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(54) Titre : COMPOSITIONS DE CEMENTS POUR PROTHESE DENTAIRE
(54) Title: DENTURE ADHESIVE COMPOSITIONS

(57) **Abrégé/Abstract:**

The present invention relates to a denture adhesive composition comprising mixed salts of an alkyl vinyl ether-maleic acid or anhydride copolymer and/or terpolymer with isobutylene wherein the mixed salt contains a cationic salt function comprising at least about 1 % strontium cations, from about 1 % to about 40 % of zinc cations, from 0 % to about 2.5 % of a cation selected from the group consisting of iron, boron, aluminum, vanadium, chromium, manganese, nickel copper, yttrium, titanium, and mixtures thereof; from about 36 % to about 60 % free acid component; and from 0 % to about 65 % of a cation selected from the group consisting of magnesium, calcium, and mixtures thereof. In addition, the present invention relates to denture adhesive compositions comprising the above composition and at least one non-adhesive self-supporting layer. The present invention further relates to a method of increasing the adhesion of dentures to the oral cavity by applying the above compositions to dentures, directly to the oral cavity, palate or ridge of the oral cavity, or applying it to both, and thereafter securing the dentures to the ridge or palate of the oral cavity.



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(54) Title: DENTURE ADHESIVE COMPOSITIONS (57) Abstract The present invention relates to a denture adhesive composition comprising mixed salts of an alkyl vinyl ether-maleic acid or anhydride copolymer and/or terpolymer with isobutylene wherein the mixed salt contains a cationic salt function comprising at least about 1 % strontium cations, from about 1 % to about 40 % of zinc cations, from 0 % to about 2.5 % of a cation selected from the group consisting of iron, boron, aluminum, vanadium, chromium, manganese, nickel copper, yttrium, titanium, and mixtures thereof; from about 36 % to about 60 % free acid component; and from 0 % to about 65 % of a cation selected from the group consisting of magnesium, calcium, and mixtures thereof. In addition, the present invention relates to denture adhesive compositions comprising the above composition and at least one non-adhesive self-supporting layer. The present invention further relates to a method of increasing the adhesion of dentures to the oral cavity by applying the above compositions to dentures, directly to the oral cavity, palate or ridge of the oral cavity, or applying it to both, and thereafter securing the dentures to the ridge or palate of the oral cavity.		

DENTURE ADHESIVE COMPOSITIONS

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BACKGROUND OF THE INVENTION

10 Ordinary removable dentures, dental plates and the like, comprise teeth mounted in a suitable plate or base. Denture stabilizers are used to fill the interstices between the dentures and the gums or tissues. Prior to placement of the denture in the oral cavity, a denture stabilizer is applied to the denture-plate surface which, for a perfect fit, should uniformly contact the gums and mucous tissues. The denture stabilizer is formulated not
15 only for its adherent properties, but also to provide a cushion or gasket between the denture and the gums or tissues, thereby positioning the denture securely in the oral cavity.

Considerable effort has been made over the years to develop improved denture adhesive compositions. Both synthetic and natural polymers and gums have been used
20 singly, in combination, and in combination with various adhesives and other materials in an attempt to lessen certain deficiencies. These deficiencies include inadequate holding power, oozing of the adhesive from under the dental plate during insertion and throughout the wearing period, and messiness and difficulty of removing the residual adhesive from the mouth and dentures. Additionally, food may become trapped between the denture and
25 the oral cavity of the wearer.

Alkyl vinyl ether-maleic copolymers and salts thereof are known in the art for use in denture adhesive compositions. Such disclosures include: U.S. Patent 3,003,988 to Germann et al., issued October 10, 1961; U.S. Patent 4,980,391 to Kumar et al., issued
December 25, 1990; U.S. Patent 5,073,604 to Holeva et al., issued December 17, 1991;
30 U.S. Patent 5,900,470, Prosise et al., issued May 4, 1999; U.S. Patent 5,037,924, Tazi et al., issued August 6, 1991; U.S. Patent 5,082,913, Tazi et al., issued January 21, 1992; and

U.S. Patent 5,525,652 to Clarke, issued June 11, 1996. In addition strip or insert denture adhesives are also known. For example, U.S. Patent 4,880,702 to Homan et al., issued 11/14/89 discloses a denture stabilizer in the form of a strip consisting of three layers. The two outside layers consist of a polymer selected from the group consisting of polyethylene oxide having an average molecular weight of about 200,000 to 10,000,000, sodium carboxymethylcellulose, polyvinyl alcohol, and mixtures thereof. The inside layer consists of microcrystalline wax and a polymer sufficient to adhere the inside layer to gums and a denture base, after contact with water, when the outside layers have been dissolved. European Patent Application 0,353,375 to Altwirth published 2/7/90, discloses an adhesive insert for dentures consisting of a adhesive permeated fibrous fleece and an adhesive consisting of a pasty mixture of alginate and/or carboxymethylcellulose, polyvinyl acetate and an alcoholic solvent. Despite the above-noted technologies as well as many others, a need still exists for denture stabilizing compositions providing improved hold.

In accordance with the present invention, it has been discovered that denture adhesive compositions comprising alkyl vinyl ether-maleic acid copolymers and/or terpolymers with isobutylene comprising strontium and zinc mixed salts together with specific levels of free acid, provide superior denture hold. The present denture adhesive compositions may also be effectively used as a wound dressing, underwater adhesive, a bioadhesive, a delivery vehicle for other actives, and/or any other application that requires adhesion to skin or tissue, including cosmetic pore cleansing strips.

An object of the present invention is to provide alkyl vinyl ether-maleic copolymers and/or terpolymers comprising strontium and zinc mixed salts, together with specific levels of free acid for adhesive compositions for firm hold, which may be used with dentures. These composition effectively hold dentures in place for a prolonged period of time.

SUMMARY OF THE INVENTION

The present invention relates to a denture adhesive composition comprising mixed salts of an alkyl vinyl ether-maleic acid copolymer and/or terpolymer with isobutylene wherein the mixed salt contains a cationic salt function comprising, or in the alternative consists essentially of, at least about 1% strontium cations, from about 1% to about 40%

of zinc cations, from 0% to about 2.5% of a cation selected from the group consisting of iron, boron, aluminum, vanadium, chromium, manganese, nickel, copper, yttrium, titanium, and mixtures thereof; from about 36% to about 60% free acid component; and from 0% to about 65% of a cation selected from the group consisting of magnesium, calcium, and mixtures thereof. In addition, the present invention relates to denture adhesive compositions comprising the above composition and at least one non-adhesive self-supporting layer. The present denture adhesive compositions can optionally comprise one or more additional adhesive components. The present invention further relates to a method of increasing the adhesion of dentures to the oral cavity by applying the above compositions to dentures, directly to the oral cavity, palate or ridge of the oral cavity, or applying it to both, and thereafter securing the denture to the ridge or palate of the oral cavity.

