrylamides and mixtures thereof in amounts between about 5 and about 50% by weight, and substantially free of unpolymerized surfactant or protective colloid.

Among other preferred subclasses of resin for use
35 prior to treatment according to this invention are dispersions of copolymers of about 50 to about 90% by weight of butyl acrylate and about 1 to about 2% by weight of sulfoethyl methacrylate based on the total weight of

polymer. Another preferred subclass of polymers are the latexes of vinylidene chloride-containing polymers internally stabilized with sulfoethyl methacrylate and free of surfactant, and including optionally vinyl chloride and one or more acrylic comonomers.

Another preferred vinylidene chloride-containing copolymer is one comprising about 15 to about 20 weight % vinyl chloride, about 2 to about 5 weight % butyl acrylate, about 3 to about 10 weight % acrylonitrile, about 1 to about 2 weight % sulfoethyl methacrylate. This particular copolymer will have less than 70% by weight vinylidene chloride copolymer based upon total weight of comonomers (including the sulfoethyl methacrylate) used in the emulsion polymerization.

The amount of the resin comprising the coating composition can vary over a wide range. The lower concentration limit of the resin particles in the composition is dictated by the amount of resin needed to provide sufficient material to form a resinous coating. The upper limit is dictated by the amount of resin particles which can be dispersed in the acidic aqueous composition. In general, the higher the amount of resin particles in the composition, the heavier the coating formed, other factors being the same. Although coating compositions can be formulated with a range of about 5 to about 550 g/l of resin solids, the amount of the resin solids will tend to vary depending on the other ingredients comprising the composition and also on the specific latex or resin used. For many applications, good results can be achieved by utilizing about 50 to about 100 g/l of resin solids in the composition.

Optional ingredients can be added to the composition as desired. For example, it is believed that the present invention will be used most widely in applications where it is desired to apply pigmented coatings to the metallic substrate. For this purpose, suitable pigments can be included in the composition. Examples of pigments that

can be used are carbon black, phthalocyanine blue, phthalocyanine green, quinacridone red, benzidine yellow, and titanium dioxide. The pigment should be added to the composition in an amount which imparts to the coating the
5 desired color and/or the desired depth or degree of hue. It should be understood that the specific amount used will be governed by the specific pigment used and the color of coating desired. Excellent results have been achieved by using the aqueous dispersion in an amount
10 such that the composition contains about 0.2 to about 3 g of furnace black/100 g of resin solids.

Many pigments are available in aqueous dispersions which may include surfactants or dispersing agents for maintaining the pigment particles in dispersed state.
15 When utilizing such pigment dispersions, they should be selected so that the surfactant concentration in the aqueous phase of the composition is below the critical micelle concentration ("CMC"), preferably below the surfactant concentration which corresponds to the inflection
20 point on a graph of surface tension versus the logarithm of surfactant concentration in the composition. Suitable pigmented compositions are illustrated in examples herein.

Colored coatings can be produced also by the use of
25 dyes, examples of which include rhodamine derived dyes, methyl violet, safranin, anthraquinone derived dyes, nigrosine, and alizarin cyanine green. These are but a few examples of dyes that can be used.

Examples of other additives that may be used in the
30 autodepositing composition are those generally known to be used in formulating paint compositions, for example, UV stabilizers, viscosity modifiers, etc.

If a surfactant is added to the composition, either
35 as a component of the latex, or with a pigment dispersion, or with other ingredients or additives, the total amount of surfactant in the aqueous phase of the composition should be maintained below the CMC. Preferably, the

aqueous phase of the composition contains little or no surfactant.

5 In case a surfactant is utilized, the preferred surfactants are the anionic surfactants. Examples of suitable anionic surfactants are the alkyl, alkyl/aryl or naphthalene sulfonates, for example, sodium dioctylsulfosuccinate and sodium dodecylbenzene sulfonate.