3a

In accordance with an aspect of the present invention, there is provided a denture adhesive composition comprising a mixed salt of a copolymer of alkyl vinyl ether and maleic acid or anhydride, the mixed salt containing a cationic salt function comprising:

- a) from 1% to 50% strontium cations;
- b) from 1% to 40% zinc cations;
- c) from 0% to 2.5%, of a cation selected from the group consisting of iron, boron, aluminum, vanadium, chromium, manganese, nickel, copper, yttrium, titanium, and mixtures thereof;
- d) from 42.5% to 60% free acid component; and
- e) from 0% to 55% of a cation selected from the group consisting of magnesium, calcium, and mixtures thereof;

wherein the copolymer is crosslinked only with inorganic metal cations.

In one embodiment of the present invention, the specific viscosity of the copolymer is greater than 5.5.

In accordance with a further aspect of the present invention, there is provided a denture adhesive composition comprising a mixed salt of a terpolymer of alkyl vinyl ether, maleic acid or anhydride, and isobutylene, the mixed salt containing a cationic salt function comprising:

- a) from 1% to 50% strontium cations;
- b) from 1% to 40% zinc cations;
- c) from 0% to 2.5% of a cation selected from the group consisting of iron, boron, aluminum, vanadium, chromium, manganese, nickel, copper, yttrium, titanium, and mixtures thereof;
- d) from 42.5% to 60% free acid component; and
- e) from 0% to 55% of a cation selected from the group consisting of magnesium, calcium, and mixtures thereof;

wherein the terpolymer is crosslinked only with inorganic metal cations.

In a further embodiment of the present invention, the denture adhesive composition further comprises one or more ingredients selected from the group consisting of

3b

additional adhesive components, plasticizers, colorants, preservatives, thickeners, vehicles, flavors, fragrances, sensates, and mixtures thereof.

In a further embodiment of the present invention, the denture adhesive composition further comprises an additional adhesive component selected from the group consisting of natural gums, synthetic polymers, mucoadhesive polymers, hydrophilic polymers, natural polymers, saccharide derivatives, cellulose derivatives, mixed salts of AVE/MA copolymers or terpolymers, AVE/MA copolymer acid, alkyl vinyl ether-maleic anhydride copolymer; and mixtures thereof.

In a further embodiment of the present invention, the cellulose derivatives are selected from the group consisting of hydroxyethylcellulose, hydroxypropylcellulose, hydroxypropylmethyl-cellulose, sodium carboxymethylcellulose, and mixtures thereof.

In a further embodiment of the present invention, the vehicle is selected from the group consisting of petrolatum, liquid petrolatum, mineral oil, glycerin, and mixtures thereof.

In a further embodiment of the present invention, the composition further comprises at least one non-adhesive self-supporting layer.

In a further embodiment of the present invention, the non-adhesive self-supporting layer is selected from the group consisting of polyester, polypropylene, nylon, rayon, cellulose acetate, non-adhesive cellulose derivatives, cloth, fibrous fleece, paper, plastic, leather, microcrystalline wax, synthetic fibers, natural fibers, and mixtures thereof.

In a further embodiment of the present invention, the denture adhesive composition comprises a percentage of strontium cations from 1% to 25%.

In a further embodiment of the present invention, the denture adhesive composition comprises a percentage of zinc cations from 10% to 30%.

3c

In a further embodiment of the present invention, comprises from 0.01% to 2% of a cation selected from the group consisting of iron, boron, aluminum, vanadium, chromium, manganese, nickel, copper, yttrium, titanium, and mixtures thereof.

In a further embodiment of the present invention, the denture adhesive composition comprises from 5% to 55% of a cation selected from the group consisting of magnesium, calcium, and mixtures thereof.

In a further embodiment of the present invention, the denture adhesive composition comprises from 15% to 50% of a cation selected from the group consisting of magnesium, calcium, and mixtures thereof.

In a further embodiment of the present invention, the cation is selected from the group consisting of iron, boron, aluminum, vanadium, chromium, manganese, nickel, copper, yttrium, titanium, and mixtures thereof, is iron.

In accordance with a further aspect of the present invention, there is provided a method of increasing the adhesion of dentures to the oral cavity by applying the denture adhesive composition to dentures, the oral cavity, or both, and thereafter securing the denture to the oral cavity.

DETAILED DESCRIPTION OF THE INVENTION

The denture adhesive compositions of the present invention comprise strontium
15 and zinc mixed salts of an alkyl vinyl ether-maleic copolymer and/or terpolymer with
isobutylene (preferably copolymers) with specific free acid levels, and optionally
comprising additional cations selected from the group consisting of iron, boron,
aluminum, vanadium, chromium, manganese, nickel, copper, yttrium, titanium, calcium,
magnesium, and mixtures thereof. The adhesive compositions may be in the form of a
20 powder, cream, paste, liquid, aerosol, and/or wafer. Powder forms are sprinkled on a
dental prosthesis, moistened and then inserted into the oral cavity. The compositions may
also be combined with various conventional delivery vehicles to form liquids or pastes
which are applied to a dental prosthesis and inserted into the oral cavity. These
compositions can optionally comprise at least one non-adhesive self-supporting layer.
25 Denture adhesive compositions with a self supporting layer are thoroughly moistened and
applied to dentures. A detailed description of essential and optional components of the
present invention is given below.

Definitions

The term "safe and effective adhesive amounts", as used herein, means an amount
30 sufficient to provide adherence to the oral cavity and/or adherence of a dental prosthesis

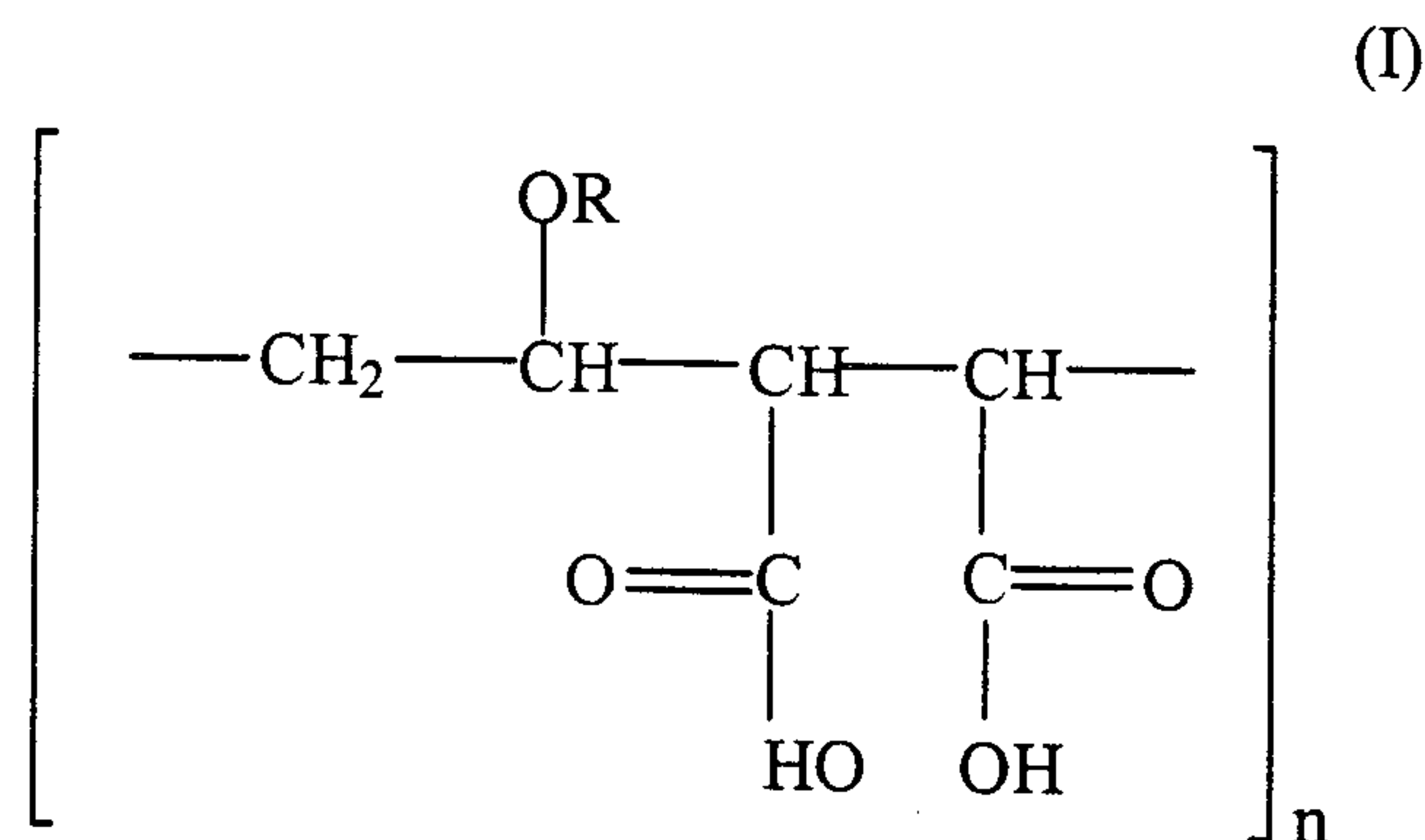
to the palate and ridge of the oral cavity, without toxicity to the user, damage to oral tissue, and alteration of the denture material.

The term "AVE/MA" as used herein refers to alkyl vinyl ether-maleic acid copolymer. The term "AVE/MA/IB" refers to terpolymers with alkyl vinyl ether, maleic acid or anhydride, and isobutylene. The term "mixed polymer salts" or "mixed salts", as used herein, refers to salts of AVE/MA or AVE/MA/IB where at least 2 different cations are mixed on the same polymer with each other or with other ester functions. The present invention comprises mixed polymer (copolymer or terpolymer) salts containing as essential salts: zinc and strontium cations. The term "free acid" ("FA") component as used herein refers either to the unreacted carboxyl groups (-COOH) of AVE/MA and/or AVE/MA/IB plus any other monovalent cations of carboxyl groups; e.g., -COONa, of the polymer. Monovalent cations include Group IA cations, such as sodium, potassium, hydrogen, etc. Preferably, the term "free acid" refers to the unreacted carboxyl groups (-COOH) of AVE/MA and/or AVE/MA/IB plus sodium and potassium cations. More preferably, the term "free acid" refers only to the unreacted carboxyl groups (-COOH) of the AVE/MA and/or AVE/MA/IB. The term "additional adhesive component", as used herein, refers to adhesives other than those described as essential AVE/MA and/or AVE/MA/IB mixed salts of the present invention.

The percentages used herein to describe the mixed salt function of the copolymers or terpolymer are defined as the stoichiometric percent of the total initial carboxyl groups reacted on the polymer. All other percentages used herein are by weight unless otherwise indicated.

Polymer

The alkyl vinyl ether-maleic acid copolymer comprises the repeated structural unit:



wherein R represents an alkyl radical, preferably a C₁ to C₅ alkyl radical, n is an integer greater than one representing the number of repeated occurrences of the structural unit in a molecule of the polymer.

The "terpolymer" or "terpolymer with isobutylene" means a terpolymer of maleic anhydride or acid, an alkyl vinyl ether (preferably with a C₁-C₅ alkyl radical), and isobutylene, having a structure of (A-B)_n where A is maleic anhydride or acid and B is alkyl vinyl ether (preferably with a C₁-C₅ alkyl radical), and/or isobutylene. The specific viscosity of the starting anhydride or acid of the terpolymer is preferably at least about 5.5, preferably at least about 6, preferably measured as a 1% weight/volume solution of methyl ethyl ketone at 25°C. The terpolymer must contain at least some isobutylene.

The present denture adhesive compositions comprise mixed salts of an AVE/MA copolymer and/or terpolymer wherein the mixed salt contains a cationic salt function comprising (or in the alternative consists essentially of) from about 1% to about 50% strontium cations, from about 1% to about 40% zinc cations, from 0% to about 2.5% of a cation selected from the group consisting of iron, boron, aluminum, vanadium, chromium, manganese, nickel, copper, yttrium, titanium, and mixtures thereof, and from 0% to about 65% of a cation selected from the group consisting of calcium, magnesium, and mixtures thereof, of the total initial carboxyl groups reacted, the mixed salt containing from about 36% to about 60% free acid component. The AVE/MA copolymers have a range of specific viscosities. For example, the specific viscosity is preferably from 1.2 to 14, as preferably measured as a 1% weight/volume solution of the starting anhydride or acid of the copolymer, in methyl ethyl ketone at 25°C. Other methods and solvents can be

used to measure the specific viscosity such as a 1% weight/volume solution in DMF (dimethyl formamide) at 25°C and a 1% weight/volume solution in 2-butanone at 25°C.