10 In preparing the autodepositing composition, the constituents thereof can be admixed in any suitable way, for example, as described in U. S. Patent No. 4,191,676. In preparing a bath of pigmented coating composition for use on an industrial scale, it is preferred that the bath be prepared by admixing:

15 A) an aqueous concentrate comprising about 350 to about 550 g/l of resin particles, preferable the aforementioned vinylidene chloride-containing resin particles, and about 10 to about 550 g/l of pigment; and

20 B) an aqueous concentrate prepared from about 0.4 to about 210 g/l of HF and a water soluble ferric-containing compound in an amount equivalent to about 1 to about 100 g/l of ferric iron.

25 The bath can be prepared by stirring water into concentrate (A) and thereafter admixing therewith the required amount of concentrate (B) with stirring to provide a homogenous composition.

30 Various steps of the overall coating process in which the present invention is used can be like those of the prior art, except as noted herein. For example, cleaning of the metallic surface prior to coating can be in accordance with the teachings of U.S. Patent No. 4,191,676. With respect to contacting the metallic surface with the autodepositing composition, it is believed that, for most applications, desired coating thicknesses
35 can be obtained by immersing the metallic surface in the composition for a period of time within the range of about 30 seconds or even less to about 3 minutes. Good

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5 results have been achieved utilizing a time of immersion
of not more than about 90 to about 120 seconds with com-
positions containing about 5 to about 10 wt % of resin
solids. However, it should be understood that longer or
shorter periods of time can be used. Agitating the com-
position aids in maintaining it uniform and in improving
the uniformity of the coatings formed. With other fac-
tors held constant, heating of the composition will re-
sult in heavier coatings. However, satisfactory results
10 can be obtained by operating the coating process at am-
bient temperature, and this is generally preferred for
convenience.

In a typical industrial process, the freshly applied
coating is rinsed with water after the coated surface has
15 been withdrawn from the composition and before signifi-
cant drying of the wet coating takes place. Such water
rinsing is effective in removing therefrom residuals,
such as acid and other ingredients of the composition
that adhere to the coated surface. If such residuals are
20 allowed to remain on the coated surface, they may ad-
versely affect the quality of the coating. Improvements
in rendering the cured form of the coating more imperme-
able to water, as provided by the present invention, are
not realized by simply water rinsing the freshly formed
25 coating.

Exemplary means for applying an adhesion and corro-
sion resistance promoting solution according to this in-
vention to the freshly formed coating include spray,
mist, and immersion, with the preferred means of applying
30 such solution being immersion of the uncured coated sur-
face in the solution for a period of time of about 5 sec-
onds to about 5 minutes.

The most preferred substrate for treatment according
to this invention is a conventional automobile leaf
35 spring made of high carbon steel and shot blasted on only
one side. Such shot blasting is believed to have at
least a slight effect on the electrochemical activity of

the steel, and the difference in such activity between the shot blasted and non shot blasted sides may have caused some of the difficulties noted in earlier attempts to use autodeposition for springs of this type.

5 The preferred activating system comprises a ferric ion-containing compound and hydrofluoric acid. Thus, a preferred autodepositing composition comprises a soluble ferric ion containing compound in an amount equivalent to about 0.025 to about 3.5 g/l ferric iron, most preferably
10 about 0.3 to about 1.6 g/l of ferric iron, and hydrofluoric acid in an amount sufficient to impart to the composition a pH within the range of about 1.6 to about 5.0. Examples of the ferric-containing compounds are ferric nitrate, ferric chloride, ferric phosphate, ferric oxide, and ferric fluoride, the last mentioned being preferred.
15

 It is preferable, as already note, if the alkaline components of the treatment solutions according to the invention are volatile or "fugitive". Aqueous ammonium
20 hydroxide and ammonium bicarbonate exemplify such fugitive bases, but the latter is less preferred, because when using it there is greater danger of blisters in the autodeposited coating after oven curing.

 After treatment according to this invention, the
25 coating should be cured. Fusion of the resinous coating renders it continuous, thereby improving its resistance to corrosion and its adherence to the underlying metallic surface.