Preferably, the cationic salt function contains from about 2.5% to about 45% strontium cations, more preferably from about 5 % to about 25% strontium cations, of the initial carboxyl groups reacted. In addition, preferably the cationic salt function contains from about 10% to about 40% zinc cations , more preferably from about 10% to about 30% zinc cations, of the initial carboxyl groups reacted; preferably from about 0.001% to about 2.5 %, more preferably from about 0.01% to about 2% of a cation selected from the group consisting of iron, boron, aluminum, vanadium, chromium, manganese, nickel, copper, yttrium, titanium, and mixtures thereof, more preferably from about 0.01% to about 2% iron cations, of the initial carboxyl groups reacted; preferably from 5% to about 60% of a cation selected from the group consisting of calcium, magnesium, and mixtures thereof, of the total initial carboxyl groups reacted, more preferably from about 10% to about 55%, even more preferably from about 15% to about 50%. Preferably the cationic salt function contains from about 37.5% to about 55% free acid component, more preferably from about 37.5% to about 50%, of the total initial carboxyl groups.

The alkyl vinyl ether maleic anhydride copolymers are obtained by copolymerizing an alkyl vinyl ether monomer, such as methyl vinyl ether, ethyl vinyl ether, divinyl ether, propyl vinyl ether and isobutyl vinyl ether, with maleic anhydride to yield the corresponding alkyl vinyl ether-maleic anhydride copolymer which is readily hydrolyzable to the acid copolymer. Suitable copolymers may be prepared by well-known methods of the prior art, for example US 2,782,182, and US 2,047,398.

Both anhydride and acid forms are also available from commercial suppliers. For example, the ISP Corporation, Wayne, N.J. provides both the polymeric free acid form (I) and the corresponding anhydride form under its "GANTREZ" trademark as the "GANTREZ S Series" and "GANTREZ AN Series", respectively. When the anhydride copolymer dissolves in water, the anhydride linkage is cleaved so that the highly polar, polymeric free acid (I) is formed. Accordingly, the anhydride form, which is relatively less expensive than the acid form, may be used as a convenient and cheaper precursor for the acid. Elevated temperatures may be advantageously employed to enhance the rate of anhydride-to-acid hydrolysis. The

terpolymers can be made by the methods discussed in U.S. Patent 5,900,470, Prosise et al., issued May 4, 1999; U.S. Patent 5,037,924, Tazi et al., issued Aug. 6, 1991; and U.S. Patent 5,082,913, Tazi et al., issued Jan. 21, 1992.

5 The salt form of the subject polymers may be prepared by the interaction of the AVE/M anhydride or acid copolymer or terpolymer with at least one cationic salt function, such as strontium, zinc, calcium, sodium, potassium, iron, or ammonium compounds having a functional group typical of reactants of a carboxylic acid, such as, for example, the hydroxide, oxide, acetate, halide, lactate, etc. in an aqueous medium. In
10 a preferred embodiment, the zinc oxide and strontium carbonate are utilized. Mixed polymer salts comprising iron cations can be prepared by the interaction of the AVE/M anhydride/acid polymers with iron compounds, in the form of a salt, such as iron sulfate n-hydrate.

Ions that form toxic, irritating or contaminating by-products should be avoided, or
15 special precautions and treatment provided to assure the removal and absence of such by-products from the polymeric salt end-product. The particular compound used should be substantially pure to assure obtaining a substantially pure, polymeric salt end-product.

If the salt form of the polymer is desired, then an aqueous dispersion of particulate zinc oxide is combined with strontium carbonate and, optionally, ferric sulfate n-hydrate.
20 This is combined with the powder polymer, in the form of a slurry, in an amount sufficient to provide the desire cationic content desired in the end-product. This is done at ambient temperature and then slowly heated to 70°-95°C with continuous vigorous mixing so as to prevent localized precipitation of the cationic polymeric salt; mixing is continued to ensure that all the salt forming compound is reacted with the copolymer.

25 Alternatively, the polymer is hydrolyzed and neutralized in an aqueous mixture or slurry of one or more divalent and/or monovalent metal bases by heating the polymer/base mixture to a temperature ranging from about 45°C to about 100°C. Reaction of the mixed polymer salt with iron cations is obtained through addition of iron salts to the hydrolyzed and neutralized mixed salt of the polymer. Completion of the reaction with iron cations is
30 indicated by an increase in viscosity to stabilization. Alternatively, iron salts may be

blended with the polymer/metal base mixture prior to the hydrolysis and neutralization reactions.

In either of the above processes, the resulting slurry or solution is transferred to shallow stainless steel drying trays and placed in a forced air mechanical convection oven at 60-70°C for a time sufficient to evaporate the reaction medium (water) and remove water from the polymer (about 18-24 hours). Alternatively, the resulting slurry or solution can be drum-dried at 100° to 200°C with hot steam to evaporate the water content and recover the polymer in the flake form. After drying, the polymer forms brittle flakes which can easily be peeled off from the trays or drum surface and ground to a fine powder as desired to provide satisfactory denture stabilizing properties. Methods of making these mixed salts of AVE/MA polymers are further disclosed in U.S. Patent Nos. 5,073,604, Holeva et al., issued Dec. 17, 1991; 5,872,161, Liang et al., issued Feb. 16, 1999; 5,830,933, Synodis et al., issued Nov. 3, 1998.

The mixed salt polymers have exceptional adhesive qualities when contacted with water or saliva such that they are extremely useful as denture adhesive materials in denture compositions. The compositions of the present invention comprise a safe and effective adhesive amounts of the mixed salt polymers, preferably at least 20 percent by weight, and more preferably at least 30 percent by weight of the composition, as the sole adhesive component or as a co-adhesive in joint usage with other adhesive components.