 The conditions under which the curing and/or fusion
30 operation is carried out depend somewhat on the specific resin employed. In general, it is desirable to apply heat to fuse the resin, although some of the vinylidene chloride-containing resins described above can be cured at room temperature. Generally, the corrosion resistance, hardness and solvent resistance properties of coatings fused at elevated temperatures have been observed to
35 be better than coatings which have been air dried. How-

ever, there are applications where air dried coatings can be used satisfactorily. The fusion of the coating should be carried out under temperature and time conditions which do not adversely affect the desired properties of the coating. Exemplary conditions used in fusing the vinylidene chloride-containing coatings are temperatures within the range of about 20° C to 120° C for periods of time within the range of about 10 to 30 minutes, depending on the mass of the coated part. Baking the coating for a period of time until the metallic surface has reached the temperature of the heated environment has been used effectively.

When baked in an oven, the coating reaches the proper "curing" or heating temperature for the full development of coating properties when the metal part reaches that temperature. For this reason, parts that are constructed of thicker steel require longer times to reach the required temperature. For massive parts, it may not be possible to reach the required temperature without deleteriously affecting the coating and causing it to degrade.

In some cases, it is possible to overcome this problem by resorting to infrared radiation curing. In this case, it is possible to cure the coating without simultaneously raising the temperature of the metal to the required temperature. However, infrared radiation curing is practicable only for simple geometric shapes, since the area to be cured must be exposed to the infrared. In using infrared radiation curing, all coated surfaces must be accessible to the infrared source, that is, the entire coated surface must "see" the infrared.

The practice of this invention may be further appreciated from the following non-limiting examples and comparison examples.

Examples and Comparison Examples

The substrates coated for these examples were panels of high carbon spring steel as used for conventional aut-

omobile leaf springs. One side only of each panel had been shot blasted in a manner typical for the treatment of conventional automobile leaf springs before coating treatment was begun. The process sequence used was:

- 5 1. Spray clean for 75 seconds ("sec") at 60° C with a conventional aqueous alkaline cleaner having a free alkalinity of 6 - 15 milliliters ("ml") and a total alkalinity not more than 3 times the free alkalinity when a sample of 10 ml of the cleaner is titrated with 0.1 N HCl
- 10 solution, using phenolphthalein indicator for free alkalinity and bromphenol blue indicator for total alkalinity.
2. Allow to drain for 60 sec.
3. Dip clean for 150 sec at 65.6° C with a conventional
- 15 aqueous alkaline cleaner having a free alkalinity of 2 - 13 milliliters ("ml") and a total alkalinity not more than 3 times the free alkalinity when a sample of 10 ml of the cleaner is titrated with 0.1 N HCl solution, using phenolphthalein indicator for free alkalinity and
- 20 bromphenol blue indicator for total alkalinity.
4. Allow to drain for 60 sec.
5. Rinse with a tap water mist at 7 - 10 ° C for 30 sec.
6. Allow to drain for 15 sec.
- 25 7. Rinse with a deionized water mist at ambient temperature for 17 sec.
8. Allow to drain for 135 sec.
9. Dip coat for 145 sec in an autodeposition bath containing 1.8 grams per liter ("g/L") of ferric fluoride, 5
- 30 g/L of AQUABLACK™ 255 carbon black pigment (commercially available from Borden Chemical Company), sufficient solids from SARAN™ 143 latex to yield 5.2 ± 0.2 w/o of total solids in the bath, sufficient hydrogen peroxide to maintain an oxidation potential of 350 ± 20 millivolts more
- 35 oxidizing than a silver-saturated silver chloride reference electrode on a platinum measuring electrode immersed in the bath, and sufficient hydrofluoric acid to maintain

a reading of 250 ± 25 microamps on a LINEGUARD™ 101 Meter. (Note: For Comparison Example 2, a different autodeposition bath containing {styrene-acrylate} copolymer latex instead of poly{vinylidene chloride} was used in this step.)