Optional Non-Adhesive Self-Supporting Layer

The present denture adhesive compositions optionally comprise at least one non-adhesive self-supporting layer. The non-adhesive self-supporting layer is characterized by its ability to maintain strength and provide integrity for the adhesive composition in the presence of water and/or saliva. The non-adhesive self-supporting layer may include materials such as polyester, polypropylene, nylon, rayon, cellulose acetate, non-adhesive cellulose derivatives, cloth, fibrous fleece, paper, plastic, leather, microcrystalline wax, synthetic fibers, natural fibers, and mixtures thereof. Preferred are non-adhesive cellulose derivatives, polyester, polypropylene, nylon, rayon, cloth, paper, microcrystalline wax, and mixtures thereof. More preferred are polyester, polypropylene, rayon, nylon, cloth and paper.

The non-adhesive self-supporting layer may be in any physical form suitable for providing strength and/or integrity to the present adhesive compositions. Such physical forms include non-woven, woven, continuous, chopped, and combinations thereof. In addition, the non-adhesive self-supporting layer may be formed by any process commonly known in the art. Such processes include un-bonded, spraybonded, spun-bonded, needle-punched, carded, thermal bonded hydroentangled, meltblown, aperture print bonded, needled, wet-laid, dry-laid, and combinations thereof.

Other Adhesive Components

The present invention compositions may also include other adhesive components. These adhesive components, if present, are used in a safe and effective adhesive amounts. In general, the other adhesive components may be present at a level of from about 0% to about 90%, preferably from about 10% to about 70%, and most preferably from about 20% to about 50%, by weight of the composition.

Suitable adhesive components include a water-soluble hydrophilic colloid or polymer having the property of swelling upon exposure to moisture to form a mucilaginous mass. Such adhesive materials include natural gums, synthetic polymeric gums, AVE/MA copolymer acid, AVE/MA copolymer anhydride, AVE/MA/IB, synthetic polymers, mucoadhesive polymers, hydrophilic polymers, saccharide derivatives, cellulose derivatives, and adhesive materials commonly employed in denture stabilizing compositions and compatible with the polymers of the present invention, and mixtures thereof. Examples of such materials include karaya gum, guar gum, gelatin, algin, sodium alginate, tragacanth, chitosan, polyethylene glycol, acrylamide polymers, carbopol,* polyvinyl alcohol, polyamines, polyquaternary compounds, polybutenes, silicones, ethylene oxide polymers, polyvinylpyrrolidone, cationic polyacrylamide polymers.

Preferred are cellulose derivatives, polyethylene glycol, polyethylene oxide, karaya gum, sodium alginate, chitosan, polyvinyl alcohol, and mixtures thereof. Most preferred are cellulose derivatives such as methylcellulose, carboxymethylcellulose, hydroxyethylcellulose, hydroxypropylcellulose, hydroxy-propylmethylcellulose, and mixtures thereof.

Other Ingredients

*trade-mark

The present invention may also comprise a reducing agent. The reducing agent aids in removal of the denture from the oral cavity after application of the mixed polymer salt of the present invention to the denture. The preferred reducing agent for use herein is ascorbic acid and its water soluble salts. The reducing agent may also be used in
5 combination with a chelating agent. Preferred chelating agents include citrate, tartrate, lactate, and the like. The reducing agent and/or chelating agent may also be delivered in a composition by carriers known in the art which are safe for oral administration (i.e., non-toxic and approved for use in humans). Such carriers include surfactants, solvents, etc.

The reducing agent and/or chelating agents are used in safe and effect amounts.
10 The term "safe and effective amount", as used herein, means an amount sufficient to aid in releasing the denture hold in the oral cavity without toxicity to the user, damage to oral tissue, and alteration of the denture material. Thus, a denture wearer applies the mixed polymer salt adhesive composition to dentures or oral tissue and inserts the denture into the oral cavity. When removal is desired, the wearer swishes in the mouth, a denture
15 releasing composition comprising a reducing agent and/or chelating agents and suitable solvent(s) which aids in releasing the denture hold.

In addition one or more toxicologically-acceptable plasticizers may also be included in the present compositions. The term "toxicologically-acceptable", as used herein, is used to describe materials that are suitable in their toxicity profile for
20 administration to humans and/or lower animals. Plasticizers that may be used in the present compositions include dimethyl phthalate, diethyl phthalate, dioctyl phthalate, glycerin, diethylene glycol, triethylene glycol, Igepal®, Gafac®, sorbitol, tricresyl phosphate, dimethyl sebacate, ethyl glycolate, ethylphthalyl ethyl glycolate, o- and p-toluene ethyl sulfonamide, and mixtures thereof. Plasticizers may be present at a level of
25 from about 0% to about 70%, preferably from about 1% to about 30%, by weight of the compositions.

The present denture adhesive compositions which also comprise a non-adhesive self-supporting layer may also comprise a coating which is sticky to dry dentures and, if present, will be placed on one side of the denture adhesive composition. Compositions
30 suitable for use as this type of adhesive layer include polybutenes, silicones, rubbers, petrolatum, natural polymers, synthetic polymers, and mixtures thereof. The adhesive

layer may be present at a level of from about 0% to about 70%, and preferably from about 0.5% to about 20%, by weight of the composition.

Other suitable ingredients include colorants, preservatives such as methyl and propyl parabens; thickeners such as silicon dioxide, and polyethylene glycol; and vehicles
5 such as liquid petrolatum, petrolatum, mineral oil and glycerin. Preferred are polyethylene glycol, silicon dioxide, and petrolatum. Colorants, preservatives, thickeners and vehicles may be present at levels of from about 0% to about 20%, by weight of the composition.

The compositions of the present invention may also include one or more
10 components which provide flavor, fragrance, and/or sensate benefit (warming or cooling agents). Suitable components include natural or artificial sweetening agents, menthol, menthyl lactate, wintergreen oil, peppermint oil, spearmint oil, leaf alcohol, clove bud oil, anethole, methyl salicylate, eucalyptol, cassia, 1-menthyl acetate, sage, eugenol, parsley oil, oxanone, alpha-irisone, marjoram, lemon, orange, propenyl guaethol, cinnamon,
15 vanillin, thymol, linalool, cinnamaldehyde glycerol acetal known as CGA, and mixtures thereof, as well as coolants. The coolant can be any of a wide variety of materials. Included among such materials are carboxamides, menthol, ketals, diols, and mixtures thereof. Preferred coolants in the present compositions are the paramenthan carboxyamide agents such as N-ethyl-p-menthan-3-carboxamide, known commercially as
20 "WS-3", N,2,3-trimethyl-2-isopropylbutanamide, known as "WS-23," and mixtures thereof. Additional preferred coolants are selected from the group consisting of menthol, 3-1-menthoxypropane-1,2-diol known as TK-10 manufactured by Takasago, menthone glycerol acetal known as MGA manufactured by Haarmann and Reimer, and menthyl lactate known as Frescolat® manufactured by Haarmann and Reimer. The terms menthol
25 and menthyl as used herein include dextro- and levorotatory isomers of these compounds and racemic mixtures thereof. TK-10 is described in U.S. Pat. No. 4,459,425, Amano et al., issued 7/10/84. WS-3 and other agents are described in U.S. Pat. No. 4,136,163, Watson, et al., issued Jan. 23, 1979.