- 5 10. Allow to drain for 135 sec.
11. Dip rinse in tap water at ambient temperature for 75 sec.
12. Allow to drain for 135 sec.
- 10 13. Dip for 75 sec at ambient temperature into an adhesion and corrosion resistance promoting treatment ("ACRPS") according to the invention or prior art, as specifically noted below.
14. Allow to drain for 180 sec.
- 15 15. Dry and cure in an oven at 110° for 25 minutes.

ACRPS compositions and test results are shown in Table 1.

TABLE 1

Ex. No.	ACRPS		Adhesion Test ² Results				Salt Spray Test ³ Results		Scribe/Scab Test ⁴ Results	
	Conc. ¹	pH	Initial		Final		S ⁵	N ⁶		
			S ⁵	N ⁶	S ⁵	N ⁶				
(Comparison) Examples with Uncured Coating Thickness 25-28 μ										
5	C1	7		5	10	12	38	VF+9	VF9	0.9
								0-1	0-1	
10	C2	8		0	0	0	0	N	VF+9	1.1
									R9.5	
	1	1.5	9.0	0	4	0	14	N	N	0.3
	2	1.5	7.5	5	6	3	23	N ⁹	10	0.9
	3	1.0	9.0	2	7	13	37	N ¹¹	0-1	1.0
15	4	0.5	8.2	8	9	5	16	N	0-2	0.7
(Comparison) Examples with Uncured Coating Thickness 18-21 μ										
	C3	12		5	2	48	26	VF9	n.m.	n.m.
	C4	13		76	12	82	17	N	n.m.	n.m.
	5	0.5	7.5	17	7	50	10	N ¹⁴	n.m.	n.m.
20	6	0.5	8.0	2	3	12	12	N	n.m.	n.m.
(Comparison) Examples with Uncured Coating Thickness 20-26 μ										
	C5 ¹⁵	16		100	40	100	35	n.m.	n.m.	n.m.
	C6 ¹⁵	17		50	15	100	25	N	n.m.	n.m.
	C7	7		0	5	15	5	N-VF8	VF-F6	n.m.
25	7	0.1	8.5	75	25	85	70	n.m.	n.m.	n.m.
	8	0.25	8.5	80	10	65	20	n.m.	n.m.	n.m.
	9	1.0	8.5	0	0	0	4	N ¹⁸	N ¹⁹	n.m.
	10	1.5	8.5	0	2	0	1	N	N-VF8	n.m.
	11	1.0	8.5	4	2	3	4	N	VF8	n.m.
30	12 ¹⁵	0.75	8.5	0	10	15	0	N	N	n.m.
	C8	20		5	5	40	15	N-VF8	n.m.	n.m.
	C9 ¹⁵	21		100	100	25	95	n.m.	n.m.	n.m.

(Notes for Table 1 are on the following page.)

Footnotes for Table 1

- 1 For the examples according to the invention (with numbers not prefixed by "C"), the concentration is in w/o of 1,1-hydroxyethylidene-1,1-diphosphonic acid for examples 1 - 6, in w/o of ammonium citrate for examples 7 - 10, in w/o of ammonium tartrate for example 11, in w/o of ammonium oxalate for example 12, and in w/o of sodium citrate for example 13. For the comparison examples (with numbers prefixed by "C"), the nature of the ACRPS is described in individual footnotes below.
- 2 Tested according to ASTM D0870-87 (Water Soak).
- 3 Tested according to ASTM B117-85, except that blistering ratings only were determined in some cases.
- 4 Tested according to Ford Motor Company "APG" test.
- 5 Measured on the shot peened side.
- 6 Measured on the non shot peened side.
- 7 ACRPS was about 0.1 N NaOH solution in water.
- 8 ACRPS was about 4 w/o sodium dichromate solution in water.
- 9 One of the three panels tested was 0-3 instead.
- 10 Three panels ranged from 0-1 to 0-5.
- 11 One of three panels tested blistered.
- 12 ACRPS was about 0.1 N NaOH solution in water.
- 13 ACRPS was about 0.1 N NH_4HCO_3 solution in water.
- 14 One of the three panels tested was rated VF9 instead.
- 15 Only one panel was tested for each of the values reported for this example, comparison example, or condition.
- 16 No solution was used; the samples were cured without rinsing.
- 17 ACRPS was deionized water.
- 18 One of the three panels tested for this condition was VF9 instead.
- 19 One of the three panels tested for this condition was VF6 instead.
- 20 ACRPS was 0.5 w/o sodium citrate solution in water, with a pH of 5.7.
- 21 ACRPS was 0.5 w/o aqueous solution of 1,1-hydroxyethylidene-1,1-diphosphonic acid, with a pH of about 2.

35

Other Notes for Table 1

"Initial" Adhesion was measured after drying but without any water soak according to GM 9071P method.

"Final" Adhesion was measured after soaking dried panels for 2 hours in water at 38° C.

40

"n.m." means not measured.

Values reported are for two or more panels for each test condition unless otherwise noted.

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CLAIMS:

1. A process for forming an autodeposited organic coating on metallic parts of the surface of an object, said process comprising steps of contacting the metallic surface to be coated with a liquid autodepositing composition to produce an uncured intermediate coating thereon and subsequently drying said uncured intermediate coating to produce the final autodeposited organic coating, characterized by contacting the uncured intermediate coating, before drying it, with an aqueous adhesion and corrosion resistance promoting solution ("ACRPS") having a pH between 7 and 11 and comprising from 0.05 to 5 w/o of anions of multifunctional organic acids having two or more electron rich functional groups selected from the group consisting of carboxyl; carbonyl; hydroxyl; ether; simple and substituted, but not quaternized, amino and amido; and phosphonyl groups.
2. The process according to claim 1, wherein the ACRPS comprises at least 0.05 w/o of anions derived from acids selected from the group consisting of 1,1-diphosphonic acids, citric acid, tartaric acid, and oxalic acid.
3. The process according to claim 2, wherein the ACRPS comprises from 0.2 to 2 w/o of anions derived from acids selected from the group consisting of 1,1-diphosphonic acids, citric acid, tartaric acid, and oxalic acid.
4. The process according to claim 3, wherein the ACRPS comprises from 0.5 to 1.5 w/o of anions derived from acids selected from the group consisting of citric acid, tartaric acid, oxalic acid, and 1-hydroxy-ethylidene-1,1-diphosphonic acid.

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5. The process according to claim 4, wherein the liquid autodepositing composition used comprises 1.8 g/L of ferric fluoride, 5 g/L of carbon black pigment, sufficient solids from a poly{vinylidene chloride} based latex to yield
5 from 5.0 to 5.4 w/o of total solids in an autodeposition bath comprising the liquid autodepositing composition, hydrogen peroxide in such an amount as to produce an oxidation potential of from 330 to 370 millivolts more oxidizing than a silver-saturated silver chloride reference
10 electrode on a platinum measuring electrode immersed in the autodeposition bath, and sufficient hydrofluoric acid to impart to the autodeposition bath a pH within the range from 1.6 to 5.0.

6. The process according to claim 4 or 5, wherein the
15 ACRPS comprises water, ammonia, ammonium ions, and multifunctional organic acid anions.

7. The process according to any one of claims 1 to 3, wherein the ACRPS comprises water, ammonia, ammonium ions, and multifunctional organic acid anions, and optionally,
20 bicarbonate and carbonate anions.

8. The process according to any one of claims 1, 2 and 7, wherein the metallic surface to be coated includes at least a portion which is a surface of high carbon spring steel or shot blasted carbon steel.

25 9. The process according to any one of claims 3 to 7, wherein the metallic surface to be coated is the surface of a leaf spring suitable for use in a conventional automobile.