These agents may be present at a level of from about 0% to
30 about 50%, by weight of the composition.

The present compositions may be used as a denture adhesive and/or used as a bioadhesive on wet tissue such as mucosal tissues, wounds, oral mucosa, etc. The present adhesive compositions can be used to deliver one or more therapeutic actives suitable for topical administration to mucosal or wet tissues. The phrase "therapeutic actives", as used herein, describes agents which are pharmacologically active when absorbed through wet tissue or mucosal surfaces of the body such as the oral cavity, wounds, or applied to the surfaces of the skin. Therapeutic actives may be present at a level of from about 0% to about 70%, by weight of the composition.

Therapeutic actives that are useful in the present compositions include antimicrobial agents such as iodine, sulfonamides, bisbiguanides, or phenolics; antibiotics such as tetracycline, neomycin, kanamycin, metronidazole, or clindamycin; anti-inflammatory agents such as aspirin,* acetaminophen, naproxen and its salts, ibuprofen, ketorolac, flurbiprofen, indomethacin, eugenol, or hydrocortisone; dentinal desensitizing agents such as potassium nitrate, strontium chloride or sodium fluoride; anesthetic agents such as lidocaine or benzocaine; anti-fungals; aromatics such as camphor, eucalyptus oil, flavors, fragrances, or sensates (warming or cooling agents), and aldehyde derivatives such as benzaldehyde; insulin; steroids; and anti-neoplastics. It is recognized that in certain forms of therapy, combinations of these agents in the same delivery system may be useful in order to obtain an optimal effect. Thus, for example, an antimicrobial and an anti-inflammatory agent may be combined in a single delivery system to provide combined effectiveness.

Process for Preparation of the Composition

A process for preparing denture adhesive compositions of the present invention (creams, powders, wafers, liquids, aerosols, pastes) comprises conventional methods disclosed in the art. Conventional methods are taught in US 5,525,652, issued June 11, 1996, Clarke et al.; US 3,003,988, issued Oct. 10, 1961, Germann et al.; US 5,073,604, Holeva et al., issued Dec. 17, 1991; and US 5,872,161, Liang et al., issued Feb. 16, 1999.

A process for the preparation of the present denture adhesive compositions optionally comprising a non-adhesive self-supporting layer, comprises coating a weighed amount of the adhesive components onto the non-adhesive self-supporting layer. This

*trade-mark

process is disclosed in US 5,877,233, Liang et al, issued March 2, 1999; US 5,872,160, issued 2/16/99, Liang et al.; US 5,880,172, Rajaiah et al., filed Oct. 25, 1996.

The term "mixture", as used in this "Process for Preparation the Composition"
5 section, refers to a solution, slurry, or suspension.

The adhesive components may be coated on the non-adhesive self-supporting layer using various methods. These include: (a) wetting the non-adhesive self-supporting layer with water, uniformly sifting the adhesive component powder(s) onto the wet layer and then rewetting the layer with water; (b) dissolving the adhesive component(s) in
10 water and/or other solvent(s) and coating the resulting mixture on the layer; (c) coating the layer with the mixture produced during AVE/MA polymer processing; (d) incorporating the adhesive component(s) into the layer as the layer is being formed; and (e) dissolving the adhesive component(s) in water and/or other solvent(s), wetting/coating the resulting mixture onto the layer, and uniformly sifting one or more adhesives in
15 powder form onto the wet/coated layer and optionally re-coating/re-wetting the layer with the mixture and/or water; (f) the method of step (e) repeated multiple times; and (g) any combination of the methods in (a) through (f) above.

As disclosed above, the adhesive components may be dissolved in water and/or other solvents and the resulting mixture coated onto the layer. Solvents for the polymers
20 include water and/or alcohols such as methanol, propanol, isopropanol, ethanol, butanol, 1,4-butanediol, cyclohexanol, and diethylene glycol; ethers or ether alcohols such as tetrahydrofuran, ethylene glycol monomethyl ether, diethylene glycol monomethyl ether, dioxane, and ethyl ether; esters such as methyl acetate, ethyl acetate and sec-butyl acetate; aldehydes, ketones or ketone-alcohols such as benzaldehyde, formaldehyde solution,
25 methyl ethyl ketone, diacetone alcohol, acetone, cyclohexanone, mesityl oxide, and methyl isobutyl ketone; lactams or lactones such as N-methyl-2-pyrrolidone, N-vinyl-2-pyrrolidone, 2-pyrrolidone, and butyrolactone; hydrocarbons such as benzene, toluene, xylene, hexane, mineral spirits, mineral oil, and gasoline; chlorinated hydrocarbons such as carbon tetrachloride, chlorobenzene, chloroform, ethylene dichloride, methylene
30 chloride; nitroparaffins such as nitroethane, and nitromethane; mercaptans such as

thiophenol and 2-mercapto-1-ethanol; and others such as acetic acid, pyridine and dimethyl formamide.

Preferred solvents for the polymers are water, methanol, propanol, isopropanol, tetrahydrofuran, methyl acetate, benzaldehyde, formaldehyde solution, methyl ethyl
5 ketone, diacetone alcohol, N-methyl-2-pyrrolidone, N-vinyl-2-pyrrolidone, dimethyl formamide, and mixtures thereof. Compounds commonly used as plasticizers can also be used as solvents for the polymers. Such plasticizers include dimethyl phthalate, diethyl phthalate, dioctyl phthalate, glycerin, diethylene glycol, triethylene glycol, Igepal® CO-630, Gafac ® RE-610, sorbitol, tricresyl phosphate, dimethyl sebacate, ethyl glycolate,
10 ethylphthalyl ethyl glycolate, and p-toluene ethyl sulfonamide.

Solvents for other adhesives such as CMC, which may be optionally included in the adhesive compositions include mixtures of water and water-miscible solvents such as ethanol and acetone. Solutions of low concentration can be made with up to 40% acetone and/or 50% alcohol. Other solvents which made be used include ethanolamines; ethylene
15 glycol; glycerol; 1,2,6-hexanetriol; mono-, di-, and triacetin; 1,5-pentanediol; polyethylene glycol (molecular weight 600 or less); propylene glycol; and trimethylolpropane.

When the adhesive compositions are prepared by dissolving the adhesive component(s) in water and/or other solvents, various embodiments of the process
20 includes: dissolving the polymers in one or more of the solvents for polymers; dissolving an optional adhesive in a suitable solvent and coating the resulting mixture onto the non-adhesive self-supporting layer and then optionally sifting one or more adhesives onto the coated layer. Coating the layer can be achieved by techniques commonly known in the art including extrusion, doctor blading, spraying, dipping, etc.

25 After the polymer has been deposited on the layer by one of the means described above, the layer is then dried. Next, the denture adhesive composition is mechanically softened by running it through a ring-roller or micro-cracker or any other suitable means. The composition is then pressed smooth in a hydraulic press or flat-roller or other suitable means. The composition is then die-cut into denture shapes. These shapes may facilitate
30 application of the composition to the dentures.

The following examples further describe and demonstrate embodiments within the scope of the present invention. The examples are given solely for the purpose of illustration and are not to be construed as limitations of the present invention. Many variations of these are possible without departing from the spirit and scope of the invention.

EXAMPLE I (A-O)

The following examples A to O describe various mixed salts and the process for making these mixed salts of AVE/MA polymers:

	A	B	C	D
Component	(In Grams)	(In Grams)	(In Grams)	(In Grams)
Water	2850	2850	2841	2850
Calcium Hydroxide		17.08	17.08	8.54
Strontium Carbonate	39.71	5.67	11.35	39.71
Zinc Oxide	12.51	12.51	10.95	3.13
AVE/MA Anhydride Polymer ¹	120	120	120	120
Resulting Mixed Salt	Sr(35%)/Zn(20%)/FA(45%)	Ca (30%)/Sr (5%)/Zn(20%)/FA(45%)	Ca(30%)/Sr (10%)/Zn(17.5%)/FA(42.5%)	Ca(15%)/Sr (35%)/Zn(5%)/FA(45%)

	E	F	G	H
Component	(In Grams)	(In Grams)	(In Grams)	(In Grams)
Water	2842	2850	2850	2850
Calcium	14.24	5.13		2.7

¹ Gantrez AN 169 available from ISP Corporation, and/or AVE/MA copolymer with specific viscosity of 6.3 and/or 7.3.

Hydroxide				
Strontium Carbonate	11.35	39.71	39.71	11.35
Zinc Oxide	12.51	12.51	3.13	12.51
Ferric Sulfate, N-Hydrate				0.34
Sodium Hydroxide				6.15
Magnesium Oxide				3.1
AVE/MA Anhydride Polymer ¹	120	120	120	120
Resulting Mixed Salt	Ca (25%)/Sr (10%)/Zn (20%)/FA(45%)	Ca (9%)/Sr (35%)/Zn (20%)/FA(36%)	Sr (35%)/Zn (5%)/FA(60%)	Ca (4.75%)/Sr (10%)/Zn (20%)/Fe (0.25%)/FA (55%)/Mg(10%)
	I	J	K	L
Component	(In Grams)	(In Grams)	(In Grams)	(In Grams)
Water	2844	2835	2837	2832
Calcium Hydroxide	14.24	9.97	11.39	
Strontium Carbonate	5.67	22.69	22.69	36.87
Zinc Oxide	12.51	12.51	3.13	10.95
Magnesium Oxide	3.10			
AVE/MA Anhydride Polymer ¹	120	120	120	120
Resulting	Ca (25%)/Sr	Ca (17.5%)/Sr	Sr (35%)/Zn	Sr(32.5%)/Zn(17

Mixed Salt	(5%)/Zn (20%)/Mg(10%)/ FA(40%)	(20%)/Zn (20%)/FA(42.5%)	(5%)/FA(60%)	.5%)/FA(50%)
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	M	N	O
Component	(In Grams)	(In Grams)	(In Grams)
Water	2842	2836	2823
Calcium Hydroxide	19.93	8.54	
Strontium Carbonate	5.67	22.69	51.06
Zinc Oxide	12.51	12.51	6.25
AVE/MA Anhydride Polymer ¹	120	120	120
Resulting Mixed Salt	Ca (35%)/Sr (5%)/Zn (20%)/FA(40%)	Ca (15%)/Sr (20%)/Zn (20%)/FA(45%)	Sr (45%)/Zn (10%)/FA(45%)

Weigh the above components and add to a 4 liter reaction vessel while mixing. Use 15% of water to pre-slurry all powders except the AVE/MA. Wash down the residual powders from the wall of the vessel. Then react the mixture at 80 to 95 °C. Dry the solution in trays in an oven at 60 to 70 °C. Mill all dried flakes to a fine powder. Equivalent mixed salts of AVE/MA/IB can be obtained by the same method as above, substituting the AVE/MA/IB for AVE/MA, at various levels.

EXAMPLE II

Denture stabilizing compositions in cream form can be made by blending together the following ingredients:

	weight (grams)
White Mineral Oil	89.74
Petrolatum, White	82.01
Carboxymethylcellulose Sodium	75.00
Silicon Dioxide, Colloidal	4.28
Colorant (Opatint* Red Dye)	0.23
Any AVE/MA mixed salt of A-O or any AVE/MA/IB mixed salt of A-O.	123.75

- Weigh, heat and mix the red dye, petrolatum, and mineral oil in a glass jar at 50 to 60 °C until visually uniform. Then weigh and shake-blend the powders (colloidal silicon dioxide, CMC, AVE/MA copolymer mixed salt or AVE/MA/IB mixed salt) together in a container.
- 5 Thereafter, mix the powders into the liquid with a spatula until visually a uniform pink cream. The above cream can be modified by increasing or decreasing the level of AVE/MA mixed salt or AVE/MA/IB mixed salt by 0 to 18.75 grams, petrolatum by 0 to 18.75 grams, and/or the CMC by 0 to 18.75 grams. The above cream composition can also be modified by using mixtures of the various AVE/MA mixed polymer salts and/or AVE/MA/IB mixed
- 10 salts. The subject places from 0.1 to 2 grams of the cream composition on the denture. Then the subject inserts the denture into his/her mouth and presses it into place.

EXAMPLE III

Denture stabilizing compositions in powder form can be made by blending together the following ingredients:

15

	weight (grams)
Carboxymethylcellulose Sodium	40.00
Any AVE/MA mixed salt of A-O or any AVE/MA/IB mixed salt of A-O	60.00

*trade-mark

Blend all components together. The above compositions can be modified by increasing or decreasing the AVE/MA mixed salt or AVE/MA/IB mixed polymer salt by 0 to 50 grams and/or the CMC by 0 to 40 grams. The above powder compositions can also be modified by using mixtures of the various AVE/MA mixed salts and/or AVE/MA/IB mixed salts.

- 5 The subject places from 0.1 to 2 grams of the composition on a pre-moistened denture, allowing it to hydrate briefly. Then the subject inserts the denture into his/her mouth and presses it into place.

EXAMPLE IV

- 10 Denture stabilizing compositions in wafer form can be made by wetting a 58" by 20" non-woven polyester (non-adhesive self-supporting layer) with water. Uniformly coat this wet sheet with the compositions listed below. Thereafter, rewet the layer with water. Dry the layer. Mechanically soften the composition by ring-roller, and then smooth the composition on a hydraulic press. Die-cut the composition into desired shapes. Moisten and apply these wafer compositions to the dentures. Then insert the denture into the
- 15 mouth and press it into place.

	weight (grams)
Carboxymethylcellulose Sodium	60.00
Any AVE/MA mixed salt of A-O or any AVE/MA/IB mixed salt of A-O.	90.00

- These wafer compositions can be modified by increasing or decreasing the AVE/MA mixed polymer salt or AVE/MA/IB mixed polymer salt by 0 to 60 grams and/or CMC by
- 20 0 to 60 grams. The above wafer compositions can also be modified by using mixtures of the various AVE/MA mixed salts and/or AVE/MA/IB mixed polymer salt.

WHAT IS CLAIMED IS:

1. A denture adhesive composition comprising a mixed salt of a copolymer of alkyl vinyl ether and maleic acid or anhydride, the mixed salt containing a cationic salt function comprising:
 - a) from 1% to 50% strontium cations;
 - b) from 1% to 40% zinc cations;
 - c) from 0% to 2.5%, of a cation selected from the group consisting of iron, boron, aluminum, vanadium, chromium, manganese, nickel, copper, yttrium, titanium, and mixtures thereof;
 - d) from 42.5% to 60% free acid component; and
 - e) from 0% to 55% of a cation selected from the group consisting of magnesium, calcium, and mixtures thereof;wherein the copolymer is crosslinked only with inorganic metal cations.
2. The denture adhesive composition according to Claim 1 wherein the specific viscosity of the copolymer is greater than 5.5.
3. A denture adhesive composition comprising a mixed salt of a terpolymer of alkyl vinyl ether, maleic acid or anhydride, and isobutylene, the mixed salt containing a cationic salt function comprising:
 - a) from 1% to 50% strontium cations;
 - b) from 1% to 40% zinc cations;
 - c) from 0% to 2.5% of a cation selected from the group consisting of iron, boron, aluminum, vanadium, chromium, manganese, nickel, copper, yttrium, titanium, and mixtures thereof;
 - d) from 42.5% to 60% free acid component; and
 - e) from 0% to 55% of a cation selected from the group consisting of magnesium, calcium, and mixtures thereof;wherein the terpolymer is crosslinked only with inorganic metal cations.

4. The denture adhesive composition according to Claim 1 or 3 further comprising one or more ingredients selected from the group consisting of additional adhesive components, plasticizers, colorants, preservatives, thickeners, vehicles, flavors, fragrances, sensates, and mixtures thereof.
5. The denture adhesive composition according to Claim 4 further comprising an additional adhesive component selected from the group consisting of natural gums, synthetic polymers, mucoadhesive polymers, hydrophilic polymers, natural polymers, saccharide derivatives, cellulose derivatives, mixed salts of AVE/MA copolymers or terpolymers, AVE/MA copolymer acid, alkyl vinyl ether-maleic anhydride copolymer; and mixtures thereof.
6. The denture adhesive composition according to Claim 5 wherein the cellulose derivatives are selected from the group consisting of hydroxyethylcellulose, hydroxypropylcellulose, hydroxypropylmethyl-cellulose, sodium carboxymethylcellulose, and mixtures thereof.
7. The denture adhesive composition according to Claim 4 wherein the vehicle is selected from the group consisting of petrolatum, liquid petrolatum, mineral oil, glycerin, and mixtures thereof.
8. The denture adhesive composition according to Claim 1 or 3 wherein the composition further comprises at least one non-adhesive self-supporting layer.
9. The denture adhesive composition according to Claim 8 wherein the non-adhesive self-supporting layer is selected from the group consisting of polyester, polypropylene, nylon, rayon, cellulose acetate, non-adhesive cellulose derivatives, cloth, fibrous fleece, paper, plastic, leather, microcrystalline wax, synthetic fibers, natural fibers, and mixtures thereof.

10. The denture adhesive composition according to any one of claims 1 to 9 comprising a percentage of strontium cations from 1% to 25%.
11. The denture adhesive composition according to any one of claims 1 to 10 comprising a percentage of zinc cations from 10% to 30%.
12. The denture adhesive composition according to any one of claims 1 to 11 comprising from 0.01% to 2% of a cation selected from the group consisting of iron, boron, aluminum, vanadium, chromium, manganese, nickel, copper, yttrium, titanium, and mixtures thereof.
13. The denture adhesive composition according to claim 3, comprising from 5% to 55% of a cation selected from the group consisting of magnesium, calcium, and mixtures thereof.
14. The denture adhesive composition according to any one of claims 1 to 13, comprising from 15% to 50% of a cation selected from the group consisting of magnesium, calcium, and mixtures thereof.
15. The denture adhesive composition according to claim 1 or claim 2 wherein the cation selected from the group consisting of iron, boron, aluminum, vanadium, chromium, manganese, nickel, copper, yttrium, titanium, and mixtures thereof, is iron.
16. A method of increasing the adhesion of dentures to the oral cavity by applying the composition of any one of claims 1 to 15 to dentures, the oral cavity, or both, and thereafter securing the denture to the oral cavity